Lipoxin A₄ Stimulates Calcium-Activated Chloride Currents and Increases Airway Surface Liquid Height in Normal and Cystic Fibrosis Airway Epithelia

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Introduction

Cystic Fibrosis (CF) is a genetic disease characterised by a deficit in epithelial Cl⁻ secretion which in the lung leads to airway dehydration and a reduced Airway Surface Liquid (ASL) height. The endogenous lipoxin LXA₄ is a member of the newly identified eicosanoids playing a key role in ending the inflammatory process. Levels of LXA₄ are reported to be decreased in the airways of patients with CF. We have previously shown that in normal human bronchial epithelial cells, LXA₄ produced a rapid and transient increase in intracellular Ca²⁺. We have investigated, the effect of LXA₄ on Cl⁻ secretion and the functional consequences on ASL generation in bronchial epithelial cells obtained from CF and non-CF patient biopsies and in bronchial epithelial cell lines. We found that LXA₄ stimulated a rapid intracellular Ca²⁺ increase in all of the different CF bronchial epithelial cell tested. In non-CF and CF bronchial epithelia, LXA₄ stimulated whole-cell Cl⁻ currents which were inhibited by NPPB (calcium-activated Cl⁻ channel inhibitor), BAPTA-AM (chelator of intracellular Ca²⁺) but not by CFTRinh-172 (CFTR inhibitor). We found, using confocal imaging, that LXA₄ increased the ASL height in non-CF and in CF airway bronchial epithelia. The LXA₄ effect on ASL height was sensitive to bumetanide, an inhibitor of transepithelial Cl⁻ secretion. The LXA₄ stimulation of intracellular Ca²⁺, whole-cell Cl⁻ currents, conductances and ASL height were inhibited by Boc-2, a specific antagonist of the ALX/FPR2 receptor. Our results provide, for the first time, evidence for a novel role of LXA₄ in the stimulation of intracellular Ca²⁺ signalling leading to Ca²⁺-activated Cl⁻ secretion and enhanced ASL height in non-CF and CF bronchial epithelia.

Abstract

Cystic Fibrosis (CF) is a genetic disease characterised by a deficit in epithelial Cl⁻ secretion which in the lung leads to airway dehydration and a reduced Airway Surface Liquid (ASL) height. The endogenous lipoxin LXA₄ is a member of the newly identified eicosanoids playing a key role in ending the inflammatory process. Levels of LXA₄ are reported to be decreased in the airways of patients with CF. We have previously shown that in normal human bronchial epithelial cells, LXA₄ produced a rapid and transient increase in intracellular Ca²⁺. We have investigated, the effect of LXA₄ on Cl⁻ secretion and the functional consequences on ASL generation in bronchial epithelial cells obtained from CF and non-CF patient biopsies and in bronchial epithelial cell lines. We found that LXA₄ stimulated a rapid intracellular Ca²⁺ increase in all of the different CF bronchial epithelial cell tested. In non-CF and CF bronchial epithelia, LXA₄ stimulated whole-cell Cl⁻ currents which were inhibited by NPPB (calcium-activated Cl⁻ channel inhibitor), BAPTA-AM (chelator of intracellular Ca²⁺) but not by CFTRinh-172 (CFTR inhibitor). We found, using confocal imaging, that LXA₄ increased the ASL height in non-CF and in CF airway bronchial epithelia. The LXA₄ effect on ASL height was sensitive to bumetanide, an inhibitor of transepithelial Cl⁻ secretion. The LXA₄ stimulation of intracellular Ca²⁺, whole-cell Cl⁻ currents, conductances and ASL height were inhibited by Boc-2, a specific antagonist of the ALX/FPR2 receptor. Our results provide, for the first time, evidence for a novel role of LXA₄ in the stimulation of intracellular Ca²⁺ signalling leading to Ca²⁺-activated Cl⁻ secretion and enhanced ASL height in non-CF and CF bronchial epithelia.

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Competing Interests: The authors have declared that no competing interests exist.

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normal human bronchial epithelial cells are a biological target for LX4. The receptor for LX4 (ALX/FPR2) is expressed in the bronchial epithelial cell line 16HBE14o- and LX4 stimulates an intracellular Ca\(^2+\) mobilisation in these cells [7]. Intracellular Ca\(^2+\) is a major regulator of Cl\(^-\) transport and the stimulation of epithelial Cl\(^-\) secretion would be of major therapeutic benefit in CF to restore efficient airway clearance. We have investigated the effect of LX4 on epithelial Cl\(^-\) secretion and its functional consequences on ASL height using bronchial epithelial cells obtained from CF and non-CF patient biopsies and in a variety of bronchial epithelial cell lines commonly used as models for CF ion transport and immunological studies.

Results

**LX4 Effects on Intracellular Ca\(^2+\) in Normal and CF Bronchial Epithelial Cells**

LX4 induced a rapid increase of intracellular Ca\(^2+\) in all human airway epithelial cell types tested. These results are summarised in typical records obtained in NuLi-1 and CuFi-3 cell lines (Figure 1A) and NHBE and CFBE primary cultures (Figure 1B) and on corresponding histograms (Figure 1C). There was no significant difference in the maximum increase in calcium induced by LX4 (100 nM) between non-CF and CF bronchial epithelial cell lines (Figure 1A and 1C). However the maximum increase obtained in CF bronchial epithelial primary culture (CuFi-3) was significantly higher than in non-CF (NHBE) bronchial epithelial primary cultures (Figure 1B and 1C). The kinetics of the Ca\(^2+\) responses were markedly different between CF and non-CF cells. In the non-CF bronchial epithelial cells (NHBE, primary cultures and NuLi-1 cell line), the intracellular Ca\(^2+\) rise induced by LX4 was fast and transient with a recovery to basal values within 2 to 3 min (Figure 1C). In contrast, in the CF bronchial epithelial cells (CFBE primary cultures and CuFi cell lines), LX4 induced a slower increase in Ca\(^2+\) and a delayed (or absent) recovery toward basal values (Figure 1C). Thus the total amount of Ca\(^2+\) mobilised in the cytosol upon LX4 exposure was higher in CF cells than in non-CF bronchial epithelial cells.

In order to investigate the origin of the calcium signal induced by LX4, we tested the effect of LX4 on intracellular Ca\(^2+\) mobilization in the absence of extracellular Ca\(^2+\) in non-CF and CF airway epithelial cells. The results presented in Figure 1A show that in NuLi-1 cells bathed in a normal Ca\(^2+\)-free solution, the response to LX4 was not different from control conditions (plain line). In NuLi-1 cells, there was no significant difference in the maximum Ca\(^2+\) increase obtained after LX4 exposure with or without external Ca\(^2+\) (F340/F380 : control 8.49±0.48 and external Ca\(^2+\)-free 8.57±0.32) and in the value measured 2 min after the peak (5.22±0.43 in control condition and 3.93±0.36 in external Ca\(^2+\)-free (n = 4, p>0.1)). These results indicate that in normal airway epithelial cells, LX4 generates a calcium signal mainly due to the release of Ca\(^2+\) from intracellular stores rather than Ca\(^2+\) entry. As shown in Figure 1A, there was no difference in the maximum Ca\(^2+\) increase in CuFi-3 cells obtained after LX4 exposure with or without external Ca\(^2+\) (F340/F380: control 8.76±0.55 and external Ca\(^2+\)-free: 8.99±0.49 (n = 4, p>0.1)). However, in external Ca\(^2+\)-free medium, the calcium response to LX4 in CuFi-3 cells was more transient with a rapid recovery to basal values. The F340/F380 ratio values obtained 2 min after the peak Ca\(^2+\) response were 7.17±0.43 in control conditions and 2.29±0.46 (n = 4, p<0.05) in an external Ca\(^2+\)-free solution. Taken together, these results suggest that, in CF airway epithelia, in addition to the Ca\(^2+\) mobilisation from intracellular stores, LX4 also stimulates Ca\(^2+\) entry and this response is absent in non-CF bronchial epithelial cells (Figure 1A).

**Role of the ALX/FPR2 Receptor in the Calcium Response to LX4**

The role of the ALX /FPR2 receptor in the Ca\(^2+\) response to LX4 was investigated using the specific inhibitor, Boc-2 (Figure 2). The effect of LX4 (100 nM) on intracellular Ca\(^2+\) was completely abolished after treatment with Boc-2 (10 μM) in both NuLi-1 (n = 5) and CuFi-3 (n = 4) cells (figure 2A and 2B). However, ATP (100 μM), a known stimulator of intracellular Ca\(^2+\) mobilisation via purinergic receptor stimulation, produced a Ca\(^2+\) signal in cells treated with Boc-2 (Figure 2A). These results support the involvement of the ALX/FPR2 receptor in the Ca\(^2+\) signalling response to LX4.

**LX4 Effects on Whole-cell Currents in Non-CF and CF Bronchial Epithelial Cells**

Since intracellular Ca\(^2+\) is a regulator of Cl\(^-\) transport and we have shown that LX4 regulates Ca\(^2+\), we investigated the effect of LX4 on ion transport using whole cell patch-clamp recording techniques. The whole-cell current-voltage relationships were obtained from whole-cell patch-clamp recordings in non-CF (NHBE primary culture and NuLi-1 cell line) and CF (CFBE primary culture and CuFi-3 cell line) human bronchial epithelial cells.

As shown from the current-voltage curves (Figure 3 and 4) and in Table 1, the whole-cell currents under control conditions were outwardly rectified in non-CF and CF cell lines and in primary airway epithelial cells. The reversal potentials (Erev) obtained in the cell lines (Erev NuLi-1 = -23.8±3.2 mV, Erev CuFi-3 = -21.3±6.6 mV) and primary cultures (Erev NHBE = -22.4±3.9 mV and Erev CFBE = -22.4±1.9 mV) indicate that Cl\(^-\) (ECl = -39 mV) is the main charge carrier under these conditions (Table 1).

LX4 exposure stimulated the whole-cell currents in non-CF and CF airway epithelial cell lines and in normal and CF primary cultures, in a time-dependent and dose-dependent manner, as illustrated in Figure 3 and Figure 4, respectively.

Typical whole-cell current-voltage relationship in NHBE and CFBE primary cultures recorded over variable duration of LX4 (100 nM) exposure illustrate the time-dependence of the response (Figure 3A and 3B). The statistical significance of the time dependence of current responses have been investigated in non-CF and CF airway epithelial cell lines and primary cultures as shown in Figure 3C. The maximum stimulatory effect on the inward conductance (for outward flux of Cl\(^-\) from the cell) in non-CF airway epithelial cells (NuLi-1 and NHBE) was obtained after 10 min exposure to LX4 and declined thereafter to control levels (Figure 3C). In CF airway epithelial cells (CuFi-3 and CFBE), the increased inward conductance induced by LX4 was sustained without recovery to basal values over the 15 min period of observation (Figure 3C).

The dose dependence of the response to LX4 was investigated in the non-CF and CF cell lines. The stimulatory effect of LX4 on membrane conductance was found to be dose-dependent as illustrated on the typical whole-cell current-voltage relationship obtained from NuLi-1 and CuFi-3 cell lines upon exposure to different LX4 concentrations (figure 4A and 4B). In both cell lines, significant responses were observed at concentrations as low as 1 μM and the maximum response achieved at 10 nM LX4 (Figure 4C).
As reported in Table 1, the inward and outward conductances were significantly increased by LXA₄ (100 nM, 10 min) without any change in the reversal potential in all cell types studied including primary cultures of CF and non-CF bronchial epithelia and CF and non-CF cell lines.

Role of the ALX/FPR2 Receptor in the Whole Cell Conductance Responses to LXA₄

The inhibitory effect of the Boc-2 antagonist on calcium responses to LXA₄ indicate a role for the ALX/FPR2 receptor in transducing LXA₄ responses in airway epithelial cells. This is also true for lipoxin effects on membrane ionic currents where treatment of Nuli-1 and CuFi-3 cells with Boc-2 (10 μM) completely abolished the stimulatory effect of LXA₄ on the whole-cell current. As indicated in figure 4C, Boc-2 did not affect the basal inward conductance but significantly prevented the increase in inward conductance induced by LXA₄ (Figure 4C).

Role of Intracellular Ca²⁺ in the Whole-cell Conductance Responses to LXA₄

The role of intracellular Ca²⁺ in the LXA₄ induced whole-cell currents was evaluated using BAPTA-AM in the patch pipette as a

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Figure 1. LXA₄ effect on intracellular Ca²⁺ activity in non-CF and CF bronchial epithelial cells. (A) Typical effect of LXA₄ (100 nM) on the cytosolic Ca²⁺ (ratio F340/F380) measured in Nuli-1 and CuFi-3 cell lines in control (plain line) and in external Ca²⁺ -free conditions (dotted line). (B) Typical effect of LXA₄ (100 nM) on the cytosolic Ca²⁺ in normal and CF primary cultures of bronchial epithelial cells (NHBE and CFBE). (C) Mean values of the maximum increase in Ca²⁺ (peak) and measured 2 and 5 minutes after the peak, in Nuli-1 (n = 6), CuFi-1 (n = 6), CuFi-3 (n = 6), CuFi-4 (n = 4) cell lines and in NHBE (n = 4) and CFBE (n = 4) bronchial epithelial cells in primary culture (* p<0.05, ** p<0.01). doi:10.1371/journal.pone.0037746.g001
LXA4 Increases Cl− Secretion and ASL in CF Airways

Figure 2. Effect of Boc-2 on the intracellular Ca2+ signal induced by LXA4. (A) Representative effect of LXA4 (100 nM) and ATP (100 μM) on cytosolic Ca2+ (ratio F340/F380) in NuLi-1 cells in control conditions (upper panel) and after 24 hours of pre-treatment with Boc-2 (10 μM) a specific inhibitor of ALX/FPR2 (lower panel). (B) Mean values corresponding to the effect of Boc-2 on the Ca2+ response to LXA4 in NuLi-1 (n = 5) and in CuFi-3 (n = 4) cell lines (** p < 0.001).

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chelator of intracellular Ca2+. Under these conditions of low intracellular Ca2+, the basal whole-cell conductances remained unchanged, whereas the lipoxin-stimulated currents were absent in BAPTAM-treated cell. The use of BAPTA-AM demonstrated the absolute requirement for an increase in intracellular calcium levels to transduce the effect of LXA4 on whole-cell inward conductance in both NuLi-1 cells and in CuFi-3 cells (Figure 4C).

LXA4 Effect on Cl− Secretion

The contribution of Cl− channels to the LXA4 induced current, was tested using chloride channel inhibitors; NPPB (calcium-activated Cl− channel inhibitor) and CFTR-inh172 (CFTR Cl− channel inhibitor), and recording their effect on membrane current and conductance responses to LXA4 in CF and non-CF cells (Figure 5). Pre-treatment of CFBF primary cultured cells for 2 min with NPPB (1 μM) prior to LXA4 exposure, completely inhibited the stimulatory effect of LXA4 on whole-cell current (p > 0.5, n = 3). Furthermore, the addition of NPPB (1 μM) 10 min after exposure to LXA4 (100 nM), immediately inhibited the whole-cell currents in non-CF and CF cells. Figure 5A illustrates a typical experiment performed in NHBE primary cells showing the inhibitory effect of NPPB treatment after LXA4 stimulation of the whole cell currents. A summary of the inhibitory effects of NPPB on inward conductance in the CF and non-CF primary cultures, and CF cell line (CuFi-3) are presented in figure 5A. As an additional proof for the role of Cl− channels in driving the LXA4 induced current, we found that in the absence of an electrochemical driving force for chloride ions (equimolar Cl− replacement in the bath and patch pipette), LXA4 did not produce a change in whole-cell current in any of the cell types tested (data not shown).

The specific CFTR inhibitor, CFTR-inh172, had no effect of CFTR-inh172 on basal or LXA4-stimulated membrane current and conductance in CFBE and CuFi-3 cells (data not shown), consistent with the absence of functional CFTR. Moreover, the CFTR inhibitor did not affect the stimulation of the whole-cell current by LXA4 in NuLi-1 cells (Figure 5B). When NuLi-1 cells were pre-treated with CFTR-inh172 (5 μM), subsequent LXA4 exposure stimulated the outward and inward conductances by 2.1±0.15 fold and 2.5±0.14 fold (n = 3), respectively (Figure 5B). These results indicate that the stimulatory effect of LXA4 on whole-cell current and conductance in non-CF and CF bronchial cells is mainly due to activation of NPPB-sensitive Cl− channels and does not involve CFTR channels.

LXA4 Effects on Airway Surface Liquid Height

The consequence of LXA4 stimulation of Cl− channels and transepithelial Cl− secretion on ASL height was investigated in the cell lines (NuLi-1, CuFi-1 and CuFi-3) and in primary cultures of non-CF and CF airway epithelial cells grown on permeable supports under an air/liquid interface. The ASL measurements were carried out after a period of 24 hours exposure to apical fluid (8 μl of PBS) to allow fluid absorption/secretion and ASL height to reach a steady state. Under control conditions (without LXA4), the non-CF epithelia from NuLi-1 cell and primary cultures displayed a continuous unbroken ASL layer (not shown) whereas CF epithelia from cell lines and primary cultures presented a disrupted and thinner ASL layer (figure 6). Exposure to LXA4 (100 nM, 15 min) increased the ASL height in both non-CF and CF cell lines and primary cultures (figure 6). LXA4 treatment produced a maximum ASL height increase from 7.25±0.07 μm (n = 6) to 9.9±0.1 μm (n = 6) in NuLi-1 epithelia (p < 0.001) and from 4.6±0.20 μm (n = 6) to 11.1±0.20 μm (n = 6) in CuFi-1 epithelia (p < 0.001), and from 6.4±0.1 μm (n = 6) to 9.5±0.2 μm (n = 6) in CuFi-3 epithelia (p < 0.001) and from 4.9±0.32 μm (n = 9) to 9.8±0.10 μm (n = 9) in CFBE primary cultures (p < 0.001). The ASL height response to LXA4 obtained in CFBE primary cultures from 3 different CF patients was found to be robust and similar to the ASL response obtained in CuFi-1 differentiated cells. In all of the CF bronchial epithelia, the disrupted appearance of the ASL layer was absent following LXA4 treatment (Figure 6 B, C, D).

Role of Cl− Transport in the ASL Height Responses to LXA4

The Na/K/2Cl co-transporter inhibitor bumetanide was used to investigate the contribution of Cl− secretion to the generation of the ASL. Bumetanide (1 μM) treatment significantly decreased the basal ASL height (control 7.25±0.07 μm, bumetanide 5.9±0.1 μm (p < 0.001, n = 6)) in NuLi-1 monolayers but had no significant effect on ASL height in CuFi-1 epithelia (control 4.6±0.1 μm, bumetanide 4.82±0.10 μm (p > 0.5, n = 6)), in CuFi-3 epithelia (control 6.4±0.1 μm, bumetanide 6.3±0.3 μm (p > 0.5, n = 6)) and in CFBE (control 4.9±0.32 μm, bumetanide...
4.9±0.25 μm (p>0.5, n=8)). These data indicate that Cl− secretion contributes to the generation of the basal ASL height in the non-CF epithelium. Furthermore, bumetanide (1 μM) significantly abolished the ASL height increase induced by LXA4 in cell types tested. The ASL height measured after LXA4 exposure in the presence of bumetanide was significantly decreased in NuLi-1 (5.6±0.11 μm, n=5, p<0.05), in CuFi-1 (4.86±0.1 μm, n=6, p<0.001) and in CuFi-3 (6.2±0.1 μm, n=6, p<0.001) cell lines and in CFBE primary cultures (4.7±0.2 μm, n=9, p<0.001) compared to LXA4 alone (Figure 6). Taken together, these results demonstrate that the stimulatory effect of LXA4 on ASL height mainly involves Ca2+-dependent Cl− secretion via NPPB-sensitive channels in both CF and non-CF epithelia grown from cell lines and primary cultures.

Role of the ALX/FPR2 Receptor in the ASL Height Responses to LXA4

We tested the effect of the ALX/FPR2 receptor antagonist Boc-2 on the ASL height response to LXA4. Boc-2 significantly reduced the effect of LXA4 on ASL height in NuLi-1, CuFi-1 and CuFi-3 bronchial epithelial cell lines and CFBE primary cultures without affecting the basal ASL height (Figure 6). These data support the conclusion that the LXA4 receptor ALX/FPR2 mediates the effect of LXA4 on airway surface liquid height as well.
as on calcium mobilization and Cl\(^{-}\) secretion in bronchial epithelium.

**Discussion**

This is the first study to report a novel effect of the endogenous lipoxin LXA\(_4\) to stimulate an increase in Airway Surface Liquid height, by enhancing Ca\(^{2+}\)-activated Cl\(^{-}\) transport in bronchial epithelial cells obtained from patients with CF and non-CF patients and in airway cell lines.

In the healthy lung, the ASL forms a thin layer of fluid on the surface of the bronchial epithelium which allows cilia to beat effectively [8]. Maintenance of an optimal ASL height for ciliary beat is crucial for the efficacy of mucociliary clearance [8–12]. Bronchial epithelial ion transport regulates the ASL height, mainly by generating osmotic gradients which provide the driving force for transepithelial water movement [13]. In CF, the lack of functional CFTR leads to a reduced ASL height, resulting in an impaired mucociliary clearance that promotes chronic bacterial infection of the airways [3]. In a previous study, we reported that LXA\(_4\) stimulated an intracellular Ca\(^{2+}\) mobilization in a normal human airway epithelial cell line 16HBE14o- [7]. Here, we tested the hypothesis that LXA\(_4\) induces a Ca\(^{2+}\) signal to drive an
increased Cl\textsuperscript{−} secretion and ASL height in CF epithelium. Lipoxin A\textsubscript{4} thus stimulates a compensatory calcium-activated chloride secretory mechanism which overcomes the lack of CFTR mediated Cl\textsuperscript{−} transport and enhances airway lumen hydration. Other studies have shown that LXA\textsubscript{4} exerts biological actions on human airway epithelial cells, with a maximal effect observed at 100 nM. LXA\textsubscript{4} (100 nM) inhibited IL-8 production by airway epithelial cells [14], stimulated an intracellular Ca\textsuperscript{2+} signaling [7], increased ZO-1 expression and transepithelial electrical resistance [15], and enhanced epithelial repair after an acid injury [16]. We report here, novel effects of LXA\textsubscript{4} on ion and fluid transport in normal and CF bronchial epithelia.

Our studies show that LXA\textsubscript{4} induces an intracellular Ca\textsuperscript{2+} mobilization in normal and CF epithelia. The Ca\textsuperscript{2+} response to LXA\textsubscript{4} involves signal transduction via the ALX/FPR2 receptor, since the FPR2 receptor antagonist Boc-2 inhibited the LXA\textsubscript{4} effect. This result supports our previous study which suggested that the Ca\textsuperscript{2+} signal induced by LXA\textsubscript{4} was mediated by the ALX/FPR2 receptor since the Ca\textsuperscript{2+} response to LXA\textsubscript{4} was only obtained in the 16HBE14o- airway epithelial cell line that express the receptor whereas LXA\textsubscript{4} did not produce any Ca\textsuperscript{2+} response in the A549 cell line which does not express ALX/FPR2 [7].

Our results indicate that in non-CF airway epithelial cells, LXA\textsubscript{4} generates a rapid and transient calcium signal mainly arising from the release of Ca\textsuperscript{2+} from intracellular stores and not as a result of increased Ca\textsuperscript{2+} entry since the calcium signal was not affected by the removal of external Ca\textsuperscript{2+}. This is in agreement with our previous findings showing that the Ca\textsuperscript{2+} mobilization induced by LXA\textsubscript{4} was generated from thapsigargin sensitive stores [7]. In contrast, in CF airway epithelial cells, the duration of the Ca\textsuperscript{2+} signal induced by LXA\textsubscript{4} was greater than in normal airway epithelial cells. Although, in CF cells, the removal of external calcium did not affect the maximum peak calcium increase, the calcium response to LXA\textsubscript{4} became more transient. These results suggest that, in CF airway epithelial cells, in addition to the calcium release from intracellular stores, LXA\textsubscript{4} also stimulates calcium entry which leads to an overall larger calcium mobilisation than in normal airway epithelial cells. The observed differences reported in the
literature between the Ca^{2+} signal obtained upon agonist exposure in CF and non-CF airway epithelial cells are controversial. Some authors reported that expression of either CFTR or ΔF508CFTR in airway epithelial cells had no effect on intracellular Ca^{2+} [17]. However, our results are in accordance with the demonstration that Ca^{2+} signaling is abnormal in CF airway epithelial cells and that correction of the abnormal trafficking of ΔF508CFTR protein restored intracellular Ca^{2+} homeostasis [18]. Recent reports also indicate that intracellular Ca^{2+} signals induced by pro-inflammatory mediators are increased in CF airway epithelia compared to non-CF due to an expansion of the apical endoplasmic reticulum Ca^{2+} stores in CF airway epithelial cells [19]. This finding is
consistent with several studies showing that the nasal transepithelial electrical potential responses to agents that promote an intracellular Ca\(^{2+}\) mobilization and Ca\(^{2+}\)-dependent Cl conductance were higher in CF patients than in normal subjects [20–23].

Intracellular Ca\(^{2+}\) regulates several epithelial functions including ion transport, mucin secretion, and cilary beat frequency which constitute a primary mode of a non-specific cleansing process and lung protection. Our results indicate that the Ca\(^{2+}\) signal induced by LXA\(_4\) is coupled to an increased Cl\(^-\) secretion in CF epithelium. LXA\(_4\) stimulated the whole-cell current and conductance in non-CF and CF epithelial cells. The inhibitory effect of BAPTA-AM used as a chelator of intracellular Ca\(^{2+}\) demonstrated the essential role of Ca\(^{2+}\) in the stimulation of the whole-cell currents by LXA\(_4\). The sensitivity of basal and stimulated whole-cell currents to NPPB or Cl\(^-\) substitution, underlines the major contribution of Cl\(^-\) secretion to the generation of the whole-cell current. These results agree with our previous report indicating that LXA\(_4\) stimulated a Ca\(^{2+}\)-activated transepithelial Cl\(^-\) secretion in non-CF bronchial epithelial cells [7]. Since we found that LXA\(_4\) stimulation of the whole-cell currents was present in CF airway epithelia (in which CFTR is not functionally expressed), LXA\(_4\) most probably affects Cl\(^-\) channels other than CFTR. In addition, we found that the duration of the LXA\(_4\) effect on whole-cell currents was different between non-CF and CF cells, with a transient current increase in non-CF cells compared to CF cells where the current increase was more sustained. One explanation may be that the time course of the effect of LXA\(_4\) on the Cl\(^-\) currents is directly related to the time course of the intracellular calcium change induced by LXA\(_4\). Therefore, the greater and sustained effect of LXA\(_4\) on whole-cell currents in CF cells could be related to the long lasting Ca\(^{2+}\) signal obtained in CF airway epithelial cells. Finally, the ineffectiveness of CFTR inh-172 on the LXA\(_4\) stimulated whole-cell currents indicates that the effect of LXA\(_4\) on Cl\(^-\) secretion is not mediated by CFTR activation. This conclusion is strengthened by the observation that bumetanide reduces further the ASL height compared to Boc-2 treatment in Nuli-1 cells but not in CuFi-3 cells where functional CFTR is absent. If LXA\(_4\) had stimulated CFTR and Ca\(^{2+}\)-dependent Cl\(^-\) channels together we would expect equivalent inhibition of ASL height by Boc-2 and bumetanide in Nuli-1 cells. We have described a novel stimulatory effect of LXA\(_4\) on Cl\(^-\) secretion which produces an increased ASL height in both normal and CF epithelia. The CF bronchial epithelia generate a thinner ASL layer than non-CF airway epithelia. The CF bronchial epithelia generate a thinner ASL layer whereas the CF bronchial epithelia generate a thicker ASL layer. The CF bronchial epithelia generate a thinner ASL layer whereas the CF bronchial epithelia generate a thicker ASL layer. This finding is consistent with the diminished ASL in CF airways reported in the literature [12]. In addition, we observed that in control conditions, the non-CF epithelial monolayers showed a continuous ASL layer whereas in CuFi-1, CuFi-3 cell lines and CFBE primary cultures the liquid layer was disrupted. The gaps in the airway surface liquid layer result from localised de-hydration of the ASL. Following LXA\(_4\) exposure in CF bronchial epithelia, the ASL height significantly increased and appeared uniform. The inhibitory effect of bumetanide indicates that the effect of LXA\(_4\) on ASL height is mainly dependent on stimulation of transepithelial Cl\(^-\) transport. However, we cannot exclude the possibility that LXA\(_4\) can also exert its action to increase ASL height through the inhibition of ENaC activity which is known to be stimulated in CF airway and down-regulated by increased intracellular Ca\(^{2+}\).

An important outcome of this work is the comparable results found for LXA\(_4\) effects on ASL height in cell lines and primary CF cultures. We have demonstrated that using thin film airway liquid cell culture techniques, the Nuli and CuFi cell lines provide a robust model of airway liquid dynamics. Although ASL volume measurements have been published for the Calu-3 airway cell line [24], it is important to distinguish between ASL volume and height measurements and their meaning for effective mucociliary clearance. ASL height is the relevant physiological parameter as it determines the effective beating of cilia which must be covered to an optimal height with ASL. The volume of ASL on the other hand may change without revealing the true optimal ASL height covering the cilia but instead may reflect flooding of the airways which would also render cilia beat inefficient for mucociliary clearance. Although the use of primary airway epithelia tissue is the ideal for research in CF, the access to patient tissue samples, particularly children with CF and ‘normal’ non-CF controls as in our study, is non-trivial. The use of particular airway cell lines such as Nuli and CuFi which in our hands display normal and CF airway epithelium phenotype to secrete a thin ASL and mucus provides an additional validated cell model for CF research where human tissue samples are rare and difficult to obtain for research purposes. Taken together, our results provide evidence for a novel role of LXA\(_4\) in stimulating Ca\(^{2+}\)-activated Cl\(^-\) secretion and ASL generation in CF and non-CF airway epithelium. Thus LXA\(_4\) or its stable analogues may provide a novel therapeutic strategy to rehydrate the CF airway by modulating ion transport and airway surface liquid height via pathways which bypass defective CFTR. Our findings also indicate that the reduced levels of LXA\(_4\) observed in CF patients may be an additional contributory mechanism by which mucociliary clearance is diminished in CF airways.

**Materials and Methods**

**Cell Culture**

For the primary culture of human bronchial epithelium, the cells were obtained from bronchial brushing specimens obtained from 6 children (<6 years old) with CF and 5 non-CF controls through the SHIELD CF study (Study of Host Immunity and Early Lung Disease in CF). The children with CF were homozygous for the Phe508del mutation. Local ethics committee approval for the study was granted and written informed consent obtained. Bronchial epithelium brushings were washed and incubated for two hours at room temperature with 250 μg/ml amphotericin B in Phosphate Buffer Saline (PBS) without calcium and magnesium. After centrifugation, the pellet was collected and re-suspended in 500 μl of Bronchial Epithelium Basal Medium (BEBM, Clonetics, BioWhittaker, San Diego, USA) supplemented with 0.5 μg/ml human recombinant epidermal growth factor, 7.5 mg/ml bovine pituitary extract, 0.5 mg/ml epinephrine, 10 mg/ml transferrin, 5 mg/ml insulin, 0.1 mg/ml retinoic acid, 6.5 μg/ml triiodothyronine, and 50 mg/ml gentamicin (BD, Eremboedegem, Belgium) and 250 μg/ml amphotericin B (BD, Eremboedegem, Belgium). The explants were plated in a 24 well plate (Nunc, Roskilde, Denmark) previously coated with a fibronectin/collagen solution and incubated at 37°C in a humidified 5% CO\(_2\) atmosphere. Twenty four hours after seeding, the volume of media was adjusted to 400 μl. The cells were cultured under these conditions for six to nine days (confluence close to 70%) before splitting. Fibroblasts were removed by 1 minute treatment with trypsin EDTA (Gibco, Invitrogen, Paisley, UK). Epithelial cells referred in this paper as NHBE (non-CF bronchial epithelial cells) and CFBE (CF bronchial epithelial cells) were then trypsinised and re-suspended after centrifugation, in supplemented BEBM. The cells were seeded at 2500–4000 cells/cm\(^2\) in flasks (BD, Eremboedegem, Belgium).
LXA4 Increases Cl- Secretion and ASL in CF Airways

NuLi-1, CuFi-1, CuFi-3, and CuFi-4 cells were kindly donated by Prof. Zalner, University of Iowa, USA. The NuLi-1 cell line was derived from human airway epithelium of normal genotype, whereas CuFi-1, CuFi-3 and CuFi-4 cell lines were derived from CF patients with ΔF508/ΔF508, R553X/ΔF508, and G551D/ΔF508 genotypes respectively. The cell lines were transformed with a RT component of telomerase and human papillomavirus type 16 E6 and E7 genes [25]. Cells were initially grown to confluency in flasks using BEM with EGF, hydrocortisone, bovine pituitary extract, transferrin, bovine insulin, triiodothyronine, epinephrine, retinoic acid, penicillin-streptomycin (0.025 μg/ml), gentamicin (0.05 ng/ml), and amphotericin (25 μg/ml).

Airway epithelial cells were plated at 2×10^6 cells/cm² on Millicell hanging cell culture inserts (Millipore, Billerica, USA) for ASL height measurements. All inserts were pre-coated with collagen type VI and grown in BEGM medium until confluence was achieved. Once cell confluence was confirmed under visual inspection, the medium was switched to DMEM/F12 (Invitrogen, Auckland, New Zealand) to aid cell differentiation. This medium was supplemented with Uultror G (2%, Pall Biopera, Cergy-Saint-Christophe, France), which enhances ion transport [25], and penicillin-streptomycin (0.025 μg/ml), gentamicin (0.05 ng/ml), and amphotericin (25 μg/ml). Medium at the apical aspect was aspirated every 3–4 days until the establishment of an air-liquid interface. The basolateral culture medium was replaced every 2–3 days. After 4–6 weeks growth, the cells formed a polarised confluent monolayer with a high transepithelial electrical resistance (TER) of >700 Ω/cm².

Intracellular Calcium Imaging

Intracellular Ca²⁺ was measured by epifluorescence microscopy as previously described [26]. The human airway epithelial cells were cultured on fibronectin-collagen coated (for primary culture cells) and on collagen VI coated (for NuLi-1 and CuFi-1 cells) glass bottom dishes (WPI, Stevenage, UK) for 6 days until 70% of confluence was reached. Cells were loaded with 5 μM of the Ca²⁺-sensitive fluorescent probe fura-2-acetoxy-methyl ester (fura 2-AM, Invitrogen, Auckland, New Zealand) to aid cell differentiation. This medium was supplemented with Ultor G (2%, Pall Biopera, Cergy-Saint-Christophe, France), which enhances ion transport [25], and penicillin-streptomycin (0.025 μg/ml), gentamicin (0.05 ng/ml), and amphotericin (25 μg/ml). Medium at the apical aspect was aspirated every 3–4 days until the establishment of an air-liquid interface. The basolateral culture medium was replaced every 2–3 days. After 4–6 weeks growth, the cells formed a polarised confluent monolayer with a high transepithelial electrical resistance (TER) of >700 Ω/cm².

Whole-cell Patch-clamp Recording

Freshly isolated epithelial cells obtained from 4 non-CF patients and from 3 CF patients (genotype: ΔF508/ΔF508) and from the NuLi-1 and CuFi-3 cell lines were used for patch-clamp experiments. The CuFi-1 cells were not used for patch-clamp experiments since we could not reach a Giga ohm seal with these cells. Cells were patch-clamped at room temperature (25°C) on an inverted microscope (TE-300, Nikon, Badhoeve Dorp, Netherlands). Patch-pipettes were prepared from soda glass (Vitrex, Modulhom, Herlev, Denmark), pulled on a programmable puller (P90/PC, Sutter Instrument Company, USA). The whole-cell configuration was obtained from cell-attached mode after breaking the patch membrane by applying a brief negative pressure in the patch pipette. Whole-cell currents were amplified (Axopatch 200B, Axon instrument, CA) and digitized using a 16-bit data converter (Digidata 1322A, Axon instrument, CA) following low pass filtering at 5 KHz and sampled in real-time. Whole-cell current voltage (IV) relationships were analysed using Clampfit software (Axon instrument, CA).

The patch pipette was filled with a “high K⁺” solution at pH = 7.2, 290 mosm: 110 mM NaCl, 1.2 mM K-H₂PO₄, 3.46 mM 3mM KH₂PO₄, 5 mM EGTA, 6 mM HEPES, 2.78 mM CaCl₂, pH = 7.2 adjusted with KOH. The bathing solution had the following composition: 140 mM NaCl, 5 mM KCl, 6 mM HEPES, 2 mM CaCl₂, 1.2 mM K-H₂PO₄, 1.2 mM MgSO₄ and pH = 7.4. The Nernst potentials between the patch pipette and bath for K⁺ and for Cl⁻ were −77 mV and −39 mV, respectively. The access resistance (Ra) was determined by fitting the current transients produced by a 5mV voltage pulse with a single exponential. The measured Ra was 5.43±0.08Ω (n = 40).

For experiments performed in “low internal calcium”, cells were bathed in Kreb’s solution and the patch pipette contained 100 nM CaCl₂ with 5 mM EGTA and 10 μM BAPTA-AM. In these conditions the free Ca²⁺ has been estimated at 1 pM using the free software WEBMAXC http://www.stanford.edu/cpatton/webmaxc/webmaxcS.htm.

Airway Surface Liquid (ASL) Height Measurements

ASL height was measured using a protocol adapted from Tarran et al. [10], using live-cell confocal fluorescence microscopy. To label the ASL, 8 μl PBS containing 1 mg/ml Texas red® dextran (10 kD; Invitrogen, Auckland, New Zealand) was added to the apical surface of the well-differentiated airway epithelium. The epithelial cells were stained using Calcim-AM (5 μM, Invitrogen, Auckland, New Zealand) dissolved in medium culture for 30 minutes and introduced to the basolateral compartment of the insert. The Fluorimeter® electronic fluid Perfluorocarbon-72 (FC-72, 3M, St Paul, USA) was added to the apical compartment of the insert at a volume of 0.5 ml. Perfluorocarbon-72 is immiscible with the ASL and was used to prevent ASL evaporation on transferring the inserts from the incubator to the microscope stage and during the confocal scanning experiments. Epithelium were Z-scanned using a Zeiss LSM 510 Meta using a 40X objective. For each culture insert, 3 different microscope fields randomly chosen were XZ scanned. In each microscope field, the ASL height was measured using the Zeiss LSM Image analyser software (Carl Zeiss MicroImaging GmbH, Germany) in 9 separate regions randomly determined and then averaged. This method of quantification was carried out after blind analysis performed by multiple users. The n values referred to the number of culture inserts tested in a given condition.

Drugs

The lipoxin LXA₄ was purchased from Calbiochem. Aliquots of LXA₄ solution [100 μM] in ethanol were stored at −80°C to avoid degradation of the molecule. The peptide Boc-Phe-Leu-Phe-Leu-Phe (Boc-2) (Phoenyx pharmaceutical, Belmont, USA) was used as specific inhibitor of the ALX/FPR2 receptor [11]. For these latter experiments, cells were pre-incubated with 10⁻⁵ M Boc-2 for 24
hours at 37°C. BAPTA-AM (10 μM, Molecular probes, Leiden, Netherlands) was used to chelate intracellular Ca²⁺ (12). The 5-Nitro-2-(3-phenylpropylamino) benzoic acid (NPPB, Sigma, USA) used at 1 μM is an inhibitor of Ca²⁺-activated Cl⁻ channels (13). CFTRinh-172, an antagonist of the CFTR channel and bumetanide, an inhibitor of the NKCC1 co-transporter were supplied by Sigma (14).

Data Analysis

The intracellular Ca²⁺ variations were measured as the difference between the mean F340/F380 ratio during the 2 min prior to exposure to LXA₄ and the ratio measured at the peak of the Ca²⁺ response and 2 and 5 min after the peak. In each experiment, the mean ratio was obtained from all cells in the microscopic field. In whole-cell patch-clamp experiments, conductances were determined by linear regression of the current-voltage relationship obtained in n cells. For the ASL height measurements, three confocal image acquisitions were performed on each culture insert and nine regions of interest were analysed in each cell. Data are presented as the mean ± S.E.M. of n independent experiments. Measures of statistical significance were obtained using the Student’s t test for paired data. A p value <0.05 was deemed to be significant. All statistical operations were performed using Excel software (Microsoft).

Author Contributions

Conceived and designed the experiments: VU BJH JJV. Performed the experiments: VG GH MA. Analyzed the data: VU BJH JJV. Contributed reagents/materials/analysis tools: RWC RC PM. Wrote the paper: VU BJH.

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