The h-Current in the Substantia Nigra pars Compacta Neurons: A Re-examination

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Abstract

The properties of the hyperpolarization-activated cation current (I_h) were investigated in rat substantia nigra - pars compacta (SNc) principal neurons using patch-clamp recordings in thin slices. A reliable identification of single dopaminergic neurons was made possible by the use of a transgenic line of mice expressing eGFP under the tyrosine hydroxylase promoter. The effects of temperature and different protocols on the I_h kinetics showed that, at 37 °C and minimizing the disturbance of the intracellular milieu with perforated patch, this current actually activates at potentials more positive than what is generally indicated, with a half-activation potential of −77.05 mV and with a significant level of opening already at rest, thereby substantially contributing to the control of membrane potential, and ultimately playing a relevant function in the regulation of the cell excitability. The implications of the known influence of intracellular cAMP levels on I_h amplitude and kinetics were examined. The direct application of neurotransmitters (DA, 5-HT and noradrenaline) physiologically released onto SNc neurons and known to act on metabotropic receptors coupled to the cAMP pathway modify the I_h amplitude. Here, we show that direct activation of dopaminergic and of 5-HT receptors results in I_h inhibition of SNc DA cells, whereas noradrenaline has the opposite effect. Together, these data suggest that the modulation of I_h by endogenously released neurotransmitters acting on metabotropic receptors –mainly but not exclusively linked to the cAMP pathway- could contribute significantly to the control of SNc neuron excitability.

Introduction

The presence of the h-current is a hallmark of midbrain dopaminergic (DA) neurons, including those of the substantia nigra pars compacta (SNc), up to the point that its occurrence is considered by many authors the main discriminating criterion to decide if a given neuron in this area is dopaminergic or not [1]. Many studies have confirmed the close relationship between the DAergic phenotype and I_h expression [2–6]. Apart from the abundant electrophysiological evidence ([7–9], to cite a few), the presence of h-channels in SNc neurons is also supported by qualitative RT-PCR experiments on single cells, which revealed that SNc neurons co-express three of the four types of HCN subunits: HCN2, HCN3, and HCN4 [10].

As expected, the presence of a current typically associated with the pacemaking process [11] for a review suggests that it could play its archetypal role also in SNc neurons, cells characterized by autorhythmicity. However, several studies reported that I_h has neither a significant role in spontaneous pacemaker activity nor does it contribute substantially to the setting of the resting potential [9,12–15].

Overall, the present knowledge of the h-current in SNc neurons is not entirely satisfactory, and this is all the more surprising for a population of neurons which is object of so many studies. The inconsistencies in the description of I_h are probably due to the strong dependence of the kinetics of this current on experimental conditions (e.g., temperature, patch configuration, ionic composition of solutions, modulation by cytoplasmic cyclic nucleotides, protocols used, etc.). This circumstance may explain why, even for a single cell type, different kinetics were found by different laboratories, and consequently different roles were proposed. In addition, there might be a problem in the cell identification: as a rule, cells in the midbrain are identified as dopaminergic on the basis of a series of electrophysiological characteristics, confirming a posteriori the identification in few randomly chosen cells with immunohistochemistry to ascertain the presence of TH. However, some of the more commonly used identification criteria are not really discriminative. For example, the presence of I_h -considered a benchmark- can be misleading, as if the absence of this current in a midbrain neuron is a trustworthy predictor that the cell is not DAergic, its presence does not reliably predict TH co-labeling [16,17]. A novelty of this study is in the use of a transgenic line of animals that expresses a reporter protein (eGFP) under the TH promoter, allowing the exact identification of each studied neuron as DAergic.

In this work we first report a kinetic characterization of the h-current in SNc neurons as close as possible to the physiological conditions (temperature, perforated patch), showing that this current at rest is larger than the one usually obtained. Then we describe how the resting membrane potential of the dopaminergic ‘principal’ cells are affected by this current. Finally, we show that neurotransmitters physiologically released onto SNc neurons can
modulate the h-current, thereby affecting the overall excitability profile of these cells.

**Results**

The results of this study are based on observations made in 357 voltage-clamped SNc neurons, all showing inward rectification at voltages negative to the resting membrane potential.

In order to isolate the h-current, except where otherwise stated, all the other main ionic currents were blocked; in particular, the sodium current with 0.6 μm tetrodotoxin (TTX), the delayed rectifier potassium current with 20 mM tetraethylammonium (TEA), the A-type potassium current with 3 mM 4-aminoypyridine (4AP), the calcium and calcium-dependent currents with 100 μM cadmium and the K_{IR} current with 0.5 mM barium.

**Biophysical Properties**

In this first section, all the recordings were carried out in slices at 37°C in whole-cell configuration. The input resistance was 413.64±30.17 MΩ (n = 188; 5–95 percentile range 136–651 MΩ), and the access resistance was 12.9±0.3 MΩ (n = 199; 5–95 percentile range 7.9–18.0 MΩ). In normal recording medium containing 2.5 mM-K+ the resting membrane potential of SNc neurons was -60.54 ± 0.92 mV (n = 91; 5–95 percentile range -50.2–71.0 mV), calculated as the potential corresponding to the zero injected current in voltage-clamp conditions. About 95% of the neurons fired spontaneously with a frequency of 4.9±0.67 Hz (n = 57, frequency measured in cell-attached mode).

All of the examined SNc neurons showed a time-dependent sag in response to the injection of outward currents (Figure 1A, arrow), becoming evident at potentials more negative than -60/70 mV; the termination of the current pulses was usually followed by a depolarizing overshoot; the depolarizing sag became progressively bigger as the membrane was stepped to more negative potential.

**Activation.** Under voltage clamp, hyperpolarizing commands from a holding potential of -40 mV evoked slow inward relaxations over the same membrane potential range as the ones producing sags and depolarizing overshoots in hyperpolarizing electrotonic potentials (Figure 1B). The h-current activated slowly and increased the magnitude and rate of activation as the cells were progressively hyperpolarized, with no sign of inactivation. Two current components were measured during the hyperpolarizing voltage steps: (i) an instantaneous current (I_{inst}), obtained at the beginning of the step; (ii) a steady-state current (I_{ss}), obtained at the end of the step. The instantaneous current was almost linear along the explored voltage, while the steady-state current increased its magnitude as the membrane potential was made more negative; the h-current amplitude, measured as I_{ss}-I_{inst} (see methods) is plotted against voltage in Figure 1C (orange symbols).

The steady-state activation curve (Figure 1C, yellow symbols) was obtained by interpolating the relative amplitudes of the tail currents with the Boltzmann function (Eqn. 2, explanation in Methods), finding - for 2 s hyperpolarizing step - values for half-activation (V_{50}) of -84.17±1.31 mV (n = 13) and for k of 7.74±0.40 mV (n = 13).

The point of half activation of the h-current critically depends on the hyperpolarizing pulse length [18]; conditioning pulses of short duration do not allow the gating process to reach the steady-state condition, therefore the probability of opening can be seriously underestimated, leading to evaluations of V_{50} more negative of their actual value. These measurement errors are more pronounced for slow HCN channels than for the fast ones, and are highly dependent on temperature [19,20]. Therefore, we have analyzed the dependence of the midpoint of the duration of the conditioning command. In nine cells studied with the double pulse protocol described above, the first command had durations of 1, 2, 4 and 8 s; we also tried the next point in the log scale, 16 s, but the membrane did not withstand the prolonged hyperpolarizations at the more negative potentials. Increasing the duration of the conditioning pulse induces a significant shift of the steady-state activation curves in the depolarizing direction: the values of V_{50} is changed from -87.52±2.0 mV for 1 s stimuli to -78.15±1.22 mV for 8 s (Figure 1D); we did not observe any change in the corresponding slopes. The V_{50} values as a function of the duration of the first step can be described by the exponential function

\[ V_{50(t)} = A + T_{50} * e^{-t/\tau} \]

where V_{50(t)} is the value assumed by the midpoint of the steady-state activation curve for conditioning potentials lasting t seconds; T_{50} is the range of variation of V_{50} as a function of the conditioning period t, and has a value of 13.90±1.58 mV; τ is the time constant of the process, 3.21±1.42 s. Finally, A is the asymptote (i.e. the value to which V_{50(t)} tends for t → ∞; dashed line in Figure 1D). The significant value obtained from this analysis is equal to -77.05 mV ±1.54; this means that, once the channel reaches its steady-state conditions, the point of half-activation is about 10 mV more positive than what is generally believed.

**De-activation.** The de-activation time constant was measured using the envelope test [21] shown in Figure 2: from a holding potential of -40 mV, two hyperpolarizing pulses to -130 mV lasting 4 s were imposed, separated by a repolarization to -40 mV of variable length (Figure 2A). In Fig. 2B, the I_{h} de-activation at -40 mV and the envelope of re-activation records at -130 mV shown in panel A are displayed together, in order to evidence the likeness of their exponential time course. The values of the amplitudes of the tail currents recorded upon re-activation at -130 mV were normalized, plotted as a function of depolarizing step duration (Figure 2C), and the de-activation time constant was calculated by interpolating the experimental points with the exponential function.

\[ I(t) = 1 + \frac{t}{t\tau} \]

where I(t) is the normalized current amplitude at time t, t the range of change of the normalized current, τ the time constant of de-activation at the potentials indicated. In a group of 5 cells, τ was 0.45±0.07 s (Fig. 2C).

**Reversal potential.** The h-current is carried by cation channels permeable to Na⁺ and K⁺ ions [22–24]; in fact, increasing the extracellular concentration of potassium from 2.5 to 32.5 mM produced a reversible increase in the amplitude of I_{h}.

The mean amplitude in I_{h} during exposure to 32.5 mM K⁺ was 47.6% ± 50 of control (n = 5, Figure 3A).

As shown in Figure 3A (arrow), we failed to observe any increase in the instantaneous current in high K⁺ [9]. The explanation for this discrepancy is complicated by the fact that the nature of I_{inst} is not well defined yet [25], contrary to the slower component, which is certainly sustained by cations passing through the well-characterized pore of HCN channels. In addition, I_{inst} usually has a small amplitude, and is not observed in any measurement of I_{h}. Speculations on the nature of I_{inst} range from models where this current represents a leak conductance or an experimental artifact, to models in which I_{inst} is caused by a second pore within the same...
HCN or a second channel population associated with HCN channels [26–28]. Midbrain DAergic neurons also have another hyperpolarization-activated current, KIR type, that could contribute to the I_{h} amplitude, and which is enhanced by an increase of the external K^+ concentration. It cannot be excluded, therefore, that the difference in the results might be consequence of different degrees of blockage of the KIR current, in addition to possible differences in the animal species used (mouse and rat).

The classical procedure to calculate the reversal potential of a voltage sensitive conductance is from the tail currents reversal, but in SNc neurons this method was rather problematic due to the activation of several outward rectifiers in the membrane potential range over which reversal was expected.

For a more precise calculation of the I_{h} reversal potential (E_{h}) we used the method of the instantaneous (chord) and ‘steady-state’ current-voltage relationships [29]. At the membrane potential of −95 mV g_{h} is strongly activated (Figure 3C) and does not show time-dependent inactivation; the reversal potential of I_{h} is then obtained from the intersection of the instantaneous (chord) current-voltage relationships recorded at holding potentials of −60 mV and −95 mV (Figure 3D), i.e. in the absence and presence of I_{h}.

In 8 neurons the mean value obtained for the reversal potential at 37°C was −44.03±3.10 mV (range −29.4 to −56.9 mV).

**Effect of temperature.** It has been shown in various types of preparation that the kinetics of I_{h} is particularly sensitive to thermic conditions [20,30,31]. The temperature at which electrophysiological recordings are made, affecting both the amplitude and the kinetics of Ih (Figure 4A), is one of the limiting factors in comparing the results; therefore, in this study most of the reported recordings were realized in precisely controlled temperature conditions.

Figure 4B shows the effect of a 10°C temperature increment on the Ih amplitude at different potentials. I/V graphs show the mean current amplitudes recorded at 27°C ( ) and 37°C ( ) as a function of membrane potential. At −130 mV, a 10°C increase causes a rise in amplitude from −194.2±18.5 pA at 27°C (n = 19) to −569.1±48.9 pA at 37°C (n = 23). The average value of Q_{10} for I_{h} amplitude between −80 and −130 mV is 3.76±0.53. The resulting maximal conductance g_{h} at 27 and 37°C is 2.26 and 6.62 nS, respectively.

We next explored whether the increase of I_{h} at −130 mV could be explained by a shift in the voltage dependency. As seen from the graph (Figure 4C), the transition from 27°C ( ) to 37°C ( ) causes a shift of the steady-state activation curve by about +10 mV: the V_{50}, calculated fitting the Boltzmann equation to the experimental points (4 s conditioning pulses), is −94.91±1.72 mV at 27°C (n = 13) and −84.23±1.28 mV at 37°C (n = 18),

![Figure 1. Basic properties of the h-current at 37°C. A: Response of a SNc DA neuron under current-clamp condition to the injection of a 300 pA hyperpolarizing current pulse. Note the appearance of the archetypal sag (arrow) due to the activation of I_{h}; resting potential −55 mV, bath solution for this recording was standard ACSF. B: Family of responses of a SN DA neuron under voltage-clamp conditions to the application of the double-pulse protocol shown in the inset above; explanation in the text. C: Current-voltage relationship of the h-current (□, right y-axis) and voltage-dependence of the activation curve (○, left y-axis) obtained from tail analysis of double-pulse experiments as shown in panel B; mean value ± S.E. (n = 20). D: Dependence of the midpoint (V_{50}) on the duration of the hyperpolarizing pulse; the dashed line highlights the asymptotical behavior of the midpoint for a conditioning pulse of infinite duration - see text for explanation. doi:10.1371/journal.pone.0052329.g001](figure1.png)
(P<0.0001, two-tailed Student t test for unpaired data). No significant changes were observed in the steepness of the relationship: the slope is $8.02 \pm 0.37$ mV at 27°C and $7.74 \pm 0.37$ mV at 37°C.

The temperature does not affect only the total conductance of the h-current (Figure 4A) but -and to a much higher degree- also its activation kinetics, which is modified under two aspects. First, the tracings at 27°C can be satisfactorily fitted by a single exponential (Figure 4E, E'), but two exponentials are always needed for an adequate fit at 37°C (Figure 4F, F'). Second, the current development rate is strongly affected by the temperature. Since at 27°C there is only an exponential, and at 37°C two, a comparison of the time courses was possible only comparing the 10–90% rise time. Since not always the steady state was reached, due to the instability of the membrane at the more negative potentials, we used the following equations, obtained by solving equation 3 (see methods) for $y = 10$ and $y = 100$ after normalization of the amplitude to 100:

- for a single exponential $t_{90} = \tau \ln(100/10)$ and $t_{10} = \tau \ln(100/90)$, where $t_{90}$ and $t_{10}$ are the times at which the current is developed for the corresponding percentage, and $\tau$ is the time constant;
- for a double exponential the solution was less straightforward: first the amplitudes of the two exponentials ($A_1$ and $A_2$) were normalized so that their sum was 100; then, Eqn. 1 was solved numerically for $f$ [32,33] setting $f(0) = 90$ and $f(10) = 10$ (the Matlab code used can be found in the Supplementary material as Code S1), obtaining $t_{10}$ and $t_{90}$, respectively.

The de-activation time constant is also affected by temperature: the double-pulse protocol described above was applied in five cells at 27°C. The normalized amplitudes of the currents to the second pulse as a function of the delay between the two pulses can be described by the exponential function indicated in equation 2, with a time constant of 1.46±0.1 s, which would give a Q10 of 3.2 when compared with the value at 37°C indicated in a section above.

Basic Pharmacology

The h-current is sensitive to low concentrations of Cs⁺ (1–2 mM) [34] and to a certain number of organic compounds blocking selectively the h-channels, like ZD7288 [35] and S-16257 (ivabradine) [36,37]. Cs⁺ 1 mM effectively blocks the h-current (Figure 5A). However, as already observed in calf Purkinje cells [38], the action of Cs⁺ is clearly voltage-dependent: in the negative region of the I-V curve Cs⁺ induces a channel blockade, whereas at more positive potentials it is ineffective, and sometimes it can...
even produce the opposite effect, i.e. a current increase. More selective, and completely voltage-independent blockages, can be obtained with ivabradine 10 μM (Figure 5B) and ZD7288 (30 μM, Figure 5C).

Role of Ih in Autorhythmicity

One of the hallmarks of SN neurons is the autorhythmicity: these neurons fire spontaneously action potentials characterized by an unusually long duration (>2.5 ms), a rather depolarized threshold (> −40 mV) and a marked afterhyperpolarization [1,39]. The role of the h-current in spontaneous activity has been thoroughly analyzed by several authors, and the conclusion has been that it is neither a significant factor underlying the spontaneous pacemaker activity nor does it contribute substantially to the setting of the normal resting potential level of the membrane [9,12–15].

Our data only partially confirm this viewpoint: recording at 37°C and in perforated patches, the block of the h-current by focal application of ivabradine 10 μM does stop the spontaneous activity; the effect is rapid and reversible, and is paralleled by an important hyperpolarization (11.83±2.07 mV, n = 7; Figure 6A). The blockage of spontaneous activity following a membrane hyperpolarization of 10 mV is not surprising, as the cell firing is based on a delicate interplay of conductances [8,14,40–42] that can be easily disrupted by the injection of outward currents [43], and personal observation). We then tested whether this blockade represented the evidence for an essential role played by the h-current in the pacemaking mechanism, or if it was only the consequence of the hyperpolarization following the suppression of the h-current. In the presence of ivabradine, the injection of a depolarizing current sufficient to restore the membrane potential to the value antecedent the Ih block (arrowhead in Figure 6), resumes immediately the spontaneous activity (Figure 6A, right).

This proves the absence of any direct role of the h-current in autorhythmicity, but also demonstrates that this current has a relevant role in determining the membrane potential.

Since the influence of the h-current on membrane potential is somewhat controversial, and since ivabradine is a relatively new drug, for which side effects have been described on currents other than Ih [44,45], we repeated the same experiment using the more classical blocker ZD7288 (30 μM). The results were substantially similar to those obtained with ivabradine (Fig. 6B), although the hyperpolarizing effect was less pronounced (6.87±0.78 mV, n = 15) and in 7 out of 15 cases we were unable to obtain a substantial recovery after 20’ washout. Interestingly, the hyperpolarizing effect of the h-current blockade was correlated with the resting membrane potential (Fig. 6C; p value <0.02, ANOVA), as expected for a conductance whose effect is increasingly influential at more negative potentials.

Modulation of h-current

Ih modulation by intracellular cAMP. The h current is dually regulated by hyperpolarization and by cyclic AMP, directly binding to a sequence (cyclic nucleotide binding domain, CNBD) located in the C-terminal segment [46,47]. We have therefore analyzed the modulatory effect of cAMP on the h-current using a recording configuration (perforated patch with amphotericin B) minimizing the perturbations of the intracellular medium.

The first experiments were conducted in current clamp conditions to determine the effects of increased intracellular cAMP on the resting membrane potential. The addition to the extracellular solution of 10 μM forskolin, a classical activator of adenylyl cyclase [48], caused a depolarization of 3.5±1.3 mV (n = 5) after 4 min of application. Then the effect declined, vanishing completely in the next 5 min with a return to the resting...
membrane potential, even maintaining the forskolin supply (Figure 7A, yellow dots). However, when the bathing solution was further enriched with 0.1 mM IBMX, a phosphodiesterase inhibitor [49], the depolarization induced by forskolin was more prominent (7.9 ± 1.8 mV, n = 5), and persisted as long as the application of the drug was maintained (Figure 7A, orange dots).

Under voltage-clamp conditions, the bath application of 10 μM forskolin and 0.1 mM IBMX, induce a significant increase of Ih amplitude (Figure 7B): at −130 mV the current amplitude is −178.5 ± 23.5 pA in control conditions (n = 8), and −227.0 ± 34.2 pA (n = 8) in the presence of increased levels of cAMP; at any tested potentials the increase in current amplitude was statistically significant (P<0.005 level, t-test for paired data).

The effect of forskolin on the Ih-current is twofold. First, it promotes a depolarizing shifts of the steady-state activation curve V50 in the depolarizing direction of 6.33 ± 0.78 mV (n = 8; a variation significant at 0.0005 level; Figure 7C). In addition, following the increase of intracellular levels of cAMP, the Ih activation time course becomes significantly faster: from 1763 ± 340 ms in control to 1435 ± 298 ms for the 10–90% rise time (n = 8; p<0.025) at −100 mV (Figure 7D).

Figure 4. Effect of temperature on h-current amplitude and kinetics. A: Family of current tracings recorded in a single cell in response to hyperpolarizing pulses from −40 to −130 mV, repeated at the temperatures indicated. B: Comparison of the I/V curves recorded at 27 (yellow dots) and 37 °C (orange dots); n = 20; the difference, tested with two-way ANOVA and post-hoc Bonferroni test, is significant at 0.001 level for the potentials more negative than −70 mV. C: Shift of the steady-state activation curves for a change from 27 °C (yellow) to 37 °C (orange), average values from 13 cells ± S.E for the V50. D: Effect of the temperature on the h-current 10–90% risetime at 27 °C (yellow) and 37 °C (orange); see explanation in the text. E, E’: Sample fit with a single exponential of an h-current tracing obtained at 27 °C in response to a voltage step to −130 mV; below the analysis of the residuals (experimental data minus the corresponding values of the fitting curve). F, F’: Sample fit with both single (green) and double (orange) exponential of a h-current tracing obtained at 37 °C in response to a voltage step to −130 mV; below the analysis of the residuals in the same grey scale code. All the recordings shown in this figure were obtained in perforated patch conditions.

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In midbrain DA neurons, serotonin has been reported to inhibit Ih in the VTA dopaminergic neurons in concentration-dependent manner [60], and to enhance the current in the substantia nigra pars compacta [61]. Therefore, we examined whether the 5-HT (100 μM) had any effect on the h-current performing the experiments in perforated patch-clamp configuration at 37°C. Using a single-pulse protocol from −40 mV to −130 mV, following application of serotonin and in the presence of Ba2+ to avoid collateral effects on the KIR current (see discussion), we observed a reduction of the h-current amplitude in 5 out of 8 cells (Figure 8B), whereas in the remaining cells we detected no significant changes (3 cells). In the first group the reduction in Ih amplitude was about 20%, and was significant at the 0.05 level (t-test, single queued). The progression of the effect in the time domain is shown in Figure 8D - the time constant measured was of 67 s.

A diffuse network of noradrenaline (NA)-containing nerve endings in the neuropil of SNc has been demonstrated. The noradrenergic input is from the locus ceruleus [62–64] and -to a lesser extent- from other NA-containing neurons in the brainstem [65,66]. NA modulates the h-current in several type of neurons, amongst which thalamic relay neurons [67], neurons of the medial nucleus of the trapezoid body (MNTB) [68], dorsal root ganglia neurons [69] and VTA neurons [70,71]. In the thalamus, NA, acting on β-adrenergic receptors, increases the intracellular levels of cAMP and shifts the Ih steady-state activation curve to more depolarized potentials, as it does in MNTB neurons. On the contrary, activation of α2 adrenergic receptors in the DRG and VTA neurons causes a significant reduction in amplitude of Ih.

We therefore tested whether the NA (100 μM) could determine any change in the h-current of DA neurons of SNc. We performed experiments in perforated patch configuration at 37°C, applying a single hyperpolarizing pulse from −40 mV to −130 mV. Upon application of NA, in 8 out of 10 cells we observed an increase in the amplitude of the h-current by about 11.8% (Figure 8C), whereas no change was detected in the remainder of SNc cells. The increase in amplitude was statistically significant, with a P-value <0.015 using single-tailed t-test for paired data.

**Discussion**

Two classes of hyperpolarization-activated inwardly rectifying currents have been reported in SNc neurons. One type has fast kinetics, is permeable primarily to K+, is blocked by extracellular Ba2+ and Cs+, and has a voltage-dependence that is itself dependent on extracellular K+ ([K+]o) concentration. Ih (or Ih) in cardiac tissue, the second type of inward rectifier, is a mixed
Ih has a relatively slow activation kinetics, is insensitive to Ba$^{2+}$, and does not show a voltage sensitivity dependent on \([K^{+}]_o\) [25]. The sensitivity to drugs very selective for Ih like ivabradine or ZD7288, the Ba$^{2+}$ insensitivity, the slow kinetics of activation and the reversal potential suggest that the current described here belongs to the latter class.

From the methodological point of view, a first novelty of this study is in the use of a transgenic line of animals that expresses a reporter protein (eGFP) under the TH promoter, allowing the precise recognition of DA neurons in \textit{in vitro} slices. This is of some importance as, in these preparations, neurons are often identified as DAergic according to the expression of Ih, a current which is not-or much less- expressed in putative non-DA neurons [1,9,73,74]. Notably, while the absence of Ih in a DA neuron is a reliable predictor that the cell is not DA-containing, the presence of Ih does not reliably predict TH co-labeling [16,17]. In other words, since a significant number of neurons expressing the h-current is not DAergic, this means that one of the principal markers assumed to identify midbrain DAergic neurons is not associated exclusively, or even significantly, with confirmed DAergic neurons. The identification of the neurons by the expression of eGFP under the TH promoter, marks a first important difference of this study with respect to other functional studies, where the expression of TH (when done) was carried out only in few sample cells.

A second methodological hallmark of this work is the choice of working systematically at 37°C and in perforated patch configu-
ration, especially when the modulation of the h-current was studied. These working conditions are more respectful of the intracellular milieu and, in our view, lead to significant differences in the observed results.

In this preparation, Ivabradine was used for the first time as a blocker of Ih. This drug, originally developed as a bradycardic agent [44], proved to be an excellent blocker of all types of HCN subunits [37,75]. The reason of the preference accorded to this drug resides in the rapidity and complete reversibility of its action, whereas the more classical Ih blocker ZD7288 is slower and often irreversible [76,77].

Biophysical Properties

Steady-state activation. At 37°C, and a 4 s hyperpolarizing pulses, we found a midpoint of activation at −82.73±1.17 mV with a slope of 7.29±0.28 mV (n = 12); however, as first shown by ref. [18], the point of half-activation depends critically on the used voltage protocol. In particular, short voltage steps do not allow channel activation to come to completion and then the steady-state activation curve derived from tail current amplitudes is seriously underestimated. Using different pulse duration, and extrapolating to the asymptote the trend of the midpoint shift in the depolarizing direction as a function of the hyperpolarizing pulse duration (Figure 1D), we calculated a V50 of −77.05 mV, i.e. from 10 to 28 mV more depolarized than the values reported in SNc neurons at 37°C [9,78].

The functional implication of these values becomes obvious if one considers that assuming a slope of 7.43 (the slope is 7.41, 7.36, 7.29 and 7.65 at 1, 2, 4 and 8 s, respectively), at −65 mV about 16.5% of the h-channels are open, which corresponds to a conductance of 1.023 nS. Since we found an average input resistance of these cells of 414 MΩ, at −65 mV the h-current gives a contribution of +8.9 mV to the resting membrane potential, a value matching very well the hyperpolarizations found upon blocking the h-current with ivabradine in the present experiments (−21.18±2.07 mV). Such an hyperpolarization would be impossible with the kinetic coordinates reported in most of the previous papers and the value reported in literature. Since the hyperpolarizing effect of the Ih blockage is controversial, we repeated the experiments using the more classical blocker ZD7288 (30 μM), obtaining results substantially similar to those obtained with ivabradine, although the hyperpolarizing effect was less pronounced (−6.87±0.78 mV, n = 15).

There could be several explanations to justify the discrepancy of our data with others reported in literature, which do not show any effect of the h-current on the resting membrane potential [9,12–15,79].
First, it cannot be excluded that ivabradine could also act on conductances different from the h-current. In fact, high inhibitory concentrations of ivabradine have been reported to inhibit both L- and T-type calcium currents [44,45]; however, at the concentrations used in the present experiments (10 μM), to the best of our knowledge, no side effects have ever been described, although these cannot be certainly excluded, considering the still scarce knowledge of this drug in nerve cells. However, we see a hyperpolarizing effect also of ZD7288; furthermore, notice that our result (a hyperpolarization of 6.9 mV) is almost coincident with that reported in a recent paper (7 mV; [80]).

Second, as shown by previous studies, Ih amplitude correlates with calbindin (CB) expression in nigral dopamine cells, with larger currents in CB-negative cells [15,81], and dissimilar proportions of CB-positive and CB-negative cells could have been analyzed in different studies - in this regard, direct analysis by molecular methods would help to characterize calbindin expression when recording from SN neurons.

**Temperature dependence.** For what concerns current amplitude, the Q10 values reported by various authors lie in the range between 1.35 [20,82] and 6 [83], with an average around 3. We found a value of 3.76 ± 0.53, which is well aligned with most of the reports. This increase is generally ascribed to depolarizing shifts in the steady-state activation curve for the h-current which, for a 10°C temperature increase, range from 2–3 mV [82] to 12–15 mV [31,84] without change in the slope. In SNc neurons, rising the temperature from 27 to 37°C we observed a depolarizing shift in the midpoint of the steady-state activation curve (V50) of 10 mV in the depolarizing direction, without any change in the slope.

The main difference in temperature sensitivity with respect to the current literature was observed in the activation rise time, for which we found a Q10 of 6.4, while values found by other authors are around 3 [30,34,82] - a diversity that can be explained by differences in the preparation and consequently in the subunit composition of the channels.

The high temperature-dependence of the Ih in SNc neurons underscores the importance of controlling this often neglected parameter in the electrophysiological recordings of this current. This dependence can be at the basis of many of the discrepancies found in literature.

The high sensitivity of the h-current in substantia nigra neurons might at least in part explain the previously observed warming-induced increase in firing frequency, decrease in input resistance, and an inward current reversing its polarity between −5 and −17 mV, which is dependent on extracellular Na+ [85]. In this context, it might also be useful to recall that changes of up to several degrees centigrade in the brain temperature are observed not only during fever, but also during different behavioral states [86]. Furthermore, neurons in different brain regions, including the substantia nigra, were reported to show high temperature sensitivity [87], a behavior whose underlying mechanism is unknown. Our data indicate that the h-current might have a role in the process.

**Pharmacological Properties**

The fundamental involvement of Ih in the control of resting membrane potential makes the h-channel a privileged target for neurotransmitter systems aiming at the regulation of SNc neurons activity. In general, the excitability profile of a cell expressing an h-current can be controlled either by moving the membrane potential in and out of the range of Ih activation, or by moving the Ih activation within the range of membrane potentials of physiological interest by directly modulating the h-channel itself.
In fact, a number of intracellular signaling molecules can affect $I_{h}$ in SNc neurons, including -besides the cAMP- also phosphoinositides [88,89] and kinases [90]. In particular, the possibility to control the voltage dependence of the h-channels in SNc neurons appears to enable transmitters of ascending activation systems, such as serotonin, noradrenaline, and dopamine, to specifically reduce or increase the membrane impedance, and more in general the excitability profile of the cell. Some of these influences, like modulation by GABAergic input arising from other nuclei in the basal ganglia, have been carefully studied (for a review see [91]), other, as those mediated by cyclic nucleotide pathways, have been less extensively investigated. In this work we have attempted a preliminary exploration of the possible modulation of the h-current by some neurotransmitter known to act on SNc neurons.

One known possible source of error is that the modulation (inhibition) of the h-current occurs secondary to GIRK channel activation [9,78,92], since between GIRK and $I_{h}$ channels there is a close link [93,94]. We can exclude this possibility since, in experiments where the modulation of the h-current was studied, we were always in the presence of Ba$^{2+}$ at concentrations that can entirely block the inward rectifier K$^{+}$-currents, GIRK included.

**$I_{h}$ modulation by cAMP.** The $h$-channels are directly activated by cyclic nucleotides [46], but when tested in SNc neurons, the adenylate cyclase activator forskolin gave contradictory results on the amplitude of the h-current, going from a complete absence of effect [78] to a 25% increase, due to a +5.33 mV shift of the steady-state activation curve towards more positive potentials [95]. Our results are almost exactly superimposable with those of the latter authors, with a 27.5% increase in current amplitude and a +6.33 mV shift of V_{50}. It should be noted that initially we were unable to see any influence because we tested the forskolin effect 10 minutes after application, and without IBMX. In these conditions, the modification of the resting membrane potential is transient and vanishes after 7–8 min (Figure 7A), but in the presence of the phosphodiesterase inhibitor the effect is larger and stable in time, and this might explain some discrepancy in the literature about this point.

**D2 receptors.** Since its discovery by Aghajanian in 1977 [96,97], the presence of D2 receptors in midbrain DA neuron, is considered one of the hallmarks of these cells, and as such it has received much attention. DA D2-like receptors are present as auto-receptors on the DA neurons in SN and VTA and play an important role in the regulation of DA neuronal firing activity by means of auto-inhibition (for a review see [98]). These G protein-coupled receptors are activated by dendritically released DA [99,100] through a still controversial mechanism [101], and the hyperpolarization following their activation is classically described as mediated through an activation of potassium channels, GIRK type [43,78,102], and A-type [103,104]. In the present work, we show that, in addition, also the h-current is modulated by the activation of D2-like type DA receptors. This is not surprising, as D2 receptors are coupled to a G-protein $G_{q}$ which directly inhibits the formation of cAMP by inhibiting the enzyme adenylate cyclase [54]. Our observation confirms a relatively recent study, conducted in brain slices at 34°C showing that DA, released endogenously following a single action potential, hyperpolarizes neighboring DA neurons by inhibiting h-channels [105].

Thus, it is possible that DA autoreceptors are linked to different effector systems. Receptors that are coupled to h-channels may be preferentially located in the vicinity of DA release sites, whereas those coupled to G-protein-gated K$^{+}$ channels may be extra-synaptic.

**Serotonergic receptors.** The bulk of available neuroanatomical data clearly indicate that the midbrain DA- neurons receive a prominent innervation from 5-HT originating in the raphe nuclei of the brainstem [106–108]. The modulatory action of 5-HT on midbrain dopaminergic neurons is complex, and probably exerted through different pathways (for an overview see [109,110]) - here we have limited the analysis of the 5-HT to its effects on the h-current.

In general, 5-HT is reported to increase the h-current in the CNS [67,111,112]. In the midbrain DA neurons both enhancement [61] and inhibition [60] of the $I_{h}$ have been reported. Our data confirm the first action, although we cannot establish whether the effect is direct or indirect.

**Noradrenergic receptors.** Controversial data were reported concerning the effects of NA in SNc. Several studies indicate that electrical stimulation of the locus coeruleus evokes an initial excitatory response in SNc neurons, frequently followed by a period of inhibition of firing [64,113]. In *vivo* studies report an hyperpolarization in rat SNc neurons by NA [43], which slows the frequency of spontaneous action potentials [114]. In SNc neurons other authors have found that NA apparently induced an inhibition of $I_{h}$ but then they report that in the presence of 300 µM external barium or internal cesium, NA did not affect $I_{h}$, suggesting that the effect on $I_{h}$ is secondary to the activation of $K_{IR}$ channels [92]. Contrary to what was found by these authors, we can exclude an indirect effect mediated by $K_{IR}$ channels as we were systematically recording in barium 500 µM. We did not look for better specific pathway of the pathway involved in the NA stimulation of the h-current, which Cathala and Pupardin-Tritsch suggest being PKC, nor of the receptor involved, for which both α1 and α2 adrenergic types seem to be excluded [92]. Nevertheless, the observation of Cathala and Pupardin-Tritsch that NA inhibits the $K_{IR}$ current, in addition to enhance the $I_{h}$, is rather interesting, as there is an increasing evidence that also in other systems amine-activated pathways can modulate both $I_{h}$ and $K_{IR}$ acting in opposite direction. A synchronous and symmetrical control of the balance between $I_{h}$ and $K_{IR}$ was also described for example in rat spinal motoneurons, where 5-HT increases the cell excitability inhibiting an $K_{IR}$ and enhancing an $I_{h}$ [115]; in salamander motoneurons, where muscarinic modulation inhibits the $I_{h}$ and enhances the $K_{IR}$ [116]; in DAergic neurons of the olfactory bulb, where cAMP increases the $I_{h}$ and inhibits the $K_{IR}$ (personal unpublished observation).

In SNc neurons, where $I_{h}$ is enhanced and $K_{IR}$ is inhibited by NA, this synchronous and symmetrical modulation might underlay a mechanism that could shorten hyperpolarizing events. Since the activation and deactivation kinetics of $K_{IR}$ are much faster than that of $I_{h}$, the net result would be a sharper response of the neuron in response to hyperpolarizing events. The enhancement of $I_{h}$ by NA could then accelerate the recovery from hyperpolarizations, ultimately shortening their duration and increasing firing frequency, a combined action that might be shared, not necessarily with the same sign, also in other systems.

**Functional Implications and Conclusions**

A substantial amount of research has focused on determining the factors that alter the activity of substantia nigra DAergic neurons. Much of this research indicated that several mechanisms that regulate dopamine neuron activity have the capability to maintain the baseline activity of dopamine cells at a fairly constant rate [117]. However, DAergic cell activity can be finely tuned also by afferent inputs that involve an assortment of ionotropic and metabotropic receptors acting directly on the neurons themselves. In addition, the neurons can modulate the input received at local level through dendritic dopamine release that affect their own responsiveness to afferent input by controlling dendritic excitabil-
ity through D2 autoreceptors. We show that the $I_{\text{h}}$, with its role in the control of membrane potential, seems to be an important target of the afferent inputs to SNC neurones. We propose that this current may be one of the main actors responsible for the rich signaling repertoire displayed by these cells which, through their effects on forebrain dopamine levels, influences much of the functioning of the basal ganglia as a whole.

**Materials and Methods**

**Animals and Surgical Procedures**

Experimental procedures were carried out to minimize animal suffering and the number of mice used. The procedures employed were in accordance with the Directive 86/609/EEC on the protection of animals used for experimental and other scientific purposes, and were approved by the Campus Veterinarian of the Ferrara University. A total of 192 mice have been used, most of them in the 14-20 postnatal day range, and 20 over 3 months old.

All experiments were performed using the transgenic mice TH-GFP/21–31 line carrying the eGFP gene under control of the TH promoter [118,119]. Transgenic mice were identified either by PCR on the genomic DNA extracted from tail biopsies, or -at postnatal day 3 or 4- looking at the fluorescence of the olfactory bulbs transilluminated with a UV source (FBL/Basic-B & N-01; BLS, Hungary; FHS/F-01) and observed with an emission filter (FHS/EF-2G2; BLS, Budapest, Hungary). Transgenic lines were maintained as heterozygous by breeding with C57BL/6J inbred mice.

**Preparation of Midbrain Slices**

Mice were anaesthetized (intraperitoneal injection of 60 mg/kg of sodium pentobarbital) and decapitated. The brain was removed from the skull in less than 1.5 min and put into ice-cold (2–4°C) dissection solution of the following composition (in mM): 3.0 KCl, 1.25 NaH2PO4, 2 MgCl2, 1.6 CaCl2, 10.0 glucose, 21.0 NaHCO3, 215 sucrose; saturated with 95% O2 and 5% CO2.

A brain section containing the substantia nigra pars compacta was obtained as follows. Two coronal cuts were performed, the caudal to remove the cerebellum and the rostral half of the cerebral hemispheres. The resulting block was glued on the rostral surface with n-butyl cyanoacrylate adhesive (3 MTM Vetbond), Segrate, Italy) and then submerged by ice-cold dissection solution. The coronal slices (thickness, 150 μm) were cut starting from the caudal surface. Slices containing the substantia nigra pars compacta were identified by illuminating the specimen with a desk UV light source (BLS), stored in an incubation chamber containing artificial cerebrospinal fluid (ACSF) continuously bubbled with carboxygen (95% O2, 5% CO2; the osmolarity was adjusted at 305 mOsm with glucose). All EC solutions were continuously bubbled with 95% O2 and 5% CO2; the osmolality was adjusted at 305 mOsm with glucose. All EC solutions were continuously bubbled with 95% O2 and 5% CO2; the osmolality was adjusted at 305 mOsm with glucose. All EC solutions were continuously bubbled with 95% O2 and 5% CO2; the osmolality was adjusted at 305 mOsm with glucose.

**Solutions**

The solutions used had the following composition (mM):
- **standard ACSF extracellular (EC) saline:** 125 NaCl, 2.5 KCl, 26 NaHCO3, 1.25 NaH2PO4, 2 CaCl2, 1 MgCl2, and 15 glucose.
- **high K EC solution:** 115 NaCl, 12 KCl, 26 NaHCO3, 1.25 NaH2PO4, 2 CaCl2, 1 MgCl2, and 15 glucose.
- **high sucrose EC dissection solution:** 3.0 KCl, 1.25 NaH2PO4, 2 MgCl2, 1.6 CaCl2, 10.0 glucose, 21.0 NaHCO3, 215 sucrose.

All EC solutions were continuously bubbled with 95% O2 and 5% CO2; the osmolality was adjusted at 305 mOsm with glucose.

**Current and Voltage Recordings**

The temperature of the 1-ml recording chamber was controlled using a couple of 39.7 W Pelhier devices (RS Components, Milan, Italy) and measured with a high-precision, low mass thermocouple (Messergeratebau, Katlenburg-Lindau, Germany); the seal resistance was always greater than 3 GΩ. A 70–80% compensation of the series resistance and correction for junction potential was routinely used.

For perforated patches, amphotericin B was included in the recording electrode filling solution as perforating agent (200 μg/ml plus 300 μg pluronic F-127). In order to make sure of the integrity of the perforated patch, EGTA was omitted from this solution and the concentration of CaCl2 was raised to 3 mM. Data were collected after the series resistance fell to <50 MΩ.

In all IC solutions the osmolality was adjusted to 295 mOsm with glucose, and the pH to 7.2 with KOH.

IVAbradine was a generous gift from Servier (Suresnes, France).

**Analysis of Current Records**

Offline analysis was performed using version 10.2 of pClamp (Molecular Devices) and version 8 of Origin (OriginLab Corporation, Northampton, MA).

The $I_{\text{h}}$ amplitude was measured as the difference between the steady-state current at the end of test voltage pulses ($I_{\text{h}}^\text{max}$) and the instantaneous current and the beginning ($I_{\text{h}}^\text{min}$); the latter were measured extrapolating the exponential fitting the h-current (single or double, see below) to the time of the onset of the hyperpolarizing pulse, as indicated by the arrow in Figure 4E.

Rates of $I_{\text{h}}$ activation were determined using the following function (Clampfit 10.2, Molecular Devices):

$$f(t) = \sum A_i \cdot e^{-t/\tau_i} + C$$

where $i = 1$ or 2 (a single or double exponential fit), $A$ is the amplitude of the fitting component(s), $\tau$ is the time constant, and $C$ is the shift of the fitted trace from zero.

The activation curve of $I_{\text{h}}$ was constructed using a two-step protocol [120]: the $I_{\text{h}}$ was first activated to a variable degree by a conditioning step, and then fully activated by a second pulse to −130 mV (Figure 1B). The resulting tail current amplitudes were then normalized and fitted by the equation:

$$\frac{I_{\text{h}}^\text{tail}}{I_{\text{h}}^\text{tail max}} = \{1 + \exp[(V_m - V_0)/k]\}^{-1}$$

$V_m$ is the membrane voltage and $V_0$ is the voltage (assumed to be −50 mV) at which the current will reach a value of 1.5 o.d., 0.87 i.d., with filament; Hilgenberg, Malsfeld, Germany) were pulled with a Zeitz-DMZ puller (Martinsried, Germany) and had a resistance of 4–5 MΩ when filled with standard intracellular solution; the seal formation was realized with the help of an air pressure controller (MPI, Lorenz Messgeratebau, Katlenburg-Lindau, Germany); the seal resistance was always greater than 3 GΩ.
Statistical significance of the results was assessed with one-way or change of the rate analyzed.

The temperature coefficients of activation and deactivation time constant were defined as:

\[ Q_{10} = \left( \frac{\text{rate}(T_2)}{\text{rate}(T_1)} \right)^{\frac{10}{(T_2 - T_1)}} \]  

thus, for every 10°C of change in temperature there is a \( Q_{10} \) fold change of the rate analyzed.

Unless otherwise stated, data are presented as means ± s.e.m. Statistical significance of the results was assessed with one-way or two-way analysis of variance (ANOVA), or Student’s \( t \) test for paired samples, as indicated; the software used was Prism 5 (GraphPad software, Lajolla, CA). A \( P \) value <0.05 was considered significant.

**Supporting Information**

**Code S1** Matlab code for the solution of eqn. 3. (Word file).

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**Author Contributions**

Conceived and designed the experiments: OB CG. Performed the experiments: CG AP. Analyzed the data: OB CG. Contributed reagents/materials/analysis tools: OB. Wrote the paper: OB.
46. DiFrancesco D, Tortora P (1991) Direct activation of cardiac pacemaker
43. Lacey MG, Mercuri NB, North RA (1987) Dopamine acts on D2 receptors to
67. McCormick DA, Pape HC (1990) Noradrenergic and serotonergic modulation
66. Moore RY, Card JP (1984) Noradrenaline-containing neuron systems. In:
65. Lindvall O, Bjo¨rklund A (1983) Dopamine- and Norepinephrine-containing
64. Collingridge GL, James TA, MacLeod NK (1979) Neurochemical and
63. Jones BE, Moore RY (1977) Ascending projections of the locus coeruleus in the
61. Nedergaard S, Flatman JA, Engberg I (1991) Excitation of substantia nigra pars
60. Liu Z, Bunney EB, Appel SB, Brodie MS (2003) Serotonin reduces the
58. Parent M, Wallman MJ, Gagnon D, Parent A (2011) Serotonin innervation of
57. Mori S, Matsuura T, Takino T, Sano Y (1987) Light and electron microscopic
56. Imai H, Steindler DA, Kitai ST (1986) The organization of divergent axonal
54. Neves SR, Ram PT, Iyengar R (2002) G protein pathways. Science 296: 1636–
53. Levey AI, Hersch SM, Rye DB, Sunahara RK, Niznik HB et al. (1993) Localization of dopamine D2 receptor mRNA and D1 and D2 receptor binding in the brain and pituitary: an in situ hybridization-receptor autoradiographic analysis. J Neurosci 13: 2507–2600.
52. Mansour A, Meador-Woodruff JH, Bunzow JR, Civelli O, Akil H et al. (1990) Baclofen inhibition of the hyperpolarization-activated current, Ih, in substantia nigra zona compacta neurons may be secondary to potassium current activation. J Neurophysiol 76: 2262–2270.
51. Harris NC, Constanti A (1995) Mechanism of block by ZD 7288 of the hyperpolarization-activated inward rectifying current in guinea pig substantia nigra neurons in vitro. J Neurophysiol 74: 2356–2378.
50. Mirejei A, Weis A, Ramirez JM (2011) Calcium-activated non-selective cation currents are involved in generation of tonic and bursting activity in dopamine neurons of the substantia nigra pars compacts. The Journal of Physiology 589: 2497–2514.
49. Brown MTC, Henny P, Bolam JP, Magill PJ (2009) Activity of neurochemically heterogeneous dopaminergic neurons in the substantia nigra: spontaneous and driven changes in brain state. J Neurosci 29: 2915–2925.
48. Orío P, Madrid R, de la Peña E, Parra A, Messeguer V et al. (2009) Characteristics and physiological role of hyperpolarization activated currents in mouse cold thermoreceptors. The Journal of Physiology 587: 1961–1976.
47. Robinson RB, Siegelbaum SA (2003) Hyperpolarization-activated current: from molecules to physiological function. Annu Rev Physiol 65: 453–480.
46. Varogas G, Lucero MT (1999) Dopamine modulates inwardly rectifying hyperpolarization-activated current (Ih) in cultured rat olfactory receptor neurons. J Neurophysiol 81: 149–158.
45. Guatteo E, Chung KK, Bowsala TK, Bernardi G, Murer NB et al. (2005) Temperature sensitivity of dopaminergic neurons of the substantia nigra pars compacta: involvement of transient receptor potential channels. J Neurophysiol 94: 3069–3080.
44. Kiyatkin EA (2011) Brain temperature homeostasis: physiological fluctuations and pathological shifts. Front Neurosci 5: 1036–1036.
43. Stein J, Gisolfi CV, Mora F (1982) Temperature regulation and dopaminergic systems in the brain: does the substantia nigra play a role? Brain Res 234: 275–296.
42. Zilles K, Klocker N, Wenzel D, Weisner-Thoms, J, Fleischmann BK et al. (2006) Pamacem by HCN channels requires interaction with phosphoinositides. Nature 44: 1027–1036.
41. Fogel KJ, Lyashchenko AK, Turbendian HK, Tibbs GR (2007) HCN pacemaker channel activation is controlled by acidic lipids downstream of diacylglycerol kinase and phospholipase A2. J Neurosci 27: 2802–2814.
40. Zong X, Eckert C, Yuan H, Wahl-Schott C, Abicht H et al. (2005) A novel mechanism of modulation of hyperpolarization-activated cyclic nucleotide-gated channels by Src kinase. J Biol Chem 280: 34224–34232.
39. Lee CR, Tepper JM (2009) Basal ganglia control of substantia nigra dopaminergic neurons. J Neural Transm Suppl 71: 90–96.
38. Cathala L, Paupardin-Tritsch D (1999) Effect of catecholamines on the hyperpolarization-activated cationic Ih and the inwardly rectifying potassium E(Kr) currents in the rat substantia nigra pars compacta. Eur J Neurosci 11: 398–406.
37. Svoboda KR, Lupica CR (1998) Opioid inhibition of hippocampal interneurones via modulation of persistent and hyperpolarization-activated current (Ih) currents. J Neurosci 18: 7044–7049.
36. Takigawa T, Alzheimer C (1999) G protein-activated inwardly rectifying K+ (GIRK) currents in dentrites of rat neocortical pyramidal cells. J Physiol 517 (Pt 2): 383–390.
35. Cathala L, Paupardin-Tritsch D (1997) Neurotensin inhibition of the hyperpolarization-activated current (Ih) in the rat substantia nigra pars compacta implicates the protein kinase C pathway. J Physiol 503 (Pt 1): 87–97.
96. Aghajanian GK, Bunney BS (1977) Pharmacological characterization of dopamine "autoreceptors" by microiontophoretic single-cell recording studies. Adv Biochem Psychopharmacol 16: 433–438.

97. Aghajanian GK, Bunney BS (1977) Dopamine "autoreceptors": pharmacological characterization by microiontophoretic single cell recording studies. Naunyn Schmiedebergs Arch Pharmacol 297: 1–7.

98. Shi WX (2009) Electrophysiological characteristics of dopamine neurons: a 35-year update. J Neural Transm Suppl 103–119.

99. Cheramy A, Leviel V, Glowinski J (1981) Dendritic release of dopamine in the substantia nigra. Nature 289: 537–542.

100. Bernardini GL, Gu X, Viscardi E, German DC (1991) Amphetamine-induced and spontaneous release of dopamine from A9 and A10 cell dendrites: an in vitro electrophysiological study in the mouse. J Neural Transm Gen Sect 84: 183–193.

101. Kennedy MJ, Ehlers MD (2011) Mechanisms and function of dendritic exocytosis. Neuron 69: 856–875.

102. Kim KM, Nakajima Y, Nakajima S (1995) G protein-coupled inward rectifier modulated by dopamine agonists in cultured substantia nigra neurons. Neuroscience 69: 1145–1138.

103. Hahn J, Tse TE, Levitan ES (2006) D2 autoreceptors chronically enhance dopamine neuron pacemaker activity. J Neurosci 26: 5240–5247.

104. Vanderhaeghe HC, Miller JJ (1977) An anatomical and electrophysiological investigation of the serotonergic projection from the dorsal raphe nucleus to the substantia nigra in the rat. Neuroscience 2: 973–987.

105. Steinbusch HW (1981) Distribution of serotonin-immunoreactivity in the central nervous system of the rat-cell bodies and terminals. Neuroscience 6: 557–618.

106. Otijudgers JE, Werkman TR, McCreary AC, Kruse CG, Wadman WJ (2006) Modulation of midbrain dopamine neurotransmission by serotonin, a versatile interaction between neurotransmitters and significance for antipsychotic drug action. Curr Neuropharmacol 4: 59–68.

107. Di Giovanni G, Di Matteo V, Pierucci M, Eposito E (2008) Serotonin-dopamine interaction: electrophysiological evidence. In: Giuseppe Di Giovannii, WDMaEE, editors. Progress in Brain Research. Serotonin-Dopamine Interaction: Experimental Evidence and Therapeutic Relevance. Elsevier. 45–71.

108. Bobker DH, Williams JT (1989) Serotonin augments the cationic current Ih in central neurons. Neuron 2: 1535–1540.

109. Gasparini S, DeFrancesco D (1999) Action of serotonin on the hyperpolarization-activated cation current (Ih) in rat CA1 hippocampal neurons. Eur J Neurosci 11: 3093–3100.

110. Grenhoff J, North RA, Johnson SW (1993) Alpha 1-adrenergic effects on dopamine neurons recorded intracellularly in the rat midbrain slice. Eur J Neurosci 7: 1707–1713.

111. White FJ, Wang KY (1984) Pharmacological characterization of dopamine autoreceptors in the ventral rat midbrain: a microiontophoretic study. J Pharmacol Exp Ther 231: 275–280.

112. Kjaerulf O, Kiehn O (2001) 5-HT modulation of multiple inward rectifiers in motoneurons in intact preparations of the neonatal rat spinal cord. J Physiol 540: 540–553.

113. Chevalier S, Nagy F, Cabelguen JM (2006) Cholinergic control of excitability of spinal motoneurones in the salamander. J Physiol 570: 525–540.

114. Pucak ML, Grace AA (1994) Regulation of substantia nigra dopamine neurons. Crit Rev Neurobiol 9: 67–89.

115. Matsushita N, Okada H, Yasoshima Y, Takahashi K, Kischi K et al. (2002) Dynamics of tyrosine hydroxylase promoter activity during midbrain dopaminergic neuron development. J Neurochem 82: 293–304.

116. Savamoto K, Nakao N, Kakishita K, Ogawa Y, Toyama Y et al. (2001) Generation of dopaminergic neurons in the adult brain from mesencephalic precursor cells labeled with a nestin-GFP transgene. J Neurosci 21: 3993–3993.

117. Kamondi A, Reiner PB (1991) Hyperpolarization-activated inward current in histaminergic tuberomammillary neurons of the rat hypothalamus. J Neurophysiol 65: 1902–1911.