Acute Hypoxia Activates an ENaC-like Channel in Rat Pheochromocytoma (PC12) Cells

Yeon Ju Bae, Jae-Cheal Yoo, Nammi Park, Dawon Kang, Jaehee Han, Eunmi Hwang, Jae-Yong Park, and Seong-Geun Hong

Department of Physiology, Institute of Health Sciences and Medical Research Center for Neural Dysfunction, Gyeongsang National University School of Medicine, Jinju 660-751, Korea

INTRODUCTION

Hypoxia induces critical stress to cells under ischemic conditions, resulting in membrane depolarization, increased cellular Ca\(^{2+}\), alterations in pH, and finally cell death without recovery to normoxia [1-3]. Upon exposure to hypoxia, chemo-sensitive cells, such as those in the carotid body and adrenal medulla, rapidly elicit appropriate changes in both signal and cell function [4]. Oxygen-sensitive K\(^+\) (KO\(_2\)) channels have been identified in rat pheochromocytoma (PC12) cells from the adrenal medulla [5,6]. Inhibition of this current by hypoxia results in cellular depolarization as well as increased intracellular Ca\(^{2+}\) through voltage-operated Ca\(^{2+}\) channels [4-7]. The Kv1.2 gene was suggested to encode the \(\alpha\) subunit of slow-inactivating KO\(_2\) channels in PC12 cells [7]. Thus, PC12 cells have been used as a model system to study O\(_2\)-sensitive channels and their function in response to low O\(_2\) levels.

Although KO\(_2\) channels are considered to be the main modulator of depolarization in response to hypoxia, other types of ion channels may be involved. For example, hypoxia could potentially increase the background inward Na\(^+\) current. Therefore, the depolarization induced by hypoxia may not be solely due to the inhibition onset of K\(^+\) efflux but also the net influx of cations largely carried by Na\(^+\) [8]. Thus, although the initial cause of depolarization is reduction of K\(^+\) current, TTX-sensitive Na\(^+\) influx is critical for depolarization as it is unaffected by hypoxia [9,10].

Recently, amiloride-sensitive and voltage-independent Na\(^+\) channels such as ENaC, which currents are enhanced by the extracellular pH drop, have been suggested as the pro-

KEY WORDS: Acute hypoxia, Amiloride, Epithelial Na\(^+\) channel, PC12 Cells, Rats
...ton sensor involved in the transduction of ischemic signals in CNS neurons [11]. In addition to their pathophysiological role, ENaC-like Na⁺ channels are tightly regulated in non-epithelial cells in order to avoid cell death due to Na⁺ loading and a reduced electrochemical membrane gradient [12].

Depolarization through ENaC activated by hypoxia/ischemia remains unclear. A number of signaling pathways that modulate ENaC activity in response to hypoxia have been described, including the Ca²⁺-calmodulin pathway, AMPK pathway, ERK1/2 pathway, stress-activated protein kinase (also known as the p38 kinase) pathway, and phosphatidylinositol 3-kinase (PIK)-Akt pathway [13].

Among them, the PI3K/Akt pathway has diverse effects in response to hypoxia. PI3K mediates neuroprotection against cerebral infarction [14], and it produces phosphoinositides, especially PIP(3,4)P₂ and PIP(3,4,5)P₃, both of which activate ENaC [15]. As an endogenous marker of tumor hypoxia [16], the β isoform of Akt directly interacts and thereby augments ENaC activity [17]. These prior studies suggest that ENaC mediates Na⁺ influx for the maintenance or induction of depolarization in response to hypoxia. Unfortunately, there has been no report on neuronal ENaC activated by onset of hypoxia.

In this study, we observed that acute hypoxia activated ENaC, which depolarized cells by direct influx of Na⁺, in rat PC12 cells, as well as reduced K⁺ outflow through an oxygen-sensing K⁺ channel.

METHODS

Cell culture

PC12 cells were cultured in RPMI-1640 (GIBCO) medium containing 10% fetal bovine serum, 5% horse serum, 100 U/ml of penicillin, and 100 μg/ml of streptomycin under 5% CO₂ at 37°C, and they were transferred every 2-3 days. For electrophysiological recording, cells were subcultured in a 35-mm Petri dish containing round coverslips with a 10 mm diameter.

Solutions

For electrophysiological measurements, standard bath and pipette solutions contained 120 mM NaCl, 5 mM KCl, 1 mM MgCl₂, 1.8 mM CaCl₂, 25 mM NaHCO₃, 1.2 mM KH₂PO₄, and 25 mM NaHCO₃, 1.2 mM KH₂PO₄, and 10 mM HEPES (pH 7.4). Intracellular Ca²⁺ was analyzed by the Fluo-3 AM fluorescence dye detection method using commercial software (MetaMorph V7.7, Molecular Devices, USA).

Electrophysiology

Single-channel currents were recorded using a patch-clamp amplifier (Axopatch 200B, Molecular Devices, USA). Patch pipettes were made from borosilicate glass capillaries (Warner, USA) coated with Sylgard elastomer (Dow Corning, USA). Single-channel currents were sampled at 5 kHz and filtered at 2 kHz using an analog-digital interface (Digitida 1440A, Molecular Devices, USA). Voltage pulse was modified from the ramp protocol used by Choi et al. [18]. Experiments were performed at room temperature.

Electrophysiology

RT-PCR

Total RNA was isolated from PC12 cells using TRIZOL reagent (Invitrogen) according to the manufacturer's protocol. One microgram of total RNA was reverse-transcribed using SuperScript II RT (Invitrogen). 1 μl of oligo dT primer, and 2.5 mM dNTPs at 50°C for 1 h. cDNA was amplified using the following primer sequences: rENaC-α, 5'-AACTCCTTACAGATACACTGAA-3' and 5'-TTCCCTTTCTCATCTGATGGA-3'; rENaC-β, 5'-CAAGAAGAGGCCCATGTTGT-3' and 5'-GTACTGGAGGCTGGATGAAAT-3'; rENaC-γ, 5'-CAAAGATCCCTCATGGAGACG-3' and 5'-ATTTGCCAACTGAGTGTG-3'.

Protein extraction and Western blotting

Lung tissue from rat was lysed with RIPA buffer (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 5 mM EDTA, 1 mM PMSF, and 1% NP-40) containing protease-inhibitor cocktail and homogenized three times for 5 min in a homogenizer (FastPrep-24). PC12 cells were lysed with the same buffer. Whole cell lysates were incubated on ice for 30 min and cleared at 20,000xg for 20 min at 4°C. Proteins were separated by 8% SDS-PAGE and blotted onto PVDF membranes. Blots were incubated overnight at 4°C using ENaC-α antibody (1:250; Santa Cruz Biotechnology).

RESULTS

To confirm the hypoxic conditions induced by the hypoxic solution (HS) saturated with 95% N₂ and 5% CO₂, changes in intracellular Ca²⁺ ([Ca²⁺]ᵢ) and oxygen levels in PC12 cells were compared with those induced by CoCl₂, which is widely used as a chemical hypoxic agent.

When 200 μM CoCl₂ was applied for 3 min, [Ca²⁺]ᵢ was significantly elevated (266.4±39.2% compared to control, n=3). Similarly, N₂ gas mixed gas increased [Ca²⁺]ᵢ, about 2.8 fold within 3 min (280.2±10.0%, n=6) following bubbling (Supplementary Fig. 1). The oxygen level was remarkably reduced to about 40% within 3 min following onset of HS perfusion (normal: 152.76 mmHg and hypoxia: 91 mmHg, n=3). However, N₂ gas did not alter the pH level in the experimental solution. These results show that application of N₂ gas induced acute hypoxia.
Fig. 1. Single channel currents activated by acute hypoxia. (A) Single channel currents recorded at +80 mV (upper) and −80 mV (lower) under normoxic conditions. Channel currents in expanded scale were quoted from each representative recording. (B) All-point histograms analyzed from activities shown in (A). Histograms at +80 mV and −80 mV were separately presented. (C) Channel currents recorded under hypoxic conditions. These activities were recorded in the same cell after switching to hypoxic solution (HS). (D) Histograms analyzed with recording data shown in (C). Arrowheads indicate the peaks activated by infusion of HS. Current recordings were made in cell-attached (CA) configuration with voltage waveforms as presented at the top of (A). Dashed lines (in A, C) denote closed level. Calibration bars are equal to 100 ms and 2 pA.

**Acute hypoxia-induced channel with 30-pS conductance**

In this study, we tried to identify a channel responsible for direct depolarization in response to acute hypoxia. For activation of such channels, single channel currents were recorded before and after hypoxic stress using a cell-attached (CA) configuration, which can protect the environment of the channel.

After forming CA configuration under symmetrical Na⁺-rich (120 mM NaCl) conditions, diverse channels with amplitudes ranging from less than 0.5 pA to ~3.5 pA were observed. Channels could be detected at −80 mV, but not at the depolarization potential (+80 mV, refer to Fig. 1A). Hypoxic stress did not affect channel activities, except for one channel with a conductance of 30-pS and an amplitude of 2.5 pA at +80 or −80 mV (Fig. 1C). This channel was activated ~2 min after application of HS to the bath solution (arrowheads in Fig. 1D). Further, the channel could be detected at a rate of ~13% (one of seven to eight cells tested) under hypoxic conditions, and it showed long-lived opening behavior (mean open time, t_o = 310.7±52.4 ms, n=4) and open probability (P_o) was 0.88±0.02 (n=4).

**The 30-pS channel has Na⁺ selectivity**

Interestingly, the 30-pS channel was activated in response to acute hypoxia. Patch mode was switched to inside-out (IO) configuration from CA in order to identify which ion passes through. Under symmetrical Na⁺-rich conditions, channel activation induced by hypoxia was maintained after excision of the membrane patch, implying that the channel was permeable to Na⁺ or Cl⁻ (Fig. 2A and 2B). This activation disappeared after return to normoxia (data not shown), suggesting that the channel could work in the absence of cytosolic components. In the present study, we did not observe activation of the channel in IO patch mode from CA mode upon exposure to HS.

In the next step, to determine whether or not the channel is permeable to either Na⁺ or Cl⁻, the effects of chloride on channel currents were examined using glutamate. Hypoxia-induced 30-pS channels were not affected by removal of Cl⁻ (Fig. 2D). For the experiments shown in Fig. 2C and D, we clamped the cell interior at +80 mV in order to record currents elicited either by internal (bath) K⁺ or external (pipette) Cl⁻.

To examine cation selectivity, IO membrane patches were excised from cells containing 30-pS channels activated by hypoxia. Channel activity was switched off by recovery to normoxia (upper trace in Fig. 3A) but vigorously reactivated by subsequent application of HS (middle in Fig. 3A and right panel in 3B). When Na⁺ was removed by replacement with equimolar K⁺ or N-methyl-D-glucamine (NMDG) in the bath solution (i.e. cytoplasmic side), channel activity was completely abolished (Fig. 2C and 3A). As shown in the lower trace in Fig. 3A, channel activity (N*P_o) in the presence of NMDG, but not Na⁺, was remarkably reduced to 18.07±6.13% (n=3) compared to that under hypoxic Na⁺-rich conditions. Channel inhibition was also observed upon the removal of Na⁺ in the pipette solution (extracel-
Fig. 2. Hypoxia-activated channel currents impermeable to anions. (A) Preservation of channel activity induced by hypoxia in excised cell-free membrane patch. This channel was silent upon return to normoxia (upper tracings) after switching to inside-out (IO) configuration. This channel was reactivated following perfusion of HS (lower tracings). (B) All-point histograms analyzed from activities recorded in IO patch before (left) and after perfusion of HS (right). These experiments were performed on membrane patches, which were found to contain hypoxia-activated channels in CA mode. Arrowhead shows the peak activated by hypoxia. (C) Effects of potassium and chloride ions on hypoxia-activated channel. Right panel shows inhibition of 30-pS channels by application of potassium glutamate (K-glu). (D) Ineffectiveness of Cl⁻ on hypoxia-elicted activation of 30-pS channel. Right panel shows no effect without Cl⁻ in the pipette. For experiments in (C) and (D), membrane patches were clamped at +80 mV in order to record currents elicited either by internal (bath) K⁺ or external (pipette) Cl⁻. Dashed lines (in A, C) denote closed level. Calibration bars are equal to 100 ms and 2 pA.

Fig. 3. Hypoxia-activated channel selective to Na⁺. (A) Representative tracings of 30-pS channels blocked by removal of Na⁺. Upper two traces were recorded in the presence of Na⁺ in normal (upper) and hypoxic bathing solution (middle), respectively, whereas the lower one was recorded in the presence of impermeable NMDG instead of Na⁺ as indicated by the drawings. (B) Na⁺-dependence of peaks corresponding to activation of channels under hypoxic conditions. Arrowhead designates typical peak for channels with an amplitude of 2.5 pA (at 80 mV) under hypoxic conditions. Note that removal of Na⁺ abolished typical peaks in the histogram labeled as Hypoxia+NMDG-Cl. (C) Summarized bar chart representing the channel activity reduced by removal of Na⁺ under hypoxic conditions. All data were obtained from single channel recordings in IO patches clamped at +80 mV. Asterisk (*) stands for differences from control (p<0.001).
Hypoxia-Activated Na⁺ Channel

The Na⁺-selective 30-pS channel is blocked by lidocaine and amiloride

Since the channel activated by acute hypoxia in PC12 cells was a Na⁺ channel, we explored its pharmacology using neuronal Na⁺ channel blockers such as lidocaine and amiloride. Channel activity was dramatically reduced to 80% by 100 μM lidocaine (Supplementary Fig. 2).

Regarding Na⁺ channels in neurons, most voltage-independent Na⁺ channels are rapidly inactivated, with mean open times at best from 16 ms to 0.7 ms [19,20]. On the contrary, voltage-independent Na⁺ channels such as epithelial Na⁺ channel (ENaC)/degenerin family members are open longer than voltage-dependent Na⁺ channels, with relatively high Po up to 0.9 [21]. According to such simple kinetic criteria, we found that the 30-pS channel shared properties with ENaC, including conductance of 30 pS, 310 ms-long mean open time, and Po of 0.88 at −80 mV, as stated with Fig. 1, which are close to those of epithelial α-subunit Na⁺ (ENaC-α) channel [22].

To determine whether or not the 30-pS channel shares common pharmacology with ENaC, we applied amiloride, which is widely used to specifically block ENaC and acid-sensing ion channel (ASIC). As shown in Fig. 4, less than 1 min following treatment with a nanomolar amount of amiloride (200 nM), channel activity was significantly reduced to about 30.58±10.62% (n=3) compared to that of control (p<0.05). Since ASICs require just a few tens of μM amiloride to achieve 50% inhibition (IC₅₀), and also Ca²⁺-permeability [23], the 30-pS channel was determined to be an ENaC rather than the ASIC family member.

ENaC is endogenously expressed in PC12 cells

Previous results have strongly suggested that the 30-pS channel activated by hypoxia demonstrates properties resembling ENaC. Therefore, ENaC RNA and protein were confirmed by using RT-PCR and Western blotting in order to determine that ENaC is endogenously expressed in PC12 cells. Using constructed subunit primers of ENaC consisting of α, β, and γ subunits, we confirmed the presence of RNAs corresponding to three subunits in PC12 cells (Fig. 5A). Finally, we demonstrated the endogenous protein expression of ENaC using antibody specific to rat ENaC-α as well as RNA expression using RT-PCR. As shown in Fig. 5B, ENaC protein was detected by Western blotting using ENaC-α-specific antibody and lung tissue as a positive con-
trol (left lane in Fig. 5B). These results confirm that the 30-pS channel could be a member of the ENaC family activated by acute hypoxia in rat PC12 cells.

Taken together, the results in the present study suggest the presence of an ENaC-like channel activated by acute hypoxia in neuronal rat PC12 cells.

DISCUSSION

The present study provides evidence for the presence of a Na+ channel activated by acute hypoxia in PC12 cells. This result was supported by the following: the 30-pS channel was activated under hypoxic conditions and its activity disappeared upon return to normoxia (Figs. 1 and 2). Second, the channel was selective for Na+, whereas it was unaffected by Ca2+, Cl– or K+. Finally, the channel was blocked by lidocaine, a neuronal Na+ channel blocker.

In contrast to voltage-gated Na+ channels, which are rapidly inactivated, opening of the hypoxia-activated channels lasted longer than 300 ms, and the inactivation process was not observed when the recording lasted for longer than 10 sec (P<0.88, refer to Fig. 1C) [21]. The hypoxia-activated channel shared properties similar to those of ENaC, a member of the vertebrate ENaC/ASIC family [24]. On the other hand, this channel revealed differences in kinetics and pharmacology from those of ASIC. Further, it was sensitively blocked with a low dose (200 nM) of amiloride, whereas ASICs are less sensitive (up to hundreds μM) and rapidly inactivated [25,26]. Since the pH level did not respond to bubbling of anoxic N2 gas, it was identified as an ENaC rather than ASIC. This suggestion was supported by the expression of ENaC protein in PC12 cells (Fig. 5), although direct evidence is required that heterologously expressed ENaC is also sensitive to acute hypoxia.

The Na+ channel conductance activated by acute hypoxia was larger than those reported in studies on endogenous ENaCs. The conductance was about 30 pS (2.5 pA at ~80 mV) in the present study, whereas a unit conductance of ENaC has been reported at best to be less than 10 pS [27-29]. If the hypoxia-activated Na+ channel belongs to the ENaC family, then a larger conductance would conflict with those of ENaCs which are believed to form a heteromeric channel composed of four subunits α, β, γ, and δ [21]. Only a few reports have reported conductance levels of 21~40 pS from recordings performed in the planar lipid bilayer or on homomeric ENaC-α [30-32], implying that the conductance could be modulated by β- and γ-subunits as well as cytoplasmic factor(s). For instance, reduction of disulfide bonds with dithiothreitol from ENaC incorporated into planar lipid bilayers reduced single channel conductance from 40 pS to 13 pS [30]. Considering this redox modulation of ENaC conductance, it is likely that oxidation can elevate unit ENaC conductance.

The mechanism underlying ENaC activation during acute hypoxia is difficult to explain based solely on this study. At present, there is a general agreement that intermediate or long-term hypoxia inhibits ENaC activity in the alveolar epithelia, whereas the effect of acute hypoxia on neuronal ENaC unclear. This channel is upregulated by proteolytic enzymes and phosphoinositides (PIs), including P(4,5)P2 and P(3,4,5)P3. Both PIs can interact with the N terminus of ENaC-β as well as the proximal C terminus of ENaC-γ, thereby enhancing ENaC activity. Depletion of membrane P(4,5)P2 inhibits ENaC activity [33]. On the other hand, hypoxia provokes intense Ca2+ mobilization as well as the level of P(4,5)P2, stimulating calpain proteolytic activity [34]. Thus, it can be stated that acute hypoxia activates proteolytic enzymes such as calpain and proteasins, which, in turn, trigger the activation of ENaC [35]. In order to mimic hypoxia, we did not add ATP to the bath. Therefore, the effect of PIs on the channel activity was greatly limited since ATP is essential to the modulation of membrane-delimited PI kinases such as PLK or PLC. Throughout the experiments, activation of the 30-pS channel was not spontaneous and usually occurred 2~3 min after exposure to hypoxia. These channels in the excised patch were re-activated without any alteration in unit conductance in response to hypoxia, whereas they became silent under normoxia (refer to Figs. 2–4). These findings strongly suggest that channel conductance can be separately regulated along with channel activity in response to hypoxia.

There have been many studies on K+ channel activity in response to acute hypoxia [4,5,7]. In a study on Kv1.2 and Kv1.1 in PC12 cells, hypoxia inhibited oxygen-sensitive K+ (Ko2) current, resulting in membrane depolarization and peak inhibition within 2 min, which correlated well with the time course of membrane depolarization. The amplitude of depolarization depended on the grade of hypoxia, i.e., level of low P02 [6]. However, it was also revealed that exposure to hypoxia for more than 2 min caused further depolarization (see details in Fig. 2A and 2B in reference 6). For persistent hypoxia, membrane depolarization could continue due to the influx of cations through another channel in addition to the closure of Ko2 channels. In this context, 30-pS Na+ channels are proper since they are activated at least 2~3 min following exposure to hypoxia, facilitating further depolarization.

The main focus of this study was to identify a channel other than a Ko2 channel activated by acute hypoxia. Our study was performed in PC12 cells using solution half saturated with oxygen (91 mmHg O2) in normal solution (152 mmHg O2). This is a disadvantage of our study compared to those performed under almost anoxic conditions (<20% of normal). However, our conditions are more physiological, considering that the O2 level cannot reach an anoxic state within the few minutes of acute ischemic injury. Under these conditions, the present study first identified that the Na+ channel was activated by acute hypoxia and has a single channel conductance of 30 pS with ENaC-like properties. However, it is unclear whether or not this channel protects from hypoxia/ischemia. With respect to hypoxia,
further study on the ENaC-like channel can provide therapeutically implications for neurons in response to hypoxia.

SUPPLEMENTARY MATERIALS

Supplementary data including one figure can be found with this article online at http://pdf.medrang.co.kr/paper/pdf/Kjpp/Kjpp017-01-09-s001.pdf.

ACKNOWLEDGEMENTS

This study was supported by the Basic Science Research Program through the National Research Foundation of Korea (NRF) funded by the Ministry of Education, Science and Technology (No. 2012-0000300).

REFERENCES

1. Cross JL, Meloni BP, Bakker AD, Lee S, Knackey NW. Modes of neuronal calcium entry and homeostasis following cerebral ischemia. Stroke Res Treat. 2010;2010:316862.

2. Fung ML. Role of voltage-gated Na⁺ channels in hypoxia-induced neuronal injuries. Clin Exp Pharmacol Physiol. 2000; 27:569-574.

3. Ratan RR, Siddiq A, Smirnova N, Karpisheva K, Haskew-White J, Warnock DG, Eaton DC. Determination of epithelial Na⁺ channel subunit stoichiometry from single-channel conductances. Am J Physiol. 2002;283:C658-665.

4. Conforti L, Seta K, Lobayashi S, Beiter-Johnson D, Rast RT, Yuan Y, Kim KW, Kim RH, Seta K, Millhorn DE. The molecular basis of O₂-sensing and hypoxia tolerance in phaeochromocytoma cells. Comp Biochem Physiol B Biochem Mol Biol. 2001;128:187-204.

5. Conforti L, Millhorn DE. Selective inhibition of a slow-inactivating voltage-dependent K⁺ channel in rat PC12 cells by hypoxia. J Physiol. 1997;502:293-305.

6. Zhou WH, Conforti L, Czezyk-Krasieka MF, Millhorn DE. Membrane depolarization in PC-12 cells during hypoxia is regulated by an O₂-sensitive K⁺ current. Am J Physiol. 1996;271:C658-665.

7. Conforti L, Seta K, Lobayashi S, Millhorn DE. O₂-sensitive K⁺ channels: role of the Kv1.2 subunit in mediating the hypoxic response. J Physiol. 2000;524:783-793.

8. Kim D, K⁺ channels in O₂ sensing and postnatal development of carotid body glomus cell response to hypoxia. Respir Physiol Neurobiol. 2013;185:44-56.

9. López-López JR, González C, Ureña J, López-Barneo J. Low pO₂ selectively inhibits K channel activity in chemoreceptor cells of the mammalian carotid body. J Gen Physiol. 1989;93:1001-1015.

10. López-López JR, González C, Perez-Garcia MT. Properties of ionic currents from isolated adult rat carotid body chemoreceptor cells: effect of hypoxia. J Physiol. 1997;499:429-441.

11. Ji HL, Benos DJ. Degenerin sites mediate proton activation of deltabetagamma-epithelial sodium channel. J Biol Chem. 2004;279:26939-26947.

12. Wesc D, Miranda P, Alfonso-Oromas D, Althaus M, Castro-Hernández J, Dominguez J, Morty RE, Claus W, González-Hernández T, Alvarez de la Rosa D, Giraldez T. The neuronal-neuronal-specific S0K1.1 kinase regulates 5-epithelial Na⁺ channel independently of PY motifs and couples it to phospholipase C signaling. Am J Physiol Cell Physiol. 2010;299:C779-790.

13. Seta KA, Spicer Z, Yuan Y, Lu G, Millhorn DE. Responding to hypoxia: lessons from a model cell line. Sci STKE. 2002;2002:re11.

14. Abe E, Fujiki M, Nagai Y, Shiqi K, Kubo T, Ishii K, Abe T, Kobayashi H. The phosphatidylinositol-3 kinase/Akt pathway mediates geranylgeranylated-coupled neuroprotection against cerebral infarction in rats. Brain Res. 2010;1330:151-157.

15. Tong Q, Gumper N, Medina JI, Shapiro MS, Stockand JD. Direct activation of the epithelial Na⁺ channel by phosphatidylinositol 3,4,5-trisphosphate and phosphatidylinositol 3,4-bisphosphate produced by phosphoinositide 3-OH kinase. J Biol Chem. 2004;279:22654-22661.

16. Chen Y, Shi G, Xia W, Kong C, Zhao S, Gao AF, Chen FY, Yang GP, Gaecia AD, Le QT, Koong AC. Identification of hypoxia-regulated proteins in head and neck cancer by proteomic and tissue array profiling. Cancer Res. 2004;64:7302-7310.

17. Lebowitz J, Edinger RS, An B, Perry CJ, Onute S, Kleyman TR, Johnson JP. Ikappab kinase-beta (ibkbeta) modulation of epithelial sodium channel activity. J Biol Chem. 2004;279:11581-11589.
Miyakawa A, Nishijo H, Yamashita J, Kawashima S, Ono T, Yoshioka T. Transient brain ischaemia provokes Ca\textsuperscript{2+}, PIP\textsubscript{2} and calpain responses prior to delayed neuronal death in monkeys. *Eur J Neurosci.* 1996;8:1932-1944.

35. Donaldson SH, Hirsh A, Li DC, Holloway G, Chao J, Boucher RC, Gabriel SE. Regulation of the epithelial sodium channel by serine proteases in human airways. *J Biol Chem.* 2002;277:8338-8345.