Intracellular Acid-Extruding Regulators and the Effect of Lipopolysaccharide in Cultured Human Renal Artery Smooth Muscle Cells

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Abstract

Homeostasis of the intracellular pH (pHi) in mammalian cells plays a pivotal role in maintaining cell function. Thus far, the housekeeping Na+/H+ exchanger (NHE) and the Na+/HCO3− co-transporter (NBC) have been confirmed in many mammalian cells as major acid extruders. However, the role of acid-extruding regulators in human renal artery smooth muscle cells (HRASMCs) remains unclear. It has been demonstrated that lipopolysaccharide (LPS)-induced vascular occlusion is associated with the apoptosis, activating calpain and increased [Ca2+], that are related to NHE1 activity in endothelia cells. This study determines the acid-extruding mechanisms and the effect of LPS on the resting pH and active acid extruders in cultured HRASMCs. The mechanism of pH recovery from intracellular acidosis (induced by NH4Cl-prepulse) is determined using BCECF-fluorescence in cultured HRASMCs. It is seen that (a) the resting pHi is 7.19±0.03 and 7.10±0.02 for HEPES- and CO2/HCO3−-buffered solution, respectively; (b) apart from the housekeeping NHE1, another Na+/coupled HCO3− transporter i.e. NBC, functionally co-exists to achieve acid-equivalent extrusion; (c) three different isoforms of NBC: NBCn1 (SLC4A7; electroneutral), NBCe1 (SLC4A4; electrogenic) and NBCe2 (SLC4A5), are detected in protein/mRNA level; and (d) pH and NHE protein expression/activity are significantly increased by LPS, in both a dose- and time-dependent manner, but NBCs protein expression is not. In conclusion, it is demonstrated, for the first time, that four pH, acid-extruding regulators: NHE1, NBCn1, NBCe1 and NBCe2, co-exist in cultured HRASMCs. LPS also increases cellular growth, pHi and NHE in a dose- and time-dependent manner.

Introduction

Atherosclerotic renal artery stenosis (ARAS) is the most common primary disease of the renal arteries and it is associated with hypertension [1,2]. In addition to threatened renal function, ARAS-induced atherosclerotic renovascular disease with renal failure and can result in mortality [3–6]. Many important cellular functions are affected by a change in pHi. Mechanisms such as cell volume [7], the permeability of ion channels [8], enzyme catalysts [9], cell differentiation, growth and apoptosis are all sensitive to changes in pHi [10–12]. These disturbances in pHi have also recently been claimed to be responsible for the development of hypertension and vascular atherosclerosis in animal models [12,13]. However, the evaluation of the biological effect on the cell of acid-base transport in vascular cells is difficult, especially in human tissues/cells.

The pHi in mammalian cells remains within a narrow range (7.0–7.2) because of the combined operation of active transmembrane transporters and passive intracellular buffering power [14]. The membrane transporters can be divided into two main categories: acid extrusion carriers and acid loading carriers. Acid extrusion carriers, such as Na+/H+ exchanger (NHE) and Na+/HCO3− co-transporter (NBC), are activated when cells are become acidic (pH < 7.1) [12,15,16]. Net acid extrusion from vascular smooth muscle cells (VSMCs) in rat and mice mesenteric small arteries is mediated by the Na+/HCO3− co-transporter NBCn1 (slc4a7) and the Na+/H+ exchanger NHE1 (slc9a1) [12,15,17,18]. NHE mediates the electroneutral exchange of extracellular Na+ for intracellular H+ [12,19,20]. pH recovery in HEPES buffered media (HCO3−-free condition) is inhibited by the removal of extracellular Na+ or by the addition of amiloride or Hoe 694 (3-methylsulfonyl-4-piperidinobenzoyl, guanidine hydrochloride), a compound that inhibits NHE activity because of its high affinity and selectivity [21]. NHE proteins comprise a family of at least ten NHE members (NHE 1–9 and sperm NHE), each of which is expressed by a separate gene [22,23]. Different subtypes of NHE occur in different cell types. In rat/mouse VSMCs, the transporter is well defined at the molecular level as NHE1 [12,18,24].
Na⁺-HCO₃⁻-dependent transporter is largely, 4-dioisothiocyanatostilbene-2,2'-disulphonic acid (DIDS)-sensitive (56% to 91%), and it is amiloride- and HOE 694-resistant [12,16,25-29]. It is also inhibited by the removal of external Na⁺ [29]. Relevant molecular candidates for Na⁺-dependent bicarbonate transport include at least five members of the slc4 family [12,27,30]. It has been recently found that NBCn1 (SLC4A7) mediates the Na⁺-dependent bicarbonate transport that is important for acid extrusion in the smooth muscle cells of mouse mesenteric, coronary, and cerebral small arteries [12,15,17,27,31]. However, thus far there have been no related reports about active acid extruding transports in human renal artery smooth muscle cells (HRASMCs).

Lipopolysaccharides (LPS) of gram-negative bacteria play a pivotal role in septic shock syndrome in humans [32]. It has been demonstrated that mRNA and protein expression of toll-like receptor 4 (TLR4) are up-regulated by LPS in human aortic smooth muscles, in a dose- (10–1000 ng/ml) and time-dependent (0–48 hr) manner [33]. In human arterial smooth muscle, LPS (10 ng/ml) has also been found to induce mRNA and protein expression of matrix metalloproteinases-9 (MMP-9), in a TLR4/ NF-κB-dependent manner [34]. Previous studies have shown that the inhibition of NHE1 has anti-apoptotic effects [35-37]. A very recent study in HUVECs has further demonstrated that LPS inhibits NHE1 in a, time-dependent manner that is associated with an increased [Ca²⁺], which results in enhanced calpain activity and apoptosis, via NHE1-dependent degradation of Bcl-2 [38]. However, thus far, the effect of LPS on resting pH and acid-extruding regulators has not been determined in HRASMCs. In this study, apart from checking the resting pH and the possible acid extruders is also studied.

In brief, this study demonstrates, for the first time, that two different types of acid-extruders: NHE1 and NBC, functionally co-exist in cultured HRASMCs. Three different isoforms of Na⁺-coupled HCO₃⁻ co-transporter: NBCn1 (SLC4A7; electroneutral), NBCc1 (SLC4A4; electrogenic) and NBCe2 (SLC4A5), are detected in protein/mRNA level. It is also demonstrated that LPS increases cellular growth, pH and NHE in a dose- and time-dependent manner, but does not affect the activity of NBCs.

**Materials and Methods**

**Human renal artery smooth muscle cells (HRASMCs)**

With the approval of the Institutional Review Board of Tri-Service General Hospital, National Defense Medical Center (TSGHIRB No. 1-101-05-065) and with prior written informed consent of patients, 9 renal artery segments (5 male: 62.5±6.3 yrs old, 4 female: 60.6±7.2 yrs old) were collected from surgically- leftover specimens of human renal arteries during renal transplant surgery at Tri-service General Hospital, Taipei, Taiwan. Primary HRASMCs were isolated by the explant technique which has been described in detail in Fletcher et al [39] and cultured in HAM's F12K medium containing 10% fetal bovine serum (FBS) (GIBCO, Grand Island, NY, USA), 100 U/ml penicillin, 100 mg/ml streptomycin, and 200 mM L-glutamate in a humidified incubator (at 37°C and 5% CO₂). The primary HRASMCs were used for experiments between 3 and 8 passages. The preparations were then perfused with oxygenated Tyrode solution, which was either 100% O₂ for nominally bicarbonate-free Tyrode solution or 5% CO₂/95% O₂ for bicarbonate-containing Tyrode solution, at 37°C, pH 7.40±0.02 for experiments.

**Immunocytochemistry**

Cells were cultured on a 6-well plate (Macalaster Bicknell, New Haven, CT) for 1–3 days. Cells were subsequently washed twice in phosphate-buffered saline (PBS). After washing, cells were fixed in 4% paraformaldehyde for 30 min at room temperature, then washed twice in PBS and blocked and permeabilized in PBS containing 0.3% triton and 5% normal goat serum for 60 min, and finally washed in PBS and incubated overnight with primary antibodies at 4°C. Cells were washed four times in PBS and labeled with secondary antibodies for 1 h in the dark. After labeling, cells were washed with PBS and incubated with DAPI (1: 200) for 40 min. Cells were washed twice more and mounted onto slides with Gel/Mount (Biomedia Corp., Forest City, CA). Images were acquired with an OLYMPUS 200M (Japan) microscope system.

**Western blot analysis of the NHE-1~3 protein and SLC4 family of HCO₃⁻ transporters**

Cell lysates were prepared using a 300 µl/well of six-well plates of RIPA/NP-40 lysis buffer ( 5 mM Tris pH 7.4, 30 mM NaCl, 1 mM PMSF, 1 µg/ml aprotinin). A 50 µg of total protein per sample was then subjected to 10% PAGE, transferred to a PVDF membrane and blocked for 2 h with 5% fat-free milk in Tris-buffered saline/0.1% Tween 20 (TBST). After three washes with TBST, the membranes were exposed to a 1: 1000 dilution of a mouse anti-human NHE1~3 antibody (Millipore, Long Beach, CA, USA) [40] or SLC4 family of HCO₃⁻ transporters: NBCe1 (Millipore, Long Beach, CA, USA) [41,42], NBCe2 (Abgent, San Diego, CA, USA), NBCn1 (Abgent, San Diego, CA, USA) and NDCBE (GeneTex, San Antonio, TX) at 4°C overnight. Following three washes with TBST, the membrane was exposed to a 1: 15000 dilution of goat anti-mouse IgG-HRP conjugate (Millipore, Long Beach, CA, USA) for 1 h and washed repeatedly with TBST. Chemiluminescence was detected using the SuperSignal Substrate (PIERCE, Rockford, IL, USA). Loading control was assessed by the detection of β-actin. NHE-1~3, SLC4 family of HCO₃⁻ transporters: NBCe1 (Genesee Scientific, MA, USA) with 10 μg cDNA and 1 pmol of each primer (NBCe1: forward: 5'-GGTGTGCAGTTCTGATCCTGTCGTC-3' and reverse: 5'-GTTGACTGTCCTCAGTTC-3). Cells were cultured on a 6-well plate (Macalaster Bicknell, New Haven, CT) for 1–3 days. Cells were subsequently washed twice in phosphate-buffered saline (PBS). After washing, cells were fixed in 4% paraformaldehyde for 30 min at room temperature, then washed twice in PBS and blocked and permeabilized in PBS containing 0.3% triton and 5% normal goat serum for 60 min, and finally washed in PBS and incubated overnight with primary antibodies at 4°C. Cells were washed four times in PBS and labeled with secondary antibodies for 1 h in the dark. After labeling, cells were washed with PBS and incubated with DAPI (1: 200) for 40 min. Cells were washed twice more and mounted onto slides with Gel/Mount (Biomedia Corp., Forest City, CA). Images were acquired with an OLYMPUS 200M (Japan) microscope system.

**Reverse Transcription – Polymerase Chain Reaction (RT-PCR)**

Total RNA was isolated using GeneJET RNA Purification Kit (Thermo Scientific, MA, USA) from human renal artery smooth muscle cells (HRASMCs). The RNA (1 μg) was reverse transcribed using Maxima First Strand cDNA Synthesis Kit (Thermo Scientific, MA, USA). PCR (DreamTaq Master Mix, Thermo Scientific, MA, USA) with 10–50 ng cdNA and 1 pmol of each primer (NBGc1-SLC4A4: forward: 5'-GGTGTGCAGTTCTGATCCTGTCGTC-3' and reverse: 5'-GTTGACTGTCCTCAGTTC-3'; NBCe2-SLC4A5: forward: 5'-ATCTTTCATGGACAGTGGAT-3' and reverse: 5'-GGTGTGCAGTTCTGATCCTGTCGTC-3').
CAGCA- GATCAC-3'; reverse: 5'-TGCTTGCTGGCAT-CAGGAAG-3'; NBCn1-SLC4A7: forward: 5'-CAGATGCAAG-CAGCCTGTGTG-3'; reverse: 5'-GGTCCATGATG-ACCA-CAAGCTG-3'; NDCBE1-SLC4A8: forward: 5'-GCTCAAGA-AAGGCTG-TGGCTAC-3'; reverse: 5'-CATGAAGACTGAGCACATC-3) was performed for 30 cycles after 15 min at 95°C: 95°C, after which denaturation was performed for 30 s at 60°C, annealing for 30 s, and elongation at 72°C for 1 min. Negative controls included omission of cDNA. PCR for actin was performed to validate each template. PCR products were separated by 2% agarose gel electrophoresis with ethidium bromide and photographed under ultraviolet illumination. All primer pairs were confirmed by nucleotide sequencing representative PCR products.

**MTT assay**

Cells were seeded onto 24-well plates at a density of 3×10⁴ cells/well and grown for up to 24 hr with 1 ml serum free medium. The growth medium was replaced on day 2 with a different dosage of LPS (1~10000 ng/ml) for another 24 hr. Cell viability was assessed by using the 3-(4, 5-dimethyl-2-thiazolyl)-2,5-diphenyl-2H-tetrazolium bromide (MTT) assay (Sigma) according to the manufacturer’s protocol. In brief, an ELISA (Enzyme-linked immunosorbent assay) reader was used to detect the...
absorbance of 490 nm after 10% MTT had been reacted with different tested cells for 3 hr.

**Measurement and calibration of the intracellular pH**

Measurement of the pH$_i$ has been described in detail in our previous reports [16,26]. In brief, the pH$_i$ in the cultured HRASMC was measured using the pH-sensitive, dual excitation single-emission fluorescent dye, 2',7'-bis(2-carboxyethyl)-5(6)-carboxyfluorescein-acetoxyethyl (BCECF-AM) (Molecular Probes). The preparations were loaded with BCECF-AM (5 μM) by incubating them for 30 min at room temperature and exciting them alternately with 490 and 440 nm wavelength light. The BCECF fluorescence emission ratio of the 510 nm emission at 490 nm and 440 nm excitation (490/440) was calibrated using the K$^+$-nigericin method [16]. Briefly, this method consisted of exposing a BCECF-loaded cell to the six nigericin calibration solutions (listed below in the *Solution* section) that clamps pH$_i$ to the value of pH$_o$, of the calibration solution. Fig. 1A showed the emission ratio changes seen on perfusing human artery smooth muscle cells with calibration solutions with different 6 pH values (5.5–9.5) in the presence of 10 μM nigericin. The emitted ratio 510 nm emission at 490 nm and 440 nm excitations (R; R = F$_{490}$/F$_{440}$) was increased as the pH value of superfusing solution was increased. R$_{max}$ and R$_{min}$ are, respectively, the maximum and minimum ratio values for the data curve. The fluorescence of BCECF at 490 nm to 440 nm is a function of pH$_i$ and the overall sampling rate in the experiment was 0.5 Hz for the recorded fluorescence ratio (490 nm/440 nm). Using the linear regression fit of the data (shown in the Fig. 1B) obtained from 6 calibration experiments similar to that shown in Fig. 1A, the mean apparent dissociation constant (pKa) at 37°C was found to be 7.18, very close to the value determined by our previous study of the human heart, as well as the value determined by other investigators [16,43,44]. The following equation [45] was used to convert the fluorescent ratio in to pH$_i$:

$$\text{pH}_i = \text{pK}_a + \log[(R_{\text{max}} - R)/(R - R_{\text{min}})] + \log(F_{440\text{min}}/F_{440\text{max}})$$

where R is the ratio of the 510 nm fluorescence at 490 nm and 440 nm excitation, R$_{max}$ and R$_{min}$ are, respectively, the maximum and minimum ratio values from the data curve and the pK$_a$ (log of dissociation constant) is 7.18. F$_{440\text{min}}$/F$_{440\text{max}}$ and F$_{490\text{max}}$/F$_{490\text{max}}$ is the ratio of fluorescence measured at 440 nm of R$_{min}$ and R$_{max}$, respectively.

**Experimental alteration of intracellular pH- weak acid/ base pre-pulse technique**

NH$_4$Cl pre-pulse techniques were used in the present work to induce acute acid loading [44,46]. NH$_4$Cl pre-pulses were achieved with (~10 minute) extracellular exposures to 20 mM NH$_4$Cl. Briefly, the mechanism of the NH$_4$Cl prepulse technique relies upon the characteristic of incomplete dissociation. For example, the details of NH$_4$Cl prepulsing procedures, used in the present study, are given below. It can be explained in terms of four phases as shown in Fig. 2: rapid entry (see phase 1 in left part of Fig. 2A), slow recovery (see phase 2 in left part of Fig. 2A), rapid exit (see phase 3 in left part of Fig. 2A), and pH$_i$ regulation (see phase 4 in left part of Fig. 2A) that the sudden acidosis activates pH$_i$ regulatory proteins in the membrane, for instance Na$^+$-H$^+$ exchanger and Na$^+$-HCO$_3^-$ cotransporter. Throughout the whole experiment, the change of pH$_i$ induced by the tested drug was compared around the 3rd min after treating the drug (the testing pH$_i$ is around 6.90±0.15), unless otherwise stated. The background fluorescence and auto-fluorescence were small (<5%) and have been ignored.

**Chemicals and solutions**

*Standard HEPES-buffered Tyrode solution* (air equilibrated) contained (mM): NaCl, 140; KCl, 4.5; MgCl$_2$, 1; CaCl$_2$, 2.5; glucose, 11; HEPES, 20; pH adjusted to 7.4 with 4N NaOH. Unless otherwise stated, pH adjustments of all HEPES-buffered solutions were performed at 37°C (these adjustments included those where ionic-substitutions were made, see below). *Standard bicarbonate-buffered Tyrode solution* (equilibrated with 5% CO$_2$/23 mM HCO$_3^-$) was the same as above, except that the sodium chloride concentration was reduced to 117 mM, and 23 mM NaHCO$_3$ was added instead of the HEPES (pH 7.40 at 37°C).

*In-substituted solutions*: In a Na$^+$-free, HEPES-buffered Tyrode solution, (air equilibrated) contained (mM): N-methyl-D-glucamine (NMDG), 140; KCl, 4.5; MgCl$_2$, 1; CaCl$_2$, 2.5; glucose, 11; HEPES, 20, and the pH was adjusted to 7.4 with HCl. The Na$^+$-free CO$_2$/HCO$_3^-$-buffered Tyrode solution (equilibrated with 5% CO$_2$/23 mM HCO$_3^-$) contained (mM): NMDG, 140; KCl, 4.5; CaCl$_2$, 2.5; MgCl$_2$, 1; glucose, 11, and pH was adjusted to 7.4 at 37°C with HCl under the condition of saturating with 5% CO$_2$/95% O$_2$. When 20 mM ammonium chloride was used, it was added directly as a solid to solution without osmotic compensation. 4,4'-disothiocyanostilbene-2,2'-disulphonic acid (DIDS) and HOE 694 (3-methylbulyl)-1-piperidinobenzoyl, guanidine hydrochloride) were added, as solids, to solutions shortly before use.

*Nigericin calibration solutions* contained (mM): KCl, 140; MgCl$_2$, 1; 10 μM nigericin; buffered with one of the following organic buffers: 20 mM 2-(N-morpholino)ethanesulfonic acid (MES, pH 5.5), 20 mM HEPES (pH 7.5) or 20 mM
3-(cyclohexylamino)-2-hydroxy-1-propane-sulphonic acid (CAPSO), pH 9.5, and were adjusted (37°C) to the correct pH with 4N NaOH.

HOE 694 was kindly provided by Hoechst Aktiengesellschaft (Germany). All other chemicals were from Sigma (UK) and Merck (UK).

Statistics
All data are expressed as the mean ± the standard error of the mean (SEM) for n preparations. Statistical analysis was performed using one-way analysis of variance (one-way ANOVA) with Schef"e’s posterior comparison. A P value smaller than 0.05 was regarded as significant.

Results
The isolation of human renal artery smooth muscle cells (HRASMCs) from tissue
This study successfully isolated HRASMCs from human artery tissue, using the so-called explant method. The HRASMs were significantly migrate out of the artery tissue on the 10th day and the 20th day in a time-dependent manner, as shown in Fig. 1C and Fig. 1D, respectively. The immunocytochemistry technique was also used in order to determine the purity of HRASMCs. In brief, HRASMCs were stained with 2-SM-actin, the specific monoclonal antibody, as a smooth muscle differentiation marker (Fig. 1E; green color) and DAPI, the nuclei counterstained marker (Fig. 1F; blue color). The cell-pattern of Fig. 1G, combined with those from Fig. 1E and Fig. 1F, is almost the same as that of Fig. 1E and Fig. 1F. This clearly indicates that the cells are HRASMCs. Therefore, a single HRASMC was successfully derived from tissue from a human renal artery using the explant technique.

The functional existence of a Na⁺-H⁺ exchanger (NHE)
In order to determine whether an acid-extrusion mechanism exists in the cultured HRASMCs, the experiments were first performed in HEPES-buffered superfusate (nominally free of CO₂/HCO₃⁻). The steady-state pHᵢ value for the cultured HRASMCs was measured as 7.19±0.02 (n = 24), in CO₂/HCO₃⁻ buffered Tyrode solution. This steady-state pHᵢ value for HRASMCs is much lower than that in HEPES-buffered Tyrode solution and the value is similar to that reported previously for mature mammalian cells, in both animal and human models [14,16].

The left part of the traces shown in Figs. 3A and 3C show the pHᵢ recovery from an acid load that is induced in a CO₂/HCO₃⁻ environment. The removal of Na⁺ from the 5% CO₂/HCO₃⁻ Tyrode solution completely inhibits pHᵢ recovery, following NH₄Cl-induced acidosis, as shown in the right part of Fig. 3A. The histogram in Fig. 3B shows the pHᵢ recovery rate, which is estimated at pHᵢ 6.83±0.03 and is the average value for six experiments in HRASMCs. This data suggests that this HCO₃⁻-dependent acid-extrusion mechanism is also Na⁺-dependent. The pHᵢ recovery is partially blocked, as expected, in the presence of HOE 694, as shown in the second part of Fig. 3C. The second column of the histogram in Fig. 3D shows the pHᵢ recovery slope, after acid loading, for seven experiments (estimated at pHᵢ 6.81±0.08) in HRASMCs. The significant difference between the first (control) and the second column (in presence of HOE 694 in a 5% CO₂/HCO₃⁻ solution) shows that, apart from NHE, another HCO₃⁻-dependent acid-extrusion mechanism is involved in the pHᵢ recovery in 5% CO₂/HCO₃⁻ Tyrode solution. It has been demonstrated that the sulfide drug, DIDS (0.4 mM), inhibits NBC effectively [12,16,25-28], so a further test was undertaken to determine whether DIDS inhibits this HCO₃⁻-dependent, but HOE 694-independent, acid-extrusion mechanism, in the HRASMCs. As shown in the right part of Figs. 3C, a combination of 30 μM HOE 694 and 0.4 mM DIDS entirely inhibits the pHᵢ recovery, following induced intracellular acidosis with 5% CO₂/HCO₃⁻ Tyrode solution (right part of Fig. 3C), but DIDS alone does not (the third right part of Fig. 3C). The third and fourth histograms in Fig. 3D show the pHᵢ recovery rate, as an average for 7 experiments, which is similar to that shown in Fig. 3C (estimated at pHᵢ 6.80±0.08). This data demonstrates that this Na⁺- and HCO₃⁻-dependent acid-extrusion mechanism relies on the NBC. In other words, this is the first functional evidence that both NHE and NBC play an important role in pHᵢ regulation through acid extrusion in cultured HRASMCs.

The effect of LPS on the protein expression of NHE, NBC and on intracellular resting pH
In order to further identify the isoform(s) of the functional NHE and NBC observed previously, the Western blot technique (see Materials and Methods for details) was used to test the response for treating a mouse antihuman NHE1-3 antibody and the SLC4 family of HCO₃⁻ transporters: SLC4A4 (NBCe1), SLC4A5 (NBCe2), SLC4A7(NBC1) and SLC4A8 (NBCe4), respectively, in cultured HRASMCs. As shown in the left part of Fig. 4A, the isoform of NHE is purely NHE1 isoform in HRASMCs.

It is also significant that three Na⁺-coupled HCO₃⁻ co-transporters: NBC1 (SLC4A7; electroneutral), NBCe1 (SLC4A4; electrogenic) and NBCe2 (SLC4A5), co-exist in the cultured HRASMCs, as shown in the left part of Fig. 4C. This is different to the case for animal models, because only NBCn1 (SLC4A7) mediates the Na⁺-dependent bicarbonate transport that is
For the discovery of 3 different NBC isoforms in cultured HRASMCs, the molecular identity of the transporter responsible was determined, using the reverse transcription – polymerase chain reaction (RT-PCR) technique. As shown in Fig. 4E, the total mRNA of NBCe1 (SLC4A4), NBCe2 (SLC4A5), NBCn1 (SLC4A7) and NDCBE1 (SLC4A8) are clearly expressed, as shown in Fig. 4C. Note that the protein of NDCBE1 (SLC4A8) was not detected (Fig. 4C), while that of the mRNA level was detected (Fig. 4E). Whether this difference in protein and mRNA in NDCBE1 is caused by the less specificity of antibody or a scarcity of protein requires further study.

The significant effect of superfusion with LPS (1–10000 ng/ml) on the pHi of the HRASMCs is shown in Fig. 5A. In a HEPES superfusate (Fig. 5A), LPS treatment results in dose-dependent changes in the pHi, i.e. there is no change at lower doses (1–100 ng/ml), but there is a significant intracellular alkalosis for each 0.03 and 0.04 increase in the pH unit at higher doses of LPS, at 1000 ng/ml and 10000 ng/ml, respectively (p<0.05; n = 7). Note that the significant intracellular alkalosis (~ +0.04 pH unit) induced by 10000 ng/ml LPS is irreversible after washout for 30 min. The histogram in Fig. 5B shows the mean pH-induced changes for seven experiments, which is similar to that shown in Fig. 5A. The result clearly shows that the LPS-induced intracellular alkalosis is concentration-dependent between 1 and 10000 ng/ml, in HEPES-buffered superfusate.

The acute and chronic effects of LPS on the Na+-H+ exchanger activity and cellular growth

As shown in the left part of Fig. 5C, the pH recovers completely from an intracellular acidosis due to the NHE in the control. As expected, LPS causes an increase in the pH recovery slope in a concentration-dependent manner, as illustrated in the right part of Fig. 5C. At higher concentrations of 1000 and 10000 ng/ml, LPS results in a four and five fold increase in the pH activity (p<0.05; n = 6), respectively, as shown in the rightmost part of Fig. 5C. In other words, this study demonstrates that the increase in the resting pH induced by 1000 and 10000 ng/ml LPS is mainly due to its effect of increasing NHE activity. The histogram (Fig. 5D) shows the mean pH recovery slope (measured at pH = 6.89±0.09), before and after the addition of LPS, for the six experiments, which is similar to the slope shown in Fig. 5C.

In order to rule out any possible interference caused by multiple NH4Cl pre-pulses in the pH recovery shown in Fig. 5C, experiments were performed using less NH4Cl pre-pulses to determine the effect of LPS on NHE activity. As shown in the Fig. 5E, 10000 ng/ml LPS increases pH recovery, following induced alkalosis (+346±56%; n = 5) that is similar to that for 10000 ng/ml in the multiple NH4Cl experiments shown in Fig. 5C. These LPS-induced changes in pH recovery are completely reversed after washout of LPS, as shown in the far right part of Fig. 5E. This result clearly rules out any possible interference caused by multiple NH4Cl pre-pulses on the effect of LPS on NHE activity that is seen in the experiment in Fig. 5C.

The effect of LPS on NBC activity

Although the result of Western blot analysis (Fig. 4C and 4D) indicates clearly that LPS has no effect on NBCs protein expression, the possible effect of LPS on NBC activity is of interest, so experiments were performed using the superfusate of 5% CO2/HCO3- Tyrode solution in the presence of 30 μM HOE 694. Under these conditions, LPS treatment causes no change in NBC activity at higher doses (1000 and 10000 ng/ml), as shown in the middle part of Fig. 6 (p>0.05, n = 4). Note that the LPS-induced effect on NBC activity is totally different to that of LPS on NHE activity (Fig. 5C and Fig. 5E, respectively).

Figure 3. Effect of 30 μM HOE 694, Na+-free and 0.2 mM DIDS on pH recovery from induced acidosis in HRASMCs superfused with 5% CO2/HCO3- Tyrode solution. A and C: The top bar shows the buffer system used in the superfusate. The periods of application of NH4Cl and tested drugs (30 μM HOE 694, Na+-free solution, 0.2 mM DIDS and HOE 694 pulse DIDS) are shown with bars above or below the trace. The left part of traces A and C shows a typical pH recovery from an intracellular acidosis induced by a 10 min NH4Cl (20 mM) pre-pulse in 5% CO2/HCO3-. Tyrode solution (pHi = 7.4, 37°C) in HRASMCs. For details of mechanism of the pre-pulse technique, please see the Materials and Methods section. The right part of traces A and C represents experiments showing the effect of 30 μM HOE 694 (a NHE exchanger inhibitor), Na+-free solution 0.2 mM DIDS (a NBC exchanger inhibitor) and HOE 694 plus DIDS on pH recovery, respectively, in HRASMCs. B and D: Histograms, showing the pH recovery slope of acid extrusion after NH4Cl-induced intracellular acidosis averaged for several experiments similar to those shown in A and C respectively. **: p<0.01 vs. control. doi:10.1371/journal.pone.0090273.g003

Figure 4A shows the expression pattern of different SLC4 family members. It is seen that the mRNA of NBCe1 (SLC4A4), NBCe2 (SLC4A5), NBCn1 (SLC4A7) and NDCBE1 (SLC4A8) are clearly expressed, as shown in Fig. 4E. Note that the protein of NDCBE1 (SLC4A8) was not detected (Fig. 4C). This result clearly rules out any possible interference caused by multiple NH4Cl pre-pulses on the effect of LPS on NHE activity that is seen in the experiment in Fig. 5C.
Figure 4. The effect of LPS on protein expression of NHE, NBC and intracellular resting pH in HRASMCs. A: The figure shows the result of Western blot analysis for β-actin, NHE 1, 2 and 3, from the bottom to the top, respectively, before (left part) and after (right part) the 1000 ng/ml LPS treatment (n = 4). B: The histogram shows relative protein expression, as an average of 6 experiments, which is similar to that shown in A. Data is shown as the mean ± SEM (p < 0.01; n = 6). C: The figure shows the result of Western blot analysis for β-actin, SLC4A8 (NBCe1), SLC4A7 (NBCn1), SLC4A5 (NBCe2) and SLC4A4 (NBCe1), from the bottom to the top, respectively, before (left part) and after (right part) the 1000 ng/ml LPS treatment (n = 4). D: The histogram shows the protein expression, as an average of 4 experiments, which is similar to that shown in C. Data is shown as the mean ± SEM (p < 0.01; n = 4). E: Gene expression of mRNA of different members of SLC4 family: NBCe1 (SLC4A4; 336 bp), NBCe2 (SLC4A5; 650 bp and 1 kb), NBCn1 (SLC4A7; 328 bp) and NDCBE1 (SLC4A8; 243 bp) extracted from HRASMCs by RT-PCR. Actin expression was used as control (373 bp). bp denotes base pairs; M denotes marker; + denotes the presence of template; − denotes the absence of template (negative control). doi:10.1371/journal.pone.0090273.g004
The chronic effects of LPS on the Na\(^+\)-H\(^+\) exchanger activity and cellular growth

The effect of the concentration of LPS (1–10000 ng/ml) on cellular growth of cultured HRASMCs is of interest. The HRASMCs were treated with LPS for 24 hrs in a culture chamber, and then a MTT assay was used to determine the cell viability (growth). It was found that LPS increases the growth of cultured HRASMCs in a concentration-dependent manner (Fig. 7), i.e., it has no effect between 1–10 ng/ml, while cell growth increased ~2 fold or ~3 fold, respectively, at higher dosages of 100 ng/ml and 1000 ng/ml (p<0.05, n = 6). Note that the highest dosage of 10000 ng/ml LPS does increase the number of cells ~1.9 fold (p<0.05, n = 5), which is not greater than that the effect of 100 and 1000 ng/ml. These results are the first solid evidence that LPS increases cellular growth in cultured HRASMCs.

In order to verify whether the phenomenon of LPS-induced cellular growth is closely related to NHE activity, the time-dependent effect of 1000 ng/ml LPS on the NHE activity was determined, as shown in Fig. 8. The choice of a 1000 ng/ml dose is based on the result of its significant effect on NHE1 protein synthesis (Fig. 4A), NHE activity and pHi (Fig. 5). The NHE activity was observed before and after the addition of LPS (1000 ng/ml), at 6, 12, 18, 24 and 48 hrs, in the culture chamber, as shown in Fig. 8A – Fig. 8F, respectively. This study finds that NHE activity is significantly increased at 18 hr ~48 hr (Fig. 8D ~ Fig. 8F, respectively), but not significantly increased before 12 hr (Fig. 8B, Fig. 8C). The histograms in Fig. 8G show the normalized

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Figure 5. Effect of lipopolysaccharides (LPS) on resting pHi and NHE activity in HRASMCs superfused with HEPES-buffered Tyrode solution. A, C, E: The top bar shows the buffer system used in the superfusate. The periods of application of NH\(_4\)Cl and LPS (1–10000 ng/ml) are shown with bars above or below the trace. Traces A represents experiments showing the effect of different concentrations of LPS (1–10000 ng/ml) on resting pHi, in HEPES-buffered Tyrode solution in HRASMCs (pHo = 7.4, 37°C). The left part of traces C and E shows a typical pHi recovery from an intracellular acidosis induced by a 7 min NH\(_4\)Cl (20 mM) pre-pulse in HEPES-buffered solution (pHo = 7.4, 37°C) in HRASMCs. The right part of traces C and E represents experiment showing the effect of different concentrations of LPS (1–10000 ng/ml) on pHi recovery in HRASMCs. B, D: Histograms, showing the change in resting pHi and pHi recovery slope of acid extrusion after NH\(_4\)Cl-induced intracellular acidosis averaged for 7 and 6 experiments similar to those shown in A and C (measured at the range between the two dash lines of the figure), respectively. *: p<0.01 vs. control. doi:10.1371/journal.pone.0090273.g005
ward evidence that the NHE isoform is purely NHE1 (Fig. 4A) and (30 HRASMCs is also sensitive to low concentration of HOE 694 [50]. These results show that the functioning NHE in the of magnitude higher than for the other isoforms, such as NHE 2 & selectivity for cloned & expressed NHE1 that is two or more orders smooth muscle [48,49]. It is shown that HOE 694 exhibits a high ubiquitously expresses in different tissues, including heart and shows that, of the nine different members of NHE, i.e. NHE 1 highly-specific NHE-1 inhibitor [21]. Molecular biological analysis acid extruder is entirely blocked by HOE 694 (Fig. 2), which is a [16,26,47]. This conclusion is confirmed by the finding that the induced acidosis in human renal artery smooth muscle. NHE’s concentrations of LPS (1000 ng/ml and 10000 ng/ml) on pHi recovery in part of trace represents experiment showing the effect of 2 different concentrations of LPS (1000 ng/ml and 10000 ng/ml) on pH recovery in HRASMCs. doi:10.1371/journal.pone.0090273.g006

NHE activity (measured at pH$_i$ = 6.88±0.06), which is similar to that shown in Figs. 8A ~8F, respectively. Note that the maximum increase due to the 1000 ng/ml dose is observed at 24 hr (Fig. 8G).

These results provided clear pharmacological evidence that, in HEPES-buffered Tyrode solution, the underlying mechanism for LPS-induced, concentration-dependent intracellular alkalosis (Fig. 3A) in cultured HRASMCs is mainly due to its effect on NHE expression/activity (Fig. 4, Fig. 5 and Fig. 8). This is the first demonstration that the concentration-dependent LPS-induced cellular growth of culture HRASMCs (Fig. 7) is possibly due to the pH$_i$ change that is caused by an increase in NHE activity (Fig. 5 and Fig. 8).

**Discussion**

The evidence of acid extruding regulators (NHE1, NBCn1, NBCe1 and NBCe2)

Using microspectrofluorimetry, this study provides the first straightforward and convincing pharmacological evidence that NHE1 and another three HCO$_3^-$-dependent acid extruder, i.e. NBC, are functionally responsible for acid extrusion, following induced acidosis in human renal artery smooth muscle. NHE’s activity is HCO$_3^-$-independent and Na$^+$ dependent (Fig. 2) [16,26,47]. This conclusion is confirmed by the finding that the acid extruder is entirely blocked by HOE 694 (Fig. 2), which is a highly-specific NHE-1 inhibitor [21]. Molecular biological analysis shows that, of the nine different members of NHE, i.e. NHE 1~9 [22], the NHE1 protein is identified as the protein that ubiquitously expresses in different tissues, including heart and smooth muscle [40,49]. It is shown that HOE 694 exhibits a high selectivity for cloned & expressed NHE1 that is two or more orders of magnitude higher than for the other isoforms, such as NHE 2 & 3 [50]. These results show that the functioning NHE in the HRASMCs is also sensitive to low concentration of HOE 694 (30 μM) (Fig. 2). Western blot analysis also provides straightforward evidence that the NHE isoform is purely NHE1 (Fig. 4A) and lacks NHE 2 and NHE 3. Whether this data excludes a significant presence for other members of NHE (4~9) in HRASMCs is worthy of discussion. It appears that it is excluded, on the grounds that the data available for NHE 4, 5 indicates that the acid extruder is essentially relatively insensitive to HOE694, and NHE 6~9 only exists in the membrane of intracellular organelles [22]. Therefore, using pharmacological methods and a molecular probe, this study provides direct evidence that the native NHE that functions during pH$_i$-regulation in the HRASMCs is the NHE-1 isoform; not other members of the NHE proteins.

Another category extruding mechanism whose activity is HCO$_3^-$- and Na$^+$-dependent (Fig. 3A) is NBC. This is supported by another result (Fig. 3C) in this study, which shows that NBC is sensitive to DIDS, a NBC inhibitor, and insensitive to HOE 694 [16,21,28,31,47]. Relevant molecular candidates for Na$^+$-dependent bicarbonate transport include at least five members of the slc4 family, including 2 electrogenic Na$^+$, HCO$_3^-$ co-transporters (NBCe1/SLC4A4 and NBCe2/SLC4A5), 1 electroneutral Na$^+$, HCO$_3^-$ co-transporter (NBCn1/SLC4A7) and 2 Na$^+$-dependent Cl$^-$/HCO$_3^-$ exchangers (NCBE/SLC4A10 and NDCBE/ SLC4A8) [12,30,33]. Recently, both in rat and mouse smooth muscle cells, the Aalkjaer group demonstrated that the NBC is disruption of the Na$^+$, HCO$_3^-$ co-transporter NBCn1 (SLC4A7) inhibits NO- mediated vasorelaxation, smooth muscle Ca$^{2+}$-sensitivity and the development of hypertension in mice [17]. Indeed, this study demonstrates functionally that a Na$^+$ and HCO$_3^-$ dependent acid-extruding mechanism is responsible for acid extrusion in the cultured HRASMCs (Fig. 3A). Surprisingly, this study demonstrates, for the first time, that three different isoforms of NBC: NBCn1 (SLC4A7; electroneutral), NBCe1 (SLC4A4; electrogenic) and NBCe2 (SLC4A5), are detected in the protein/mRNA level (see Fig. 5B and Fig. 5E) in the cultured HRASMCs. In other words, the co-existence of 3 types NBC in this study is different to that found in mouse and rat models (c.f. Aalkjaer’s group), which is probably due to differences in species/ organs, if the specificity of the antibodies used is reliable (see Materials and Methods for details). Moreover, it is important to consider another possibility. The observed isoforms of NBC may be directly caused by the altered expression of the protein, during the culture in question, or a secondary, for instance the compensatory up- or down-regulation of other proteins, as cell culture is known to alter the expression profile [44]. Knowledge of the exact stoichiometry between HCO$_3^-$ and Na$^+$ (coupling ratio)
and the extent to which the NBC is electrogenic or electroneutral to the multiple NBC isoforms in the cultured HRASMCs will involve further study.

The potential role of inhibitors of NHE1 and NBCs in a clinic

In the HRASMCs, it is demonstrated that the activity of NHE1 and/or NBCs (n1, e1 and e2) is imperative for pH_i regulation (Fig. 2, Fig. 3 and Fig. 4). Both in rat or mouse vascular smooth cells, it is seen that NHE1 is predominantly active at lower pH_i values and that it plays a major role in acid extrusion under conditions of severe intracellular acidification [18], but NBCn1 is active at both low and near-physiological pH_i values [15,17]. Whether the percentage contributions of NHE1 and NBCs of HRASMCs are similar to that of those for rat or mouse vascular smooth cells will be a subject for further study. In order to quantify the pH_i-dependency of NHE and NBC activity, an entire set of experiments to elaborately check the intracellular total buffering power (b_tot), which comprises the intrinsic- and CO₂-related buffering power, must firstly occur [14].

According to this result, it is predicted that the alteration of the activity of NHE1 and NBC plays a vital role in maintaining many conditions.

Figure 8. Time-dependent effect of lipopolysaccharides (LPS) on NHE activity in HRASMCs. A–F: HRASMCs were incubated with different LPS (1000 ng/ml) for 0, 6, 12, 18, 24 and 48 hr, respectively. Cells were subjected to perform NH₄Cl pre-pulse method to detect the NHE activity. Top bar shows the buffer system used in the superfusate. The periods of application of NH₄Cl and tested LPS (1000 ng/ml) are shown by the bars below or above the traces. Traces A shows a typical pH_i recovery from an intracellular acidosis induced by a 10 min NH₄Cl (20 mM) pre-pulse in HEPES-buffered solution (pHe = 7.4, 37°C) in HRASMCs. Traces B–F represent experiments showing the time-dependent effect of LPS (6, 12, 18, 24 and 48 hr, respectively) on NHE activity in HRASMCs. G: Histogram, shows acid extrusion after acid loading estimated at pH_i 6.88±0.06, averaged for several experiments similar like that of A–F, respectively. **p<0.01 vs. control.
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physiological functions, such as cell differentiation, growth and apoptosis in HRASMCs, which is similar to that found for many other groups in other cell types [10,11,43,51]. It is also clear that NHE and NBCs may provide an important method for the prevention of some acute and chronic, pathological vascular illnesses in clinics, such as ischaemia-reperfusion induced damage that is caused by rapid recovery of pH [32] and ARAS-induced atherosclerotic renovascular disease [3,4,12,53,54]. This study also implies that, as well as increasing the knowledge of the basic physiological mechanism of NBCs, the development of a new and specific NBCs inhibitor is another step in the process to prevent ischaemia/reperfusion-induced cardiovascular injury.

Clinical implication of concentration- and time-dependent effects of LPS on pH, NHE and NBCs

There is a close relationship between the plasma concentration of circulating LPS and the development of multiple organ failure and death in patients with bacteriologically verified systemic meningococcal disease (SMD) [53]. A plasma LPS level of more than 700 ng/L is correlated with the development of severe septic shock (P < 0.0001), adult respiratory distress syndrome (P = 0.0035), a pathologically elevated serum creatinine level (P < 0.0001), or death, as a consequence of multiple organ failure (P = 0.0002). Initial plasma LPS levels of less than 25, 25–700, 700–10000 and greater than 10000 ng/L are also associated with a 0%, 14%, 27% and 96% risk of mortality, respectively [55]. Indeed, recently, it has been demonstrated that mRNA and protein expression of toll-like receptor 4 (TLR4) are up-regulated by LPS in human aortic smooth muscles, in a dose- (10–10000 ng/ml) and time-dependent (0–48 hr) manner [33]. In human arterial smooth muscle, LPS (10 ng/ml) has also been found to induce mRNA and protein expression of matrix metalloproteinases-9 (MMP-9) and the process depends on TLR4/NF-κB [34]. This study, determines the effect of LPS at different concentrations (1–10000 ng/ml) LPS (Figs. 4–8). This is the first evidence that LPS (1–10000 ng/ml) induces concentration-dependent, intracellular alkalosis (Fig. 5A). Fig. 5C and Fig. 8 show that the LPS increases NHE activity in both a concentration- and time-dependent manner. However, LPS has no effect on either the protein expression or activity of NBCs (n1, e1 and e2), the other main acid-extruding mechanism in HRASMCs (Fig. 4C and Fig. 6). Therefore, it is demonstrated that, in HRASMCs, the LPS-induced intracellular alkalosis is mainly due to the alternation of NHE1 activity/protein expression.

As well as from cell differentiation, growth and apoptosis are sensitive to changes in pH, [10,11]. It has been claimed that irreversible endothelial dysfunction and vascular atherosclerosis are related to a disturbance in pH [12,13]. For example, NHE1 activity has been proven to play a vital role in proliferation, both in carcinogenic and non-carcinogenic cells [56–50]. Recently, it has been found that LPS-induced vascular inflammation/occlusion and systemic organ failure are initially triggered by vascular the endothelial apoptosis that is associated with increased [Ca2+]i, which results in enhanced calpain activity and in HUVECs apoptosis, via NHE1-dependent degradation of Bcl-2, which is one of the anti-apoptotic family members [30]. Indeed, this study also shows, for the first time, that LPS increases cellular growth significantly in a concentration-dependent manner (Fig. 7). The pattern of change of LPS-induced cell growth is also closely related to those of LPS-induced changes in pH, and increases in NHE activity (Fig. 5, Fig. 7 and Fig. 8).

LPS is one of the main inflammatory mediators that exerts various atherogenic effects that involve the expression of adhesion molecules [25,60]. For example, LPS has been demonstrated to stimulate the release of INF-γ, II (interilukin) -1, II-6, II-8, TNF-α (tumor necrosis factor-α) and GM-CSF (granulocyte-macrophage colony-stimulating factor), either through direct or indirect mechanisms [25,60]. If a change in NHE1 activity also affects LPS-induced impact on cytokines, then the development of specific and potent NHE inhibitors/agonists may produce a cure for LPS induced-malfunction, such as organ failures in sepsis and septic shock, in a clinic. Therefore, determining whether these LPS-induced alternations to cytokines, peptides and co-stimulatory molecules is related to changes in pH, and NHE activity is worthy of future study.

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Author Contributions

Conceived and designed the experiments: SHL CYL YTT. Performed the experiments: SHL SJS LWC. Analyzed the data: SHL CYL YTT SJS LWC THC. Contributed reagents/materials/analysis tools: SHL CYL YTT THC CYC CST. Wrote the paper: SHL.

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