LPS decreases CFTR open probability and mucociliary transport through generation of reactive oxygen species

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ABSTRACT

Lipopolysaccharide (LPS) serves as the interface between gram-negative bacteria (GNB) and the innate immune response in respiratory epithelial cells (REC). Herein, we describe a novel biological role of LPS that permits GNB to persist in the respiratory tract through inducing CFTR and mucociliary dysfunction. LPS reduced cystic fibrosis transmembrane conductance regulator (CFTR)-mediated short-circuit current in mammalian REC in Ussing chambers and nearly abrogated CFTR single channel activity (defined as forskolin-activated Cl currents) in patch clamp studies, effects of which were blocked with toll-like receptor (TLR)-4 inhibitor. Unitary conductance and single-channel amplitude of CFTR were unaffected, but open probability and number of active channels were markedly decreased. LPS increased cytoplasmic and mitochondrial reactive oxygen species resulting in CFTR carbonylation. All effects of exposure were eliminated when reduced glutathione was added in the medium along with LPS. Functional microanatomy parameters, including mucociliary transport, in human sinonasal epithelial cells in vitro were also decreased, but restored with co-incubation with glutathione or TLR-4 inhibitor. In vivo measurements, following application of LPS in the nasal cavities showed significant decreases in transepithelial Cl secretion as measured by nasal potential difference (NPD) – an effect that was nullified with glutathione and TLR-4 inhibitor. These data provide definitive evidence that LPS-generated reactive intermediates downregulate CFTR function in vitro and in vivo which results in cystic fibrosis-type disease. Findings have implications for therapeutic approaches intent on stimulating Cl secretion and/or reducing oxidative stress to decrease the sequelae of GNB airway colonization and infection.

1. Introduction

Chronic rhinosinusitis (CRS) affects 1/8 of the adults in the US population and is responsible for annual costs (direct and indirect) of $22 billion, comprising 5% of annual healthcare expenditures in the U.S [1,2]. Patients with CRS have an abysmal quality of life and have worse scores on validated questionnaires for pain and social functioning than those afflicted with congestive heart failure, angina, or chronic obstructive pulmonary disease [3]. CRS patients also experience co-morbid depression at twice the rate of the general public [4] and experience poor sleep and severe fatigue that improves with appropriate intervention [5,6].

Patients with cystic fibrosis (CF) have a prevalence of CRS...
Infection of the sinuses with gram-negative bacteria (GNB) tends to contribute to acquired defects in the mucociliary apparatus [14]. Additionally, MCT has been shown to be decreased in non-CF CRS and indicating that a slight decrease in CFTR expression confers increased likelihood of a CRS disease phenotype [12, 13].

While the genetic mutations in individuals afflicted with CF contribute to the overlying phenotypic expression of the disease, there is now long-standing evidence that wild type (WT) CFTR processing, endocytic recycling, and function can also be markedly inhibited by various environmental insults, including high altitude/hypoxemia, cigarette smoke exposure, inflammation, and infectious agents that can contribute to acquired defects in the mucociliary apparatus [14–17]. Additionally, MCT has been shown to be decreased in non-CF CRS and restored once the sinuses are cleared of infection and inflammation [18, 19]. Infection of the sinuses with gram-negative bacteria (GNB) tends to be particularly intense and recalcitrant to treatment [20,21]. Pseudomonas aeruginosa is a GNB that commonly colonizes the sinuses and airways of patients with CF and non-CF CRS [11,22–30]. Because non-CF CRS is accompanied by phenotypic resemblances and has similar GNB to CF CRS, it is prudent to consider acquired defects in CFTR as a propagating disease factor for this population.

Respiratory epithelial cells (REC) lining the mammalian airways are the primary interface between pathogens and the host; and the sinus and nasal surfaces are a crucial site for innate immune responses [31,32]. Once P. aeruginosa enters the airways, efficient clearance of the bacteria is dependent upon the recognition of the pathogen, which mounts intracellular signaling pathways responsible for triggering an innate response for host defense [33]. P. aeruginosa and other GNB exhibit lipopolysaccharide (LPS) in the cell wall, an intensely pro-inflammatory molecule recognized by the pattern recognition receptor toll-like receptor-4 (TLR4). LPS has several important functions for GNB including aiding as a permeability barrier between the cell and the environment, preserving structural stability of the membrane, and functioning as a protective barrier against foreign particles (antimicrobial peptides, salts, enzymes, drugs, and toxic heavy metals) [34,35]. LPS is critical to the disease capabilities of pathogens and is released from the outer membrane during infection. LPS is also known to generate reactive oxygen species (ROS) and causes a rapid and dynamic translocation of nuclear transcription factor NF-kappa B [36,37]. Given the universal presence of LPS on GNB and the predominance of Pseudomonas in CF disease airway colonization and infection, we hypothesized that LPS-induced ROS generation damages CFTR by oxidative modifications resulting in an acquired CFTR and mucociliary dysfunction dysfunction previously observed in non-CF CRS patients. Further knowledge regarding the interaction of LPS with respiratory epithelium, which serves as the first line of host airway defense in the upper and lower airway, is critical to develop treatment strategies targeting the impacts of early GNB colonization and infection. The objectives of this study are to identify the mechanisms by which LPS contributes to colonization and persistence in the respiratory tract through the induction of acquired defects in the mucociliary apparatus.

2. Methods

2.1. Cell culture

2.1.1. Primary cell culture

Investigational Animal Care and Use Committee and Institutional Review Board approval was obtained from the University of Alabama at Birmingham prior to the initiation of the study. Animal tissue source for REC culture was procured from the nasal septa of C57BL/6 mice and New Zealand white rabbits. Normal human sinonasal mucosa to grow human sinonasal epithelial cultures (HSNE) was obtained intraoperatively from patients undergoing endoscopic surgery for pituitary tumors, facial trauma, benign sinonasal tumors, or lacrimal obstruction. All subjects provided informed consent and were screened for 32 mutations of CFTR, including the 23 core mutations suggested for population-based CF carrier screening by the American Congress of Obstetricians and Gynecologists and the American College of Medical Genetics (Q-ANALY GENE CFTR COM VAR, Quest Diagnostics) [38]. Patients harboring a mutation were excluded from the study. RECs were dissociated from the tissues and grown on semipermeable support membranes (Costar® Transwell® clear 24-well plate inserts, 0.4-μm pore; Corning Life Sciences, Lowell, MA, USA) submerged in culture medium as previously described [14,39–44]. The media was removed from the monolayers on day 4 after reaching confluence, and the cells fed via the basal chamber. Differentiation and ciliogenesis occurred in all mammalian REC cultures within 10–21 days. Monolayers were utilized for analysis when fully differentiated with widespread ciliogenesis and transepithelial resistances >300 Ω·cm².

2.2. Cystic fibrosis bronchoepithelial (CFBE) cell culture

CFBE41o-cells expressing recombinant human CFTR (CFBE-WT, Accession GSM2985447) or flag-tagged recombinant CFTR (DS19 CFBE-mCMV-CFTR_WT_901FLAG (UAB-corrected sequence) clone # 7) were cultured as previously described (courtesy of John Kappes, PhD) [45]. Cryopreserved cells were warmed, resuspended in Minimal Essential Media-10% Fetal Bovine Serum, and plated in a flasks. Once 80% confluent, the cells were dissociated using PETF™ (Athena Enzyme Systems, Baltimore, MD) and seeded on filters (~cell number) as described above. For patch-clamp analysis, cells were seeded on glass coverslips and grown to 30–40% confluency.

2.3. Transepithelial short-circuit current measurements

All chemicals were purchased from Sigma-Aldrich (St. Louis, Mo.)

Abbreviations used:

| Abbreviation | Description |
|--------------|-------------|
| ASL          | airway surface liquid |
| CRS          | chronic rhinosinusitis |
| CBF          | ciliary beat frequency |
| CF           | cystic fibrosis |
| CFBE         | cystic fibrosis bronchoepithelia |
| CFTR         | cystic fibrosis transmembrane conductance regulator |
| GSH          | glutathione sulfhydryl |
| GNB          | gram-negative bacteria |
| HSNE         | human sinonasal epithelia |
| LPS          | lipopolysaccharide |
| μOCT         | micro-optical coherence tomography |
| MCT          | mucociliary transport |
| MNSE         | murine nasal septal epithelia |
| PCL          | pericilliary liquid |
| PBS          | phosphate buffered saline |
| ROS          | reactive oxygen species |
| REC          | respiratory epithelial cells |
| Isc          | short-circuit current |
| TLR4         | toll-like receptor-4 |
| WT           | wild type |
until otherwise stated. Filters were incubated with *P. aeruginosa* LPS (Sigma-Aldrich, St. Louis, Mo.) dissolved in phosphate buffered saline (PBS) or vehicle control solutions. Experiments were performed with concentrations of LPS ranging from 0.1 to 100 μg/ml for dose-response testing with the majority of studies completed using an optimal concentration of 10 μg/ml based on effects in the Ussing chamber. Transwell inserts were mounted in Ussing chambers and bathed with solutions that contained (in mM) 120 NaCl, 25 NaHCO₃, 3.3 K₂HPO₄, 0.8 K₃HPO₄, 1.2 MgCl₂, 1.2 CaCl₂, and 10 glucose at a pH 7.3-7.4 [46]. Bath solutions were stirred vigorously and gassed with a mixture of 95% O₂-5% CO₂ at 37 °C. Short-circuit current (Isc) measurements were obtained by using an epithelial voltage clamp (EM-CSYS-8; Physiologic Instruments, San Diego, CA). All experiments were conducted with low Cl⁻ (6 mM) in the mucosal bath (NaCl replaced with 120 mM Na gluconate). Isc was measured at 1 data point every 10 s, with the convention that positive deflection represents the net movement of anion in the serosal to mucosal direction. Transepithelial electrical resistance (TEER) was calculated according to Ohm’s law, which is a summated value from both transcellular and paracellular resistances. The electrical impedance originating from the transcellular pathway represents the stability of the apical and basolateral plasma membranes, whereas the paracellular resistance is created when adequate cell-substrate or cell-cell contacts are formed in the RECs when cultured at an air-liquid interface. Amiloride (100 μM) (Spectrum chemical MGF corp, New Brunswick, NJ) was routinely added apically in order to abolish Na⁺ absorption. Forskolin (20 μM) (Sigma-Aldrich, St. Louis, Mo.) was then administered to all bathing solutions to stimulate cAMP-mediated Cl⁻ secretion. Cl⁻ secretory currents were identified on the basis of activation by forskolin and by sensitivity to the specific CFTR inhibitor INH-172 (10 μM) (Sigma-Aldrich, St. Louis, Mo.). Effect on the basolateral Na⁺/K⁺ ATPase pumps was evaluated in apical permeability studies utilizing 300 μM amphotericin B (Sigma-Aldrich, St. Louis, Mo.) applied to the apical membrane and ouabain (1 mM) (Sigma-Aldrich, St. Louis, Mo.) applied to the basal membrane. All drugs were added as a small volume of concentrated stock solution. Time-dependent reversibility studies were performed after washing the apical surface 3x with PBS following incubation. Blockade of TLR4 receptors was accomplished with TAK-242 (1 μM, Sigma-Aldrich, St. Louis, Mo.), while glutathione sulf-hydryl reduced ethyl ester (GSH) (1 mM, Sigma-Aldrich, St. Louis, Mo.) was used to block ROS.

### 2.4. Patch clamp analysis

CFBE cells expressing WT CFTR cells on coverslips were transferred to an experimental chamber mounted on a microscope stage (Olympus). CFTR currents were recorded in whole-cell mode of the patch-clamp technique using an Axopatch 200b amplifier interfaced with a computer through DIGITA 1440A, as previously described [47]. Recording pipettes were constructed from borosilicate glass capillaries (Warner Instruments, Hamden CT) using a Narishige PP83 microelectrode puller and fire-polished with a PP90 microforge (Narishige, Tokyo, Japan). Cells were perfused with an external solution of the following ionic composition (in mM): 145 CsCl, 2 MgCl₂, 2 CaCl₂, 5.5 Glucose, 10 HEPES pH 7.4 (1 N NaOH). The pipette resistance used for whole-cell recording ranged from 3 to 5 GΩ when filled with the following intrapipette solution (in mM): 135 CsCl, 10 KCl, 2 MgCl₂, 0.1 EGTA, 5.5 Glucose, 10 HEPES, pH 7.2 (1 N KOH). All experiments were performed at room temperature. Cells were continuously perfused with the external solution before and during the recording whole-cell configuration. The rate of perfusion was adjusted to 1 ml/min where the volume of the chamber was 0.5 ml, and the solution in the chamber was changed twice per minute. Conventional patch-clamp techniques were also adapted to record single-channel currents by cell-attached configuration. The pipettes were partially filled with a solution containing (in mmol/l) 135 CsCl, 10 KCl, 2 MgCl₂, 0.1 EGTA, 5.5 Glucose, 10 HEPES, pH 7.2 (1 N KOH) and had tip resistances of 6–8 MΩ. To obtain seals, bath solutions contained (in mmol/l) 140 NaCl, 4.0 KCl, 1.8 CaCl₂, 1.0 MgCl₂, 10 glucose, 10 HEPES, pH 7.4. After seal formation, cells were perfused with potassium solution (150 mM) of the following composition: 150 mM KCl, 2 mM MgCl₂, 10 mM HEPES, pH 7.4 (1 N KOH), 5 mM Glucose. The osmolality of the solution was adjusted to 300 mosmolos (w/w) mannitol). This solution induces cell membrane potential depolarization to zero mV allowing easy control of the patch potential which will be Vpatch = -Vpipette, since VM = 0 mV [48-50]. Cells were incubated with LPS (10ug/ml) for 4 h prior to patch formation. PBS vehicle controls were included in all studies. Single-channel recordings were analyzed using pClamp 10 software (Axon). Tracings were filtered post-acquisition at 500 Hz. Current was normalized by the capacity of the membrane (pA/μF).

### 2.5. Protein analysis

#### 2.5.1. Western blotting

Cells were washed with ice-cold PBS 2x, then lysed with ice-cold RIPA buffer (50 mM Tris-HCl, 1% Nonidet P-40, 0.5% Sodium deoxycholate, 0.1% SDS, 150 mM NaCl, and Complete Protease Inhibitor (Roche Applied Science, Indianapolis, In,)). Cell lysates were sonicated x3 for 10 s each, then spun for 30 min at 4 °C to collect the supernatant. Thirty μl of the supernatant was mixed with 4X Laemmli sample buffer +5% beta-mercaptoethanol and incubated at 37 °C for 30 min. Proteins were separated by SDS-PAGE (4–12% NuPAGE gels), transferred to polyvinylidene fluoride membranes (Bio-Rad, Hercules, CA, USA). CFTR was detected using MM13-4 mouse monoclonal antibody (Nterminal 1:500, Sigma Aldrich, St. Louis, Mo. Catalog #: 05-581), and anti-mouse IRDye labeled secondary antibodies and imaged using LI-COR system. TLR4 was detected with anti-TLR4 (Accession number O00206, 1:500, Novus Biologicals, Centennial, Co, Catalog# NB100-56579) and anti-rabbit IRDye labeled secondary antibodies and imaged using LI-COR system (Lincoln, Ne.). Densitometry [51] was performed with Image Studio Lite (Li-Cor, Lincoln, Ne.).

#### 2.5.2. Biotinylation of surface CFTR

The expression of cell surface CFTR was measured after biotin-labeling using EZ-link® Biotin-LC-hydrazide (Thermo Fisher Scientific, Waltham, Ma.) [52]. CFBE cells were washed 3 times with pre-cold PBS then incubated with 5 mM Sodium Periodate which dissolved in PBS for 30min at 4 °C. Then cells were washed 2 times with 100 mM Sodium Acetate (Ph 5.5) following by one PBS wash. 2 mM biotin-LC-hydrazide dissolved in Sodium Acetate solution was added to the cells for a 30min incubation at 4 °C before cells were lysed in RIPA buffer. Cell lysates were spun down at 4 °C for 30 min and CFTR in the supernatant was then immunoprecipitated with 24-1 anti-CFTR C-terminal antibody (mouse monoclonal, Accession number P13569, Sanofi Genzyme, Cambridge, MA). Hybidiome was purchased in 1998 and antibody produced/purified at the AAB hybidioma facility). Proteins were transferred to polyvinylidene fluoride membranes. Densitometry was performed with Image Studio Lite (Li-Cor, Lincoln, NE, USA).

#### 2.5.3. Measurement of carbonyl adducts in purified CFTR protein

WT CFBE cells with flag tagged WT CFTR grown on 10 cm Petri dishes at 90% confluence were treated with LPS (10 μg/ml), PBS, PBS + GSH(1 mM), or LPS + TAK242(1 μM) for 4 h, then washed 3X with cold PBS and treated with lysis buffer (50 mM Tris HCl, pH 7.4, 150 mM NaCl, 1 mM EDTA, and 1% Triton X-100). For a reducing agent, 2-mercaptoethanol was added to the samples to attain a final concentration of 0.74 M. Cell lysis supernatants were collected and incubated with anti-flag M2 gel (Sigma-Aldrich, St. Louis, Mo.Catalog#: A2220)at 4 °C overnight under gentle rotation. Flag-tagged CFTR protein was eluted by 3X Flag Peptide (100 μg/ml, Sigma-Aldrich, St. Louis, Mo. Catalog# F4799) for 2 h at 4 °C according to the manufacturers protocol. The presence of protein carbonyl groups was assessed using the Oxyblot protein oxidation detection kit (Sigma-Aldrich, St. Louis, Mo. Catalog#
S7150), according to the manufacturer’s protocol. Briefly, the carbonyl groups in the protein side chains were derivatized to 2,4-dinitrophenyl-hydrazone by reacting with 2,4-dinitrophenylhydrazine. Three μg of protein was used for each sample, and 2,4-dinitrophenol-derivatized protein samples were separated by polyacrylamide gel electrophoresis, as described previously [53]. Polyvinylidene fluoride membranes were incubated for 1 h in stock primary antibody (1:150 in 1% PBS/TBST buffer), and after washing, for 1 h in stock secondary antibody (1:300 in % PBS/TBST buffer). Membranes were washed 3 × in TBST and visualized, as described previously [54,55]. The abundance of protein carbonylation was assessed by densitometry of each lane.

2.6. Reactive oxygen species

Measurement of ROS was performed as previously described [56]. For time-dependent studies, CFBE cells were seeded in 96-well plates at a density of 1x10⁴ and treated with 10 μg/mL LPS up to 24 h. Time-dependent ROS production was measured from 15 min to 24 h according to the duration of LPS incubation. Cells were incubated with 25 μM carboxy-H2 dichloro-dihydrofluorescein (DCF) diacetate (Image-iT™ LIVE Green ROS Detection Kit, Thermo Fisher, Waltham, MA. Catalog# I-36007) for 30 min at 37 °C, then washed with warm HBSS 3 times and lysed with 100 μl Passive lysis buffer (Promega, Madison, WI. Catalog#E1941). ROS production was measured with cell lysis by fluorescence detection of Carboxy-DCF formation in a microplate reader excitation and emission filters centered at 485 and 525 nm. To evaluate ROS generation by confocal microscopy at the peak duration, CFBE cells were grown on glass-bottom petri dishes and treated with 10 μg/mL LPS or PBS control for 4 h. After a gentle wash with warm Hank’s buffered saline, cells were labeled with 25 μM carboxy-H2 DCF diacetate (Image-iT™ LIVE Green ROS Detection Kit, Thermo Fisher, Waltham, MA. Catalog# I-36007) for 30 min or incubated with 5 μM MitoSOX™ reagent (MitoSOX™ Red Mitochondrial Superoxide Indicator, Thermo Fisher, Waltham, MA. Catalog# M36008) for 10 min at 37 °C. Hoechst 33342 was added during the last 5 min for nuclear staining. Dyes were removed and cells were washed 3x before imaging 10x and 20x dry objectives in 1×PBSoxic staining. Dyes were removed and cells were washed 3x before imaging and 20ul 10 mg/ml Texas Red® (Invitrogen, Carlsbad, CA. Catalog#C2102) was administrated to the basolateral medium 3 h before imaging and 20ul 10 mg/ml Texas Red® (Invitrogen, Carlsbad, CA. Catalog#D-1863) in FC-70 (Flourinert FC-70, Thermo Fisher, Waltham, MA. Catalog#NC 9062226) to the apical side 30 min before imaging [62]. Imaging was performed with a Zeiss LSM 710 Confocal Microscope using Zeiss Plan Apo 20x .8 na objective in 1-μm steps at room temperature. Values for each monolayer were derived from three regions of interest. ASL depth was measured in the orthogonal (head on X-Z) image view [48,49].

2.8. Nasal potential difference assay

LPS (10 μg/ml) +/- GSH (1 mM, Sigma-Aldrich, St. Louis, Mo.) or TAK242 (1 μM, Thermo Fisher, Waltham, MA.), and PBS vehicle control solution was mixed in a thermal transition gel (PureRegen gel, Jackson- sonville, Fla.) and instilled intranasally in C57BL/6 mice for 4 h (n = 6 per condition). A modified three-step protocol was used, as described previously [63–65]. First, nasal cavities of anesthetized C57/BL/6 mice were perfused with Ringer’s solution containing 148 mM NaCl, 4 mM KCl, 1.2 mM MgCl₂, 2.25 mM CaCl₂, 2.4 mM K₂HPO₄, 0.4 mM KH₂PO₄ (pH 7.4). Second, an identical solution with amiloride 100 μM to inhibit Na⁺ absorptive pathways was perfused. Finally, a zero-CI-containing solution (NaCl in solution #1 replaced by 148 mM Na gluconate, 2.25 mM Ca gluconate, 4 mM K gluconate; pH 7.4) was perfused + forskolin 20 μM to gauge CI⁻ secretion across the nasal epithelium. The activity was measured from the stable baseline to the highest point of hyperpolarization. All traces were interpreted in a blinded fashion.

2.9. Lactate dehydrogenase (LDH) assay

Cellular toxicity was evaluated as previously described [25,66]. Relative cell viability in the presence of LPS reflects concentrations of released LDH enzyme originating from damaged MNSEs. Released LDH at 4 h after incubation was measured with a coupled enzymatic reaction that converted a tetrazolium salt (iodonitrotetrazolium) into the red color formazan (Sigma-Aldrich, St. Louis, Mo. Catalog#Tox-7). Quantification of LDH was determined from the concentration of the converted formazan within the 20 min of incubation. The resulting formazan absorbs maximally at 492 nm and can be measured quantitatively at 490 nm. Total LDH in each sample was expressed in nanograms per milliliter (ng/mL).

2.10. CFTR gene expression

Total RNA was isolated from 4-h LPS or PBS control-treated filters with RNeasy mini kit (Qiagen, Valencia, Ca) according to the manufacturer’s instructions [67]. To prevent possible DNA contamination, samples were pretreated with RNase-free DNase (Qiagen) and column purified. Sequences used for murine CFTR and 18S rRNA were purchased from Assays on Demand (Thermo Fisher, Waltham, MA.) with assay ID for murine CFTR, Mm00445197_m1. A one-step PCR protocol was used to quantify CFTR mRNA transcripts by an Prism 7500 sequence detection system according to the manufacturer’s instructions. TaqMan
OneStep PCR Master Mix Reagents Kit was used for reverse transcription and PCR. All experiments were performed in triplicates.

2.11. cAMP ELISA

Cellular cAMP was measured using an ELISA-based detection kit (Cayman Chemicals, Ann Arbor, MI) as previously described [68,69]. MNSEs were incubated with LPS or PBS control for 4 h on the apical side. Forskolin (20 μM)-stimulated cAMP levels were also measured after 15 min incubation. Apical fluid was removed, and cells were washed 3 times with ice-cold PBS. Ice-cold ethanol was used to extract cellular cAMP. The supernatants were vacuum dried, and the cAMP levels were quantified per the manufacturer’s directions.

2.12. Statistical analysis

Descriptive statistics were computed and numbers compared using Student’s t-test (two-sided) or analysis of variance followed by Tukey-Kramer multiple comparison test as appropriate. All tests were performed using GraphPad Prism (GraphPad Software, Inc., La Jolla, CA). All error bars designate standard deviation, with significance set at p≤0.05.

3. Results

3.1. LPS decreases CFTR-dependent ion transport in mammalian REC

Using exposures to physiologically relevant concentrations based on GNB infection in vivo, [70–74] LPS derived from P. aeruginosa (Sigma Aldrich, St. Louis, Mo.) was administered to the apical surfaces of primary MNSE cultures in dose and time-dependent fashion to test the effects of early exposure on forskolin-stimulated transepithelial Cl– transport. A single incubation with LPS added to the apical membrane caused a significant and dose-dependent reduction in CFTR-dependent ion transport (Fig. 1A–C) with peak effects at 4 h and a concentration of 10 μg/ml (used for all subsequent studies). There was a significant decrease in amiloride-sensitive Isc (in μA/cm²) (LPS, 2.8±0.8 vs. Control, 3.5±1.1, p≤0.05, n≥20) which was consistent with previous investigations [75]. Forskolin-stimulated Isc (LPS, 20.6±3.2 vs. Control, 26.7±2.5, p≤0.001, n≥20) and inhibition with CFTRinh-172 (LPS, 16.0±4.3 vs. Control, 19.9±6.5, p≤0.05, n≥20) were both significantly reduced, indicating an LPS-induced acquired deficiency in CFTR-mediated Cl– secretion (Fig. 1D). The impact of LPS exposure was conserved across the tested mammalian REC’s including CFBE-WT cells (LPS, 112.1±18.2 vs. Control, 133.5±20.9, p≤0.01, n=13), HSNE (LPS, 32.2±4.1 vs. Control, 38.3±6.2, p≤0.01, n=12), and rabbit nasal epithelial cells (LPS, 29.3±7.6 vs. Control, 35.9±5.4, p≤0.01, n≥12) (Fig. 1E) in primary culture. Co-incubation with the TLR4 inhibitor TAK242 eliminated the effect of LPS on Isc in MNSE confirming CFTR-mediated Cl– secretion reduction is dependent on activity of this pattern recognition receptor (LPS, 11.5±2.1 vs. LPS + TAK-242, 15.0±2.2, p≤0.05, n≥6) (Fig. 1F). A small but significant effect on the basolateral Na+/K+/ATPase was noted with apical permeabilization and ouabain treatment (Supplementary Figure 1), indicating that LPS added to the apical membrane also inhibited the basolateral Na+/K+ ATPase. LPS also did not induce cellular toxicity since there was no increase in

Fig. 1. (A–D) LPS incubation in MNSE induces acquired CFTR dysfunction. (A) Representative Ussing chamber current tracings of MNSE treated with LPS (10 μg/ml) or PBS control for 4 h. Sequential addition of amiloride (100 μM), forskolin (20 μM), and CFTRinh-172 (10 μM) are shown. (B) Change in forskolin-dependent Isc of MNSE treated with increasing concentrations of LPS or vehicle control for 4 h reveals decreased Cl– secretion with a plateau at 10 μg/ml, n≥13. (C) Time-dependent changes in forskolin-induced ΔIsc following incubation of LPS (10 μg/ml) demonstrates decreased Cl– secretion with longer incubation and a plateau at 4 h, n≥10. (D) Summary of Isc measurements at optimal time (4 h) and dosing (10 μg/ml). All values are significantly different from each other (amiloride, p≤0.05; forskolin, p≤0.001; CFTRinh-172, p≤0.05, n≥20). (E) LPS-induced acquired CFTR dysfunction was consistent across mammalian RECs. CFBE-WT, human sinonasal epithelial cells, and rabbit nasal septal epithelial cells were all significantly decreased compared to PBS vehicle control by approximately 20% of CFTR-mediated Isc (p≤0.01, n≥12). (F) Co-incubation with the TLR4 inhibitor TAK242 (1 μM) eliminated LPS-dependent inhibition of CFTR-mediated Isc.
LDH release or reduced TEER at this dose and concentration (Supplementary Fig. 2A and 2B) and normal levels of forskolin-stimulated Isc were restored by 3 h following removal of LPS (Supplementary Figure 3).

3.2. LPS abrogates CFTR currents by reducing open channel probability

CFBE-WT cells were subjected to whole-cell patch-clamp analysis to characterize impact of LPS on CFTR function at the cellular level (Fig. 2). LPS incubation resulted in near-complete inhibition of forskolin–induced currents (in pA/pF) (LPS + Forskolin, 0.34 ± 0.11 vs. Control + Forskolin, 79.06 ± 21.7, p<0.001, n = 9). Similar to Ussing chamber experiments, co-incubation with the TLR4 inhibitor TAK-242 eliminated the effect of LPS, indicating that activation of TLR4-mediated pathways is necessary for this response (Tak-242, 92.57 ± 13.1, p<0.001, n = 9). We observed more substantial inward (nearly complete) than outward (~50% at -100 mV) current inhibition. Single-channel patch-clamp analysis was performed in cell-attached mode with a holding potential of -100 mV (Fig. 3). Single-channel conductances were approximately 6.5 pS with a single event amplitude of 6.5 pA consistent with what was previously reported for CFTR[47,69]. The activity was measured after cells were incubated with or without LPS following the addition of 10 μM forskolin and 100 μM IBMX. While unitary conductances and single-channel amplitudes were unaffected by LPS, the number of active channels and their open probability (NPo) in the membrane were markedly decreased (LPS, 0.4 ± 0.1 vs. Control, 1.5 ± 1.2, p<0.01, n = 10). Taken together with the whole-cell recordings indicating uneven rectification, the decrease in channel open probability confirms that LPS treatment leads to changes in CFTR function.

3.3. Total and cell surface CFTR levels are unaffected by LPS exposure

Western blot analysis in MNSE (using the rat monoclonal anti-CFTR antibody, 3G11) and CFBE-WT (mouse monoclonal anti-CFTR antibody, MM13-4) revealed no changes in total CFTR levels (MNSE, LPS 1.16 ± 0.18 vs. control, 1.0 ± 0.34, n = 5 and CFBE-WT, LPS 1.07 ± 0.08 vs. 1.0±/-0.05, n = 6) (Fig. 4A and B). TLR4 protein levels were also unaffected by LPS incubation (CFBE-WT, 1.03 ± 0.10 vs. 1.0 ± 0.07, n = 6). CFBE cells were used to measure surface CFTR and TLR4 due to greater protein expression and availability of antibodies for both immunoprecipitation and Western blot. CFTR has been reported previously to extract and endocytose LPS[36]. Since total CFTR levels did not change, we wanted to test if there was change in the distribution of CFTR following LPS treatment. To evaluate whether reduced Isc resulted from diminished cell surface CFTR (sCFTR), cell surface proteins were biotinylated[52]. Following lysis, total CFTR was immunoprecipitated (mouse monoclonal anti-CFTR C-terminal antibody, 24-1), subjected to SDS-PAGE and Western blot. Biotinylated CFTR was detected with SA-HRP. No significant changes in cell surface CFTR levels were observed (Fig. 4C). To test the sensitivity of the cell surface biotinylation assay to detect surface CFTR, we applied cAMP to enhance surface CFTR levels[52]. Consistent with previous findings, we measured 10–15% increase in sCFTR levels in response to forskolin (20 μM) in both LPS (LPS, 1.06 ± 0.10 vs. +cAMP 1.33 ± 0.03, p<0.001, n = 6) and control samples (PBS, 1.0 ± 0.06 vs. +cAMP 1.25 ± 0.04, p<0.001, n = 6).

These results confirm that reduced transepithelial currents (Isc) at this dose and incubation are not the result of decreased CFTR protein expression at the apical surface. Furthermore, CFTR mRNA levels at 4-h LPS incubation were not significantly different (Supplementary Figure 4).
Fig. 3. (A&B) Single channel patch-clamp with 4-h LPS (10 μg/ml) incubated CFBE-WT cells shows the CFTR open probability and number of active channels were markedly decreased. (C) LPS significantly reduced the number of active channels and their open probability shown here as NPo ($p \leq 0.01$, $n = 10$).

Fig. 4. LPS treatment does not alter CFTR levels. (A) CFTR Western blot from MNSE. MNSE were grown on permeable supports, tested in Ussing chambers, followed by lysis and assessment of CFTR protein levels by WB. Biological duplicates of LPS (10 μg/ml) and PBS-treated samples are shown on top. CFTR levels are plotted relative to beta-actin; $n = 5$, $p > 0.05$ (NS). (B) CFTR and TLR4 Western blot from CFBE cells. CFBE cells were treated with LPS or PBS. Western blots of biological duplicates of CFTR and TLR4 with beta-actin as loading control are shown on top. CFTR and TLR4 levels were plotted relative to beta actin. $n = 6$. (C) Cell surface and total CFTR levels in CFBE cells following cAMP (10 min forskolin 20 μM), LPS and LPS + cAMP treatment. cAMP enhances cell surface CFTR levels equally in LPS and PBS-treated cells ($p \leq 0.001$, $n = 6$). Surface CFTR levels are shown on top, $v = $ vehicle, cAMP = cAMP treatment. Following biotinylation, only Band C CFTR was pulled down (top), Total CFTR levels are shown in the middle. Both band B and C are present. Beta actin was used as loading control for total CFTR. Surface CFTR levels were plotted relative to total CFTR Band C levels; ***: $p \leq 0.001$. 
3.4. Evidence for the contribution of LPS-induced ROS to CFTR dysfunction

LPS-induced intracellular ROS accumulation in CFBE cells (as measured by the oxidation of 20,70-DCF diacetate to fluorescent DCF) occurs in a time-dependent fashion with a single exposure at a peak effect of 4 h similar to reductions in Isc noted in the Ussing chamber (Supplementary Figure 5). Cytoplasmic and mitochondrial ROS (percentage of fluorescent cells) were significantly elevated after 4 h of LPS exposure (cytoplasmic, LPS, 38.44 ± 3.27%, p ≤ 0.001, n = 11 and mitochondrial, LPS, 55.64 ± 15.14% vs. PBS control, 36.64 ± 12.47%, p ≤ 0.01, n = 12) as visualized with confocal microscopy (Fig. 5). Further, co-incubation of cells with LPS and the ROS scavenger, glutathione (GSH), eliminated the impact of LPS exposure on CFTR in the Ussing chamber (in pA/pF) (Fig. 6B, LPS, 9.67 ± 0.45 vs. PBS, 9.15 ± 1.22; LPS + GSH, 9.01 ± 0.81; and LPS + TAK-242, 9.05 ± 0.89; p ≤ 0.05, n ≥ 9). MCT was markedly inhibited in the LPS group when compared to the control group and when administered with GSH or TAK-242 (ASI; LPS, 4.28 ± 1.21 vs. PBS, 9.15 ± 1.22; LPS + GSH, 8.65 ± 1.63; LPS + TAK-242, 8.27 ± 1.63; p ≤ 0.001, n ≥ 9; PCL; LPS, 2.98 ± 0.46 vs. PBS, 4.74 ± 0.58; LPS + GSH, 4.48 ± 0.38; LPS + TAK-242, 4.76 ± 0.31; p ≤ 0.001, n ≥ 9). Mean CBF (Hz) was also significantly reduced with exposure to LPS compared to controls and addition of GSH and TAK-242 (LPS, 7.70 ± 1.22 vs. PBS, 9.15 ± 1.21; LPS + GSH, 9.01 ± 0.81; and LPS + TAK-242, 9.05 ± 0.89; p ≤ 0.05, n ≥ 9). MCT was markedly inhibited in the LPS group when compared PBS control vehicle (0.14 ± 0.07 mm/min vs. 0.45 ± 0.33 mm/min, p ≤ 0.001, n = 12) and when co-incubated with GSH and TAK-242 (0.36 ± 0.06 mm/min and 0.36 ± 0.04 mm/min, respectively; p ≤ 0.05, n ≥ 9). In MNSE cells, ASL depth was significantly reduced according to confocal microscopy z-stacked images; thus replicating our findings (with identical conditions) using another established technique. (Red-ASL; Green-Cell marker; 3.93 ± 0.32 vs. Control 5.19 ± 0.59, p ≤ 0.001, n ≥ 11).

3.5. MCT and other markers of airway functional microanatomy are decreased following LPS exposure

It is established that inhibition of transepithelial Cl− secretion leads to airway surface dehydration and diminished ASL and PCL depth, as well as consequent effects on the mucociliary transport apparatus, such as CBF [46,76,77]. In HSNE cultures, μOCT was used to assess changes to markers of mucociliary function after LPS incubation with or without GSH or TAK-242 (Fig. 7). Mean ASL and PCL depth thickness (μm) were significantly reduced in the LPS group when compared to the control group and when administered with GSH or TAK-242 (ASL; LPS, 7.70 ± 1.22 vs. PBS, 9.15 ± 1.21; LPS + GSH, 9.01 ± 0.81; and LPS + TAK-242, 9.05 ± 0.89; p ≤ 0.05, n ≥ 9). MCT was markedly inhibited in the LPS group when compared PBS control vehicle (0.14 ± 0.07 mm/min vs. 0.45 ± 0.33 mm/min, p ≤ 0.001, n = 12) and when co-incubated with GSH and TAK-242 (0.36 ± 0.06 mm/min and 0.36 ± 0.04 mm/min, respectively; p ≤ 0.05, n ≥ 9). In MNSE cells, ASL depth was significantly reduced according to confocal microscopy z-stacked images; thus replicating our findings (with identical conditions) using another established technique. (Red-ASL; Green-Cell marker; 3.93 ± 0.32 vs. Control 5.19 ± 0.59, p ≤ 0.001, n ≥ 11).

3.6. LPS inhibits nasal potential difference in vivo

Given our in vitro findings of acquired CFTR dysfunction in MNSE, we

![Fig. 5. (A & B) Cytoplasmic (green) and mitochondrial (red) ROS were significantly increased after LPS (10 μg/ml) exposure in CFBE cells. (C) The percentage of positive cells in LPS group was 2.5x higher (p ≤ 0.001, n = 11) than the control for cytoplasmic ROS and (D) 1.52x higher (p ≤ 0.01, n = 12) for mitochondrial ROS. (**p ≤ 0.01, ***p ≤ 0.001). (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)](image-url)
next tested LPS exposure in C57BL/6 mice in vivo (Fig. 8). LPS (10 μg/ml) + GSH (1 mM) or TAK-242 (1 μM) or PBS control solution embedded in a thermal transition gel were instilled intranasally for 4 h. LPS significantly reduced Cl– secretion across the nasal epithelium measured by NPD assay in response to low Cl– forskolin solution by approximately 30% (in mV; LPS, 4.98 ± 1.46 vs. Control, 7.34 ± 1.59, p ≤ 0.05, n = 6). When co-incubated with GSH or TAK-242, the response to low Cl– forskolin was similar to control vehicle and significantly improved over LPS incubation alone (GSH, 7.84 ± 1.40 and 7.36 ± 1.20, p ≤ 0.05, n = 6). This confirmed the findings that were observed in vitro were reflective of the in vivo situation following LPS exposure.

4. Discussion

This study demonstrates that brief exposures to physiologically relevant concentrations of LPS lead to a localized, CF-like mucosal dysfunction in mammalian RECs. LPS inhibited cyclic AMP-activated CFTR Cl– currents by decreasing the number of active channels and their open probability. The decrease in channel activity may be the result of channel deactivation (due to post-translational modification or alteration of signal transduction) and/or removal of channels from apical membranes. Our biotinylation studies show that CFTR Band C levels remain unchanged after LPS treatment (Fig. 4C). This led us to propose that the LPS-induced decrease of active CFTR channels was caused by deactivation of CFTR channels in the apical membrane. In Ussing chambers, CFTR-mediated Cl– secretion was significantly reduced as early as 30 min following LPS exposure with peak effects at 4 h and 10 μg/ml concentration without inducing cell death or deleterious effect on TEER. Although almost complete inhibition of forskolin-stimulated Cl– secretion measured in cells isolated with patch clamp was observed, there was a smaller, 20% reduction in forskolin-stimulated current change in the Ussing chamber after LPS exposure occurred. In contrast to patch clamping studies, vectorial transport of Cl– ions across confluent monolayers depends on a number of transporters (such as the Na+–K-ATPase, Na+–Cl cotransporter, K channels etc.). In addition, human sinonasal epithelial cell cultures contain small percentages of other cells that may influence the results. Regardless, the degree of CFTR functional inhibition by LPS was sufficient to reduce epithelial function including ASL, PCL, CBF, and MCT. The severity of the ion transport and functional microanatomy abnormalities were intermediate compared with the congenital absence of CFTR (no Cl– transport) observed in CF epithelial cells, but were capable of producing pronounced defects in mucociliary function at levels that would elicit poor mucociliary clearance and mucus retention in vivo especially if the exposure is persistent [78]. Importantly, the in vitro results were confirmed following in vivo exposure of mice to LPS that led to a marked decrease in transepithelial Cl– transport across the murine nasal airway with 4-h incubation. Our results indicate that LPS exposure resulted in elevated cytosolic and mitochondrial ROS levels and CFTR carbonylation, a post-translational modification that may also lead to reduced CFTR protein stability, reduced function and thickened airway secretions [20,79–82]. Indeed, the production of ROS induced by nitric oxide in LLC-PK1-cells stably expressing CFTR, as well as in human airway epithelial cell monolayers, decreased wild type CFTR levels and was accompanied by CFTR nitration [83]. These changes in CFTR levels and posttranslational nitration were accompanied by reduced cAMP-activated Cl– currents [16]. Although these studies applied a more prolonged (24–48 h) exposure to nitric oxide, it is possible that the produced ROS and CFTR carbonylation following LPS treatment can also lead to reduced CFTR levels after a long exposure, but was not studied in the current project.

**Fig. 6.** CFTR dysfunction was eliminated with the ROS scavenger GSH (1 mM). LPS + GSH co-incubation significantly increased Cl-transport compared to LPS only levels (p ≤ 0.001) in both Ussing chamber (A) and whole-cell patch-clamp (B) to a level comparable to PBS controls (C&D). CFTR protein carbonylation was increased after exposure to LPS, and normalized with LPS + GSH or LPS + TAK-242 (1 μM) co-incubation to levels similar to vehicle. (*p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ns = no significance).
These observations suggest that during a chronic infection multiple pathways may contribute to reduced CFTR function and acquired CF-like symptoms.

Our evidence for functional changes to CFTR implicate early oxidative stress as the primary mechanism responsible for LPS-induced acquired CFTR dysfunction based upon: 1) decreased open probability of CFTR, 2) reduced number of active channels without a concomitant decrease in surface levels of CFTR, 3) elimination of cAMP/PKA dependent effects as a source of the reduction, and 4) co-incubation with GSH eliminating the impact on CFTR function, MCT, and carbonylation. While there was a significant decrease in amiloride-sensitive Isc, this was consistent with previous observations by Boncoeur and colleagues [75] who found that LPS decreased αENaC in alveolar epithelial cells by ~50% in as little as 4 h by induction of ROS. Changes here could be attributable in part to a small, but significant effect on basolateral Na/K + ATPase pumps noted with apical permeabilization as seen in Supplementary Figure 1. This could decrease the driving force for apical cation and anion transport. However, this does not appear to be the primary mechanism of reduced cAMP-stimulated currents and cannot explain the impact on CFTR open probability. Reactive species generated intracellularly may oxidize cysteines and cause a number of additional post translational modifications which may result in loss of CFTR function. CFTR possesses 18 cysteine residues that are potential targets, and several oxidized forms of GSH have been shown to glutathionylate and inhibit the CFTR channel [84]. The identification of carbonyl adducts only represents a correlation and not a causal relationship. To prove this, identification of specific residues modified by LPS-generated ROS require mass spectrometry studies of purified CFTR. The very small quantities of CFTR in apical membranes of respiratory epithelial cells stymied our attempts to conduct these studies. Determining the biogenetic and structural modifications that confer loss of function to CFTR protein will offer vital insight regarding critical areas of inhibition, and describe a new method utilized by human pathogens such as Pseudomonas to abrogate the normal MCT mechanism. Such information is crucial for identifying new therapeutic targets for both CF and non-CF airway disease.

Conditioned media of PAO1 strain of P. aeruginosa has also been shown to trigger changes to respiratory epithelial function, primarily through endoplasmic reticulum stress and subsequent activation of the unfolded protein response (UPR) [85]. While the authors attributed the effects to pyocyanin and other secreted factors, LPS was not evaluated. Importantly, UPR activation leads to inhibition of CFTR transcription as well as reduction in CFTR maturation [86,87]. Given the early functional response to LPS in our studies and the lack of change in total and surface CFTR protein, or mRNA levels, it is unlikely that oxidative stress or activation cellular stress responses such as UPR or the integrated stress response (ISR)-mediated CFTR expression reduction was involved in the functional changes. In other words, the markedly reduced CFTR

Fig. 7. μOCT of HSNE shows airway surface liquid, periciliary liquid, ciliary beat frequency, and mucociliary transport were all significantly reduced after LPS (10 μg/ml) incubation - an effect that was eliminated with co-incubation with GSH (1 mM) and TAK-242 (1 μM). A: Cross sectional μOCT images of HSNE treated with LPS (10 μg/ml)+/- GSH (1 mM) or TAK-242 (1 μM) or PBS vehicle control for 4 h. ASL depth (red bar) and PCL depth (yellow bar) are demarcated. B-E: Summary data showing ASL depth (B), PCL depth (C), CBF (D), and MCT (E). F: In MNSE, cells were stained to visualize overlying surface liquid. Confocal images demonstrate decreased ASL depth from inhibition of apical Cl− secretion (red-ASL; green-cell marker). G: LPS significantly decreased ASL depth (in μm) in murine nasal septal epithelial cultures (*p≤0.05, ***p≤0.001). (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)
channel open probability and diminished numbers of active channels cannot be attributed to protein degradation or turnover. While chronic activation of redox pathways by LPS would be likely to confer ER stress, UPR or IRS, these mechanisms are unlikely to be the cause of functional changes to the channels exhibited in our study. Furthermore, increased intracellular Cl\(^{-}\), shown to trigger an IL-1\(\beta\) autocrine and positive feedback loop in CF bronchial epithelial cells and ROS-mediated inflammation, \([88–90]\) was observed in a 24–48 h time frame, well outside the window of observed effects in the present study. Yet, it is possible that the changes caused by ROS (carbonylation) affected the function of CFTR, particularly since CFTR protein expression (bands B and C), as well as surface localization, were not altered. GSH, the most abundant cellular thiol antioxidant, critical to maintaining intracellular redox balance and redox-sensitive signaling events abrogated the LPS-induced acquired CFTR dysfunction and subsequent carbononylation of CFTR. Previous studies have shown GSH is chronically low in the fluid that lines the epithelial layer of lung tissue subjected to chronic \textit{Pseudomonas} infection \([91,92]\). Pulmonary GSH levels are also decreased in CF patients, but intracellular redox potential in CF and CFTR-corrected nasal epithelial cells is normal. Thus, \textit{P. aeruginosa}-derived LPS could be one of the factors responsible for oxidative stress and impairments in GSH status in the airways of CF patients \textit{in vivo}. Importantly, strategies to scavenging oxidative stress in the airways represent an important platform for disease intervention.

Acquired CFTR deficiency contributing to the propagation of chronic infection in CRS differs substantially from prevailing models regarding the pathogenesis of CRS \([93–97]\). The possibility of dysfunctional Cl\(^{-}\) transport in CRS suggests new avenues relevant to the field of rhinology and sinus disease; in particular, LPS decreasing CFTR-mediated transport has important implications for understanding impaired MCT in numerous upper and lower respiratory infectious illnesses. Regardless of the mechanism underlying LPS-mediated CFTR dysfunction, increasing transepithelial Cl\(^{-}\) secretion and apical fluid hydration are appealing options for therapeutic intervention, and the evidence for pronounced dysfunction provides an impetus for activating CFTR as a means to promote MCT in individuals with CRS. Furthermore, blunting cellular oxidative stress with ROS scavenger compounds could reverse MCT defects and improve respiratory epithelial health in the setting of early colonization and/or infection. Although this therapeutic approach could have harmful effects in patients by impacting innate immunity, tactics to utilize topical, inhaled drugs to reduce ROS are already clinically approved for use – in particular, the redox agent and free radical scavenger, N-acetyl cysteine, has been used to remove mucus plugs in CF and other respiratory diseases of mucus obstruction \([98,99]\).

In conclusion, this study demonstrated that pathophysiological concentrations of LPS adversely affect REC CFTR function through the generation of ROS and confer substantial dysfunction to markers of MCT. The major effect of LPS causes potent inhibition to active CFTR Cl\(^{-}\) transport and MCT, and thus permits colonization and propagation of infection with \textit{P. aeruginosa} and other GNB in upper and lower airways. While the findings have implications for early colonization and infection with GNB in non-CF airways, LPS is a critical factor that should also be considered in CF airways colonized or recurrently infected with \textit{Pseudomonas}, including the respiratory tract of patients with improved CFTR function due to CFTR modulator therapy. Furthermore, in the setting of chronic GNB infection in non-CF CRS, restoration of impaired CFTR function with CFTR potentiators or ROS scavengers represents a critical and innovative approach to therapy.

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Declaration of competing interest

The authors declare no competing financial interests.

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Appendix A. Supplementary data

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