Liposomes Loaded With Phosphatidylinositol 5-Phosphate Improve the Antimicrobial Response to Pseudomonas aeruginosa in Impaired Macrophages From Cystic Fibrosis Patients and Limit Airway Inflammatory Response

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Despite intensive antimicrobial and anti-inflammatory therapies, cystic fibrosis (CF) patients are subjected to chronic infections due to opportunistic pathogens, including multidrug resistant (MDR) Pseudomonas aeruginosa. Macrophages from CF patients show many evidences of reduced phagocytosis in terms of internalization capability, phagosome maturation, and intracellular bacterial killing. In this study, we investigated if apoptotic body-like liposomes (ABLs) loaded with phosphatidylinositol 5-phosphate (PI5P), known to regulate actin dynamics and vesicular trafficking, could restore phagocytic machinery while limiting inflammatory response in in vitro and in vivo models of MDR P. aeruginosa infection. Our results show that the in vitro treatment with ABL carrying PI5P (ABL/PI5P) enhances bacterial uptake, ROS production, phagosome acidification, and intracellular bacterial killing in human monocyte-derived macrophages (MDMs) with pharmacologically inhibited cystic fibrosis transmembrane conductance regulator channel (CFTR), and improve uptake and intracellular killing of MDR P. aeruginosa in CF macrophages with impaired bactericidal activity. Moreover, ABL/PI5P stimulation of CFTR-inhibited MDM infected with MDR P. aeruginosa significantly reduces NF-κB activation and the production of TNF-α, IL-1β, and IL-6, while increasing IL-10 and TGF-β levels. The therapeutic efficacy of ABL/PI5P given by pulmonary administration was evaluated in a murine model of chronic infection with MDR P. aeruginosa. The treatment with ABL/PI5P significantly reduces pulmonary neutrophil infiltrate and the levels of KC and MCP-2 cytokines in the lungs, without affecting pulmonary bacterial load. Altogether,
INTRODUCTION

Cystic fibrosis (CF) is an autosomal recessive genetic disease caused by a mutation in the gene encoding the cystic fibrosis transmembrane conductance regulator channel (CFTR) (1). The CFTR is usually expressed on the apical membrane of epithelia, and its dysfunction causes a defective chloride secretion leading to a modification in the airway surface liquid (2). The pathophysiological changes in CF result in a systemic disease, which affects the pancreas, liver, reproductive tract, and mainly the lungs (3). Here, the loss of function of CFTR causes a defective mucociliary clearance and a dramatic production of sticky mucus, which is associated with chronic infection by opportunistic pathogens, such as *P. aeruginosa* (4). Infections sustained by MDR *P. aeruginosa* in CF are increasing, reflecting cumulative exposure to antibiotic treatment (5). Moreover, the chronic bacterial infections associated with the persistent inflammation, leading to pulmonary insufficiency, represent the main cause of mortality and morbidity in CF patients (6). Today, the identification of novel host- and/or pathogen-directed therapeutic tools represents an urgent challenge for the scientific community to fight the emergence of MDR pathogens, as well as a priority also at the global level.

The defective antimicrobial response exerted by innate immune cells in CF patients has been documented and depends, at least in part, on a dysfunctional phagocytosis process (7, 8). Phagocytosis is an important innate effector mechanism deputed to the intracellular elimination of invading pathogen by the generation of highly microbialic organelles called phagolysosomes. These organelles originate from a phagosome, generated by the invagination of plasma membrane, which matures to a fully microbicidal phagolysosome, composed by 1,2-dioleoyl-sn-glycero-3-phospho-(1′-myo-inositol-5′-phosphate) (PI5P, Avanti Polar Lipids) were added to a 99:1 dodecane:silicone solution to obtain a monodispersed vesicles >0.2 μm in diameter.

The second lipid messenger phosphatidylinositol 5-phosphate (PI5P) is a minor phosphoinositide representing less than 10% of the total lipids (12). PI5P can be directly produced from phosphatidylinositol (PI) by the activity of phosphoinositide 5-kinase (PIKfyve) or by the dephosphorylation of phosphatidylinositol 3,5-bisphosphate (PI3,5P2) by myotubularin 3-phosphatases (13). PI5P is present at the cellular membrane and at the early phagosome (14), and its level resulted increased during the late stages of the phagocytosis process (15). Moreover, it can regulate endosome vesicle trafficking (16), cellular actin remodeling, and bacterial invasion (14), and can be involved in class III phosphatidylinositol 3-kinase (Vps34)-independent autophagy activation (17).

In this study, we have generated asymmetric apoptotic body-like liposomes (ABLs) composed by phosphatidylserine (PS) at the outer membrane surface resembling an apoptotic body, to target macrophages and to downmodulate inflammatory reaction (18), and by the bioactive lipid PI5P at the inner membrane surface to enhance the phagocytosis process. In particular, this study evaluates the immunotherapeutic value of ABL/PI5P *in vitro* in impaired macrophages from CF patients and *in vivo* in models of *P. aeruginosa* infection, assessed in terms of i) uptake and intracellular bacterial killing, ii) mechanisms of bactericidal activity, and iii) potentially tissue-damaging inflammatory response.

MATERIAL AND METHODS

Liposome Preparation

Apoptotic body-like liposomes (ABLs) were produced as previously described (19). Briefly, the inner monolayer lipids composed by 1,2-dioleoyl-sn-glycero-3-phospho-(1′-myo-inositol-5′-phosphate) (PI5P, Avanti Polar Lipids) were suspended in anhydrous dodecane (Sigma) at a concentration of 0.05 mg/ml. L-α-phosphatidyserine (PS, Avanti Polar Lipids) was used as outer monolayer lipid and was added to a 99:1 dodecane:silicone solution to obtain a final concentration of 0.05 mg/ml. Asymmetric liposomes were prepared by adding 2 ml of outer monolayer lipid suspension over 3 ml of cell culture medium (for *in vitro* experiments) or saline (for *in vivo* experiments). Finally, 100 μl of the inner monolayer lipid suspensions were added over the 2-ml lipid phase, and the samples were centrifuged at 120 × g for 10 min. After the centrifugation, ABLs were collected in the aqueous phase using FACSCalibur (Becton Dickinson), allowing quantification of monodispersed vesicles >0.2 μm in diameter.

Cell Culture

Primary monocyte-derived macrophages (MDMs) were prepared as previously described (17). Briefly, peripheral blood mononuclear cells (PBMCs) from healthy donors and CF patients were isolated by Ficoll density gradient, and monocytes were then positively sorted using anti-CD14...
monoclonal antibodies conjugated to magnetic microbeads (Miltenyi Biotec), according to manufacturer’s instructions. Monocytes were then suspended in complete medium and incubated for a further 5 days in 96-well plates at a concentration of 10^6 cells/ml in the presence of M-CSF (50 ng/ml, Miltenyi Biotec) to get differentiated macrophages.

**Bacteria**

MDR *P. aeruginosa* strain (ATCC® BAA-2113) was used in *in vitro* experiments and MDR-RP73 *P. aeruginosa* clinical isolate (20) was used in an *in vivo* mouse model of chronic *P. aeruginosa* infection (21, 22). The BAA-2113 single colony was collected by streaking on Trypticase soy agar (TSA, BD Difco™) and then suspended in 15 ml of Trypticase soy broth (TBS, BD Difco™). Bacteria were grown in Erlenmeyer flask at 37°C under stirring for 18 h, and their growth was monitored by measuring the optical density at a wavelength of 600 nm by Varioskan LUX Multimode Microplate Reader (Thermo Fisher Scientific). BAA-2113 was stored at −80°C until use after suspension in TSB and 30% glycerol.

For *in vivo* experiments, an aliquot of RP73 strain from glycerol stocks (TSB + 25% glycerol) was streaked for isolation on TSA and incubated at 37°C O/N. One colony was picked from the plate and used to inoculate 10 ml of TSB and placed overnight in a shaking incubator at 37°C 200 rpm. Thereafter, bacterial suspension was diluted to 0.15 OD/ml in 20 ml of TSB/flask and grown for 4 h at 37°C at 200 rpm, to reach the log phase.

**Patients**

CF patients (n = 19) were enrolled at “Bambino Gesù” Children’s Hospital in Rome, Italy. All of the CF patients were clinically stable at the time of blood donation (5 ml). Controls (n = 20) were represented by buffy coats from healthy blood donors, attending at the Blood Transfusion Unit of Policlinico “Umberto I” in Rome, Italy. Clinical and demographic features of CF patients as well as healthy controls are summarized in **Table 1**.

**Evaluation of *In Vitro* Bacterial Uptake and Intracellular Growth**

To assess bacterial uptake, MDMs from healthy donors or from CF patients were distributed in 96-well plates at a concentration of 2 × 10^5 cells/well and were stimulated with ABL/PI5P used at a ratio of 1:1 (ABL:MDM), for 30 min before infection and/or simultaneously with the infection, in the presence or absence of the CFTR inhibitor INH172 (Sigma), used at a concentration of 10 µM. Then cells were washed once and infected with MDR *P. aeruginosa* for 1 h at 37°C at an MOI of 30 in the presence or absence of INH172. Thereafter, extracellular bacilli were killed at 1 h of incubation with 400 µg/ml amikacin. Finally, cells were lysed with 1% deoxycholate (Sigma), samples were diluted in PBS-tween 80, and colony-forming units (CFUs) were quantified by plating bacilli in triplicate on TSA.

To assess intracellular bacterial growth, MDMs from healthy donors or from CF patients were distributed in 96-well plates at a concentration of 2 × 10^5 cells/well and were infected with MDR *P. aeruginosa*, for 1 h at 37°C at an MOI of 30, in the presence or absence of INH172, used at a concentration of 10 µM. Thereafter, extracellular bacilli were killed at 1 h of incubation with 400 µg/ml amikacin. Cells were then washed and incubated with ABL/PI5P, added to a ratio of 1:1 (ABL:MDM) for a further 2 h, in the presence or absence of INH172. Finally, cells were lysed with 1% deoxycholate (Sigma), samples were diluted in PBS-tween 80, and CFUs were quantified by plating bacilli in triplicate on TSA.

**Table 1** Demographic and clinical characteristics of cystic fibrosis (CF) patients and healthy donors (HD).

| CF Age (range) | Genotype | Microbiology | FEV-1 (%) | HD Age (range) |
|---------------|----------|--------------|-----------|----------------|
| 1             | 26-30    | F508del/N1303K | S.a, A.x, S.m. | 44 | 1 | 36-40 |
| 2             | 16-20    | F508del/P5L | S.a | 79 | 2 | 21-25 |
| 3             | 11-15    | G576A/R686C | S.a, E.n.c, E.a | 106 | 3 | 51-55 |
| 4             | 21-25    | F508del/F508del | S.a, B.b | 85 | 4 | 56-60 |
| 5             | 31-35    | F508del/F508del | S.a, Buc | 72 | 5 | 41-45 |
| 6             | 21-25    | F508del/621+1G>T | P.a, Buc | 50 | 6 | 36-40 |
| 7             | 41-45    | F508del/F508del | S.m, Buc | 91 | 7 | 56-60 |
| 8             | 31-35    | F508del/W1282X | S.a, P.a | 109 | 8 | 56-60 |
| 9             | 26-30    | F508del/F508del | S.a, E.sc | 115 | 9 | 41-45 |
| 10            | 21-25    | F508del/G1244E | S.a | 97-103 | 10 | 56-60 |
| 11            | 26-30    | F508del/G424X | E.f, S.a, C.g, S.m | 65 | 11 | 55-58 |
| 12            | 26-30    | DeltaF508/G856E | S.a, P.m, S.m, A.t | 82 | 12 | 46-50 |
| 13            | 6-10     | DeltaF508/R334L | S.p, S.a, H.i | 96-104 | 13 | 21-25 |
| 14            | 11-15    | R553X/3277-26A>G | S.a | 88 | 14 | 41-45 |
| 15            | 6-10     | G856E/621+1G>T | A.i, H.i | 86 | 15 | 56-60 |
| 16            | 21-25    | G1214E/T338I | H.i | 102 | 16 | 51-55 |
| 17            | 6-10     | 1717-1G>A/E831X | GAS, B.b, H.i | 119 | 17 | 41-45 |
| 18            | 16-20    | W1282X/2789+5G>A | S.a, S.ca | 85 | 18 | 31-35 |
| 19            | 26-30    | DeltaF508/DeltaF508 | S.a, P.a | 25 | 19 | 21-25 |
| 20            |          |              |             | 20 | 26-30 |

S.a, Staphylococcus aureus; A.x, Achromobacter xylosoxidans; S.m, Stenotrophomonas maltophilia; E.n.c, Enterobacter cloacae; E.a, Enterobacter asburiae; B.b, Bordetella bronchiseptica; Buc, Burkholderia cepacia; P.a, Pseudomonas aeruginosa; E.sc, Escherichia coli; E.f, Enterococcus faecalis; C.g, Candida glabrata; P.m, Proteus mirabilis; A.t, Aspergillus fumigatus; S.p, Streptococcus pneumoniae; H.i, Haemophilus influenzae; GAS, Group A Streptococcus; B.b, Branhamella catarrhalis; S.a, Scotoesporium apiospermum.
In order to evaluate the role of ROS and of phagosome acidification in intracellular bacterial killing, *P. aeruginosa*-infected cells were treated simultaneously with ABL/PI5P and either PEG-Catalase (100 U/ml) or Concanaamycin A (10 nM), respectively.

**Fluorimetric Analysis**

Phagosomal acidification was assessed by using the fluorescent probe Lysosensor green DND 189 (Molecular Probes) (23), which measures the pH of acidic organelles, such as phagosomes/lysosomes. Briefly, MDM from healthy donors were pretreated or not for 1 h with 10 µM INH172 and then exposed or not to Crimson fluorescent microbeads (1 µm Fluospheres® carboxylate-modified microspheres, LifeTechnologies), for 1 h at 37°C at a ratio of 5:1 in the presence or absence of 10 µM of INH172, in order to exclude possible differences in microbead internalization among experimental groups. Then cells were washed and incubated for a further 90 min with ABL/PI5P, added to a ratio of 1:1 (ABL:MDM), in the presence or absence of INH172. Cells were stained for 15 min at 37°C with 1 µM of Lysosensor green DND 189. pH calibration curve was obtained by incubating macrophages in calibration buffers at pH 4.5, 5.5, 6.5, and 7.5 (Intracellular pH Calibration Buffer Kit, Molecular Probes), and by labeling cells for 15 min at 37°C with 1 µM of Lysosensor green DND 189 according to the manufacturer’s instructions. pH was evaluated by fluorometry by setting the wavelength of excitation at 443 or 625 nm and emission at 505 or 645 nm, for Lysosensor green DND 189 and Crimson fluorescent microbeads, respectively.

ROS generation was analyzed by loading MDM isolated from healthy donors with the fluorescent indicator 20,70-dichloro fluorescein diacetate (DCF, Molecular Probes), used at a concentration of 10 µM, for 40 min at 37°C in the dark. Thereafter, MDM isolated from healthy donors were pretreated or not for 1 h with 10 µM INH172 and then exposed or not to Crimson fluorescent microbeads (1 µm Fluospheres® carboxylate-modified microspheres, Life Technologies), for 1 h at 37°C at a ratio of 5:1 in the presence or absence of 10 µM of INH172, in order to exclude possible differences in microbead internalization among experimental groups. Cells were then washed and incubated for a further 90 min in the presence or absence of INH172 with ABL/PI5P, added to a ratio of 1:1 (ABL:MDM). The production of ROS was evaluated by fluorometry by setting the wavelength of excitation at 443 or 625 nm and emission at 505 or 645 nm, for DCF and Crimson fluorescent microbeads, respectively. Fluorescence has been evaluated by the use of a Varioskan LUX Multimode Microplate Reader (Thermo Fisher Scientific).

**Mouse Model of Chronic Infection**

Immunocompetent C57Bl/6Ncr BR male mice (n = 16 treated with 3×10^5 ABL/PI5P and n = 16 treated with vehicle) were challenged with 3–4×10^3 CFUs of the *P. aeruginosa* MDR-RP73 embedded in agar beads for chronic infection by intratracheal (i.t.) administration. Agar beads were prepared following established procedures (21, 24). Local treatment by Penn-Century MicroSprayer® Aerosoliser with 3×10^5 ABL/PI5P started soon (5 min) after infection and was repeated daily for 6 days. Body weight and health status were monitored daily. After 6 days postinfection, lung CFUs and cell counts in the bronchoalveolar lavage fluid (BALF) were analyzed as previously described (21, 24). Finally, 6 days after infection, murine lungs were excised aseptically and homogenized in 2 ml of PBS added with protease inhibitors (Complete™ Protease Inhibitor cocktail—Roche) using the homogenizer GentleMACS™ Octo Dissociator, and the levels of TNF-α, KC, JE, and MIP-2 in the supernatant of murine lungs were measured by ELISA kit (DuoSet® ELISA Development Systems).

**Enzyme-Linked Immunosorbent Assay**

MDMs were infected or not with *P. aeruginosa* (MOI 30) in the presence or absence of INH172 and stimulated or not with ABL/PI5P at a ratio of 1:1 (ABL:MDM) for 2 h. Thereafter, supernatants were collected, cells were lysed, and both stored at −20°C until analysis. The levels of tumor necrosis factor-α (TNF-α), interleukin-1β (IL-1β), IL-6, IL-10, and transforming growth factor-beta (TGF-β) in the supernatants of MDMs were measured by human TNF-α ELISA kit (BD Biosciences), human IL-6 DuoSet® ELISA Development Systems, human IL-1β DuoSet® ELISA Development Systems, human IL-10 DuoSet® ELISA Development Systems, and human TGF-β DuoSet® ELISA Development Systems (all by R&D system) and used according to the manufacturer’s instructions. The levels of murine TNF-α, KC, JE, and MIP-2 were measured by DuoSet® ELISA Development Systems (R&D system). The activation of NF-kB transcription factor was assessed on lysed cells by “NFkB p65 (Total/Phospho) Human InstantOne™ ELISA Kit” (Invitrogen) and used according to the manufacturer’s instructions.

**Statistics**

Comparison between groups was done using Student’s t test, as appropriate, for normally distributed data. The Wilcoxon rank test sum or Mann–Whitney test was performed for data that were not normally distributed.

**Ethics Statement**

 Buffy coats from anonymized healthy donors, who gave their written informed consent to donate the nonclinically usable components of their blood for scientific research, were obtained from the Blood Transfusion Unit of Policlinico “Umberto I” in Rome. The present study, which is based on nonclinical in vitro research, did not require any specific approval from an ethical committee, according to the Italian law (decree by Ministero della Salute by February 8, 2013, published on Gazzetta Ufficiale della Repubblica Italiana no. 96 of April 24, 2013, and legislative decree no. 211 of June 24, 2003, published on Gazzetta Ufficiale della Repubblica Italiana no. 184 of August 9, 2003). Cystic fibrosis patients, giving their (or parental) informed consent to participate in the study, were enrolled at “Bambino Gesù” Children’s Hospital in Rome after having received detailed information on the scope and objectives of the study by a sanitary personnel who explained the patient...
information leaflet (ethics approval #738/2017 of “Bambino Gesù” Children’s Hospital, Rome).

Animal studies adhered to the Italian Ministry of Health guidelines for the use and care of experimental animals (IACUC #733).

Research with P. aeruginosa RP73 clinical isolate from CF patient has been approved by the Ethics Commission of Hannover Medical School, Germany. The patient and parent gave informed consent before the sample collection.

RESULTS

ABL Loaded With PI5P Improve Dysfunctional Bacterial Uptake in CF and INH172 Treated Macrophages

CF macrophages show defective P. aeruginosa internalization (25–27). Hence, we tested the capability of ABL carrying PI5P to improve phagocytosis of MDR P. aeruginosa in macrophages with disabled CFTR. Results confirmed that the bacterial uptake of MDM from CF patients or INH172-treated MDM from healthy donors was dysfunctional compared to that of untreated MDM (Figure 1A). The dysfunctional bacterial uptake capacity was significantly improved by the preventive treatment with ABL/PI5P of INH172-treated dTHP-1 cells, infected with MDR P. aeruginosa at an MOI of 30 and 10, and resulted completely restored at an MOI of 30 (Figure S1A). Moreover, this effect was specific for ABL/PI5P, as any effect was not observed when liposomes composed by either PS or PI5P only were used (Figure S1B). Bacterial internalization was also improved by the pretreatment with ABL/PI5P of primary MDM, with pharmacologically inhibited CFTR (Figure S2), and of CF MDM (Figures 1B, C). No modification of the bacterial uptake was observed when ABL/PI5P was used simultaneously with MDR P. aeruginosa infection, excluding that liposomes exerted their effect interacting with the pathogen (Figure S2).

Treatment With ABL/PI5P Rescues Impaired Phagosome Maturation and ROS Generation in Macrophages With Pharmacologically Inhibited CFTR

Dysfunctional activity of CFTR leads to impaired phagosome maturation due to unbalanced influx of chloride ions (Cl−) that does not allow intraphagosomal acidification (8). In this context, we determined basal intracellular pH and ROS production, both in the normal and CFTR-pharmacologically inhibited macrophages. MDMs with CFTR functionally inhibited by INH172 had a more basic intracellular pH than untreated MDM and, after exposure to microbeads, showed an impaired phagosome acidification (Figure 2A), which could be completely restored after 90 min of treatment with ABL/PI5P (Figure 2A). This result was confirmed by using microbeads labeled with NHS, a pH-sensitive fluorochrome, whose fluorescence decreases proportionally to acidification of phagosome microenvironment: MDMs with CFTR functionally inhibited by INH172 and treated with ABL/PI5P showed a reduction of NHS fluorescence at levels comparable to that of control MDMs (Figure S3).

Phagosome acidification and ROS generation are sequential steps leading to intracellular bacterial killing and type II NADPH oxidase (NOX-2) assemblies from component subunits on maturing phagosomes (28). On these grounds, we monitored ROS generation in MDM with or without pharmacologically inhibited CFTR following exposure to microbeads and after 90 min of treatment with ABL/PI5P. As expected, the exposure to microbeads induced a significant ROS generation in control cells (Figure 2B). On the contrary, the exposure to microbeads provoked an impaired ROS production in MDM with INH172-inhibited CFTR, which was significantly restored by the ABL/PI5P treatment (Figure 2B). Together, these results show that the inhibition of CFTR by INH172 causes an impaired phagosome acidification and a reduced ROS production that could be significantly recovered by the treatment with ABL/PI5P.

ABL/PI5P Promote Intracellular Bacterial Killing of INH172-inhibited Control Macrophages and CF Macrophages

Since ABLs/PI5Ps were shown to restore the functional intraphagosomal acidification and oxidative burst of macrophages with pharmacologically inhibited CFTR, we investigated whether an increased bactericidal activity against MDR P. aeruginosa strains could also represent a functional consequence of ABL/PI5P treatment of cells with altered CFTR function. In this context, we preliminarily tested the capability of ABL/PI5P to improve intracellular bacterial killing in dTHP-1 cells with disabled CFTR infected with MDR P. aeruginosa (BAA-2113 strain) at the MOI of 30 and 10. Results expressed in Figure S4A show a significant reduction in intracellular bacterial viability after exposure to ABL/PI5P, which was higher at an MOI of 30. Moreover, such effect was specific for ABL/PI5P, as any effect was not observed when liposomes composed by either PS or PI5P only were used (Figure S4B).

Thereafter, we investigated the effect of ABL/PI5P on primary MDMs with pharmacologically inhibited CFTR. Our results show that 2 h of ABL/PI5P treatment on INH172-treated MDM significantly enhances the intracellular killing of MDR P. aeruginosa strain (BAA-2113) (Figure 3A) as well as of a panel of additional three MDR P. aeruginosa strains (BAA-2108, BAA-2111, and BAA-2112) (Figure S5).

In order to evaluate the role of phagosome acidification and/or of ROS generation in intracellular killing of MDR P. aeruginosa induced by ABL/PI5P, we exposed P. aeruginosainfected cells to either Concanaamycin A (ConA), a specific inhibitor of V-ATPases blocking phagosome acidification, or polyethylene glycol-Catalase (PEG-Cat), which reduces hydrogen peroxide to water. Results show that intracellular killing of MDR P. aeruginosa, induced by ABL/PI5P stimulation of MDM with pharmacologically inhibited CFTR, is ROS mediated and phagosome acidification dependent, as it results ineffective in the presence of Peg-Cat and Conc A, respectively (Figure 3B).
Finally, we tested the efficacy of ABL/PI5P in MDMs from CF patients. On the basis of the efficacy of freshly isolated and nontreated CF macrophages to limit intracellular bacterial growth, we could divide patients in two groups: “impaired” and “controller,” according to intracellular bacterial replication index higher or lower than 1, respectively (Figure 4A). Notably, MDM isolated from patients of the “impaired” group were susceptible to ABL/PI5P stimulation (Figure 4C), increasing significantly their intracellular killing upon liposome treatment, whereas ABL/PI5P did not further increase the intracellular killing of MDM isolated from patients belonging to the “controller” group (Figure 4B) or from healthy donors (Figure S6).

**ABL/PI5P Treatment Modulates Anti- and Pro-Inflammatory Cytokine Production in Macrophages With Pharmacologically Inhibited CFTR**

Chronic infection, mainly due to *P. aeruginosa*, and unresolved acute inflammation are key mechanisms responsible for progressive lung destruction in CF (29) and an effective host-directed therapeutic strategy should also limit the inflammation-based immunopathology. On the basis of previous results showing the anti-inflammatory effect of ABL (18), we wanted to investigate the effect of ABL/PI5P treatment of MDM incubated or not with INH172 on NF-kB activation and on the production of a panel of pro- and anti-inflammatory cytokines after infection with MDR *P. aeruginosa*. In this model, we could show high basal levels of NF-kB activation after CFTR inhibition, which further increased following infection with MDR *P. aeruginosa*. Interestingly, the same NF-kB activation levels were significantly reduced by the treatment with ABL/PI5P (Figure 5A). The reduced activation of NF-kB was confirmed by the comparative *in vitro* measure of cytokines whose transcription depends upon NF-kB activity (TNF-α, IL-1β, and IL-6). In fact, infected macrophages with dysfunctional CFTR showed a significant increase in TNFα, IL-1β, and IL-6 secretion in comparison with control infected macrophages, and ABL/PI5P treatment reduced the levels of the same inflammatory cytokines in infected macrophages irrespective of CFTR inhibition (Figures 5B–D). On the contrary, the secretion of anti-inflammatory cytokines, such as IL-10 and TGF-β, was significantly increased in ABL-/PI5P-treated MDMs (Figures 5E, F).

**ABL/PI5P Therapeutic Treatment Reduces Inflammatory Reaction in a Murine Model of MDR *P. aeruginosa* Chronic Infection**

We wanted to test in an *in vivo* model the functional consequences of the *in vitro* observed anti-inflammatory functions of ABL/PI5P in addition to the promotion of intracellular killing of pathogens. This is particularly interesting since massive neutrophil infiltration is the main cause of chronic damage to the epithelial lung structure in the CF lung (30). Thus, we tested the efficacy of ABL/PI5P administrated by Penn-Century MicroSprayer Aerosoliser in mice, 5 min after infection with MDR *P. aeruginosa* embedded in agar beads. An evaluation of the inflammatory response and bacterial burden in lung and in BALFs was considered as read-

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**FIGURE 1 |** Dysfunctional *Pseudomonas aeruginosa* uptake in macrophages with pharmacologically inhibited or naturally mutated cystic fibrosis transmembrane conductance regulator channel (CFTR) and its enhancement by apoptotic body-like liposome/phosphatidylinositol 5-phosphate (ABL/PI5P) stimulation. (A) Monocyte-derived macrophages (MDMs) from healthy donors, treated or not with INH172, or from cystic fibrosis (CF) patients were infected with multidrug-resistant (MDR) *P. aeruginosa* (BAA-2113 strain) at an MOI of 30. (B, C) CF MDMs were stimulated or not with ABL/PI5P for 30 min before infection (B) or before and during infection (C). Cells were then infected with MDR *P. aeruginosa* (BAA-2113 strain) at an MOI of 30. The bacterial uptake was quantified by colony-forming unit (CFU) assay and indicated as phagocytosis index, calculated as the ratio between the CFUs obtained immediately after the infection and the inoculum. (A) Statistical analysis was performed by using the two-sided Mann–Whitney test and *p < 0.05; **p < 0.01* in comparison with control cells (healthy donors, n = 6; CF patients, n = 6). (B, C) Statistical analysis was performed by using the two-sided Wilcoxon rank sum test (B, n = 6) *p = 0.03* and (C, n = 9) *p = 0.004.*
out measures of ABL/PI5P treatment efficacy. Results showed a significant reduction of both KC and MIP-2 (Figures 6A, B) and no significant variations in the levels of TNF-α and MCP-1 (Figures 6C, D) in the lungs of ABL/PI5P-treated mice in comparison with vehicle-treated mice. Results also showed a significant reduction in neutrophil count in BALF (Figure 7B) of ABL/PI5P-treated mice in comparison with vehicle-treated mice. A reduction, although not significant, of BALF total cells (Figure 7A) and macrophages (Figure 7C) was observed. Of note, the significant reduction in BALF neutrophils observed in ABL/PI5P-treated mice did not significantly interfere with pulmonary bacterial burden (Figure 7D).

DISCUSSION

CF is a genetic disorder that leads to a progressive dysfunction of lung activity by predisposing patients to colonization by...
opportunistic bacterial pathogens. Infections caused by *P. aeruginosa*, particularly because of the emergence of MDR strains, represent the major cause of morbidity and mortality in CF patients (31). These evidences highlight the urgency to develop novel therapeutic approaches, which may contribute to the control of MDR pathogens, including *P. aeruginosa*. Phagocytosis and intracellular killing of extracellular pathogens are the most important effector mechanisms of innate immune cells that can be hampered in CF patients (26). Hence, strategies aimed at improving the capacity of lung resident innate immune cells to phagocytose and kill pathogens may represent a promising host-directed approach to combat bacterial lung infections in CF patients.

In the present manuscript, we show that ABLs carrying PI5Ps are able to increase, both *in vitro* and *ex vivo*, the capacity of INH172-treated and CF macrophages to internalize and kill MDR strains of *P. aeruginosa*. Moreover, in a murine model of *in vivo* *P. aeruginosa* infection, we show that ABLs carrying PI5Ps are capable of reducing neutrophil recruitment and lung inflammation, without promoting bacterial growth. In particular, we show that treatment with ABL/PI5P enhance nonopsonic *P. aeruginosa* phagocytosis in CF and INH172-treated macrophages. Several not mutually exclusive mechanisms may explain this observation. PI5P may promote actin dynamics and bacterial phagocytosis i) via recruitment and activation of the exchange factor Tiam1 and Rac1 (14), ii) by directly activating PI3K/Akt signaling pathway (32), or iii) by participating as substrate to the PI(4,5)P2 production (33), which may directly induce membrane remodeling (34) or be converted, by means of phosphoinositide 3-kinase (PI3K), in 3,4,5-tris phosphate [PI (3,4,5)3], which in turn is able to activate Akt signaling pathway (35).

We then showed that ABL/PI5P treatment restores intracellular acidification and ROS production of human macrophages, whose CFTR was pharmacologically inhibited. Following phagocytosis, phagosome maturation requires the sequential interaction with early endosomes, late endosomes, and ultimately, with lysosomes, leading to the generation of a highly microbiocidal organelle called phagolysosome. In pharmacologically inhibited- or CF-macrophages, the altered CFTR function leads to a limited phagosome acidification because of the unbalanced CI− ion distribution, which alters phagolysosome maturation and causes a defective intracellular bacterial clearance (8, 36). Together, our data indicate that ABL/PI5P treatment may rescue the impaired bacterialicidal mechanisms of macrophages with dysfunctional CFTR by restoring phagosome acidification and enhancing ROS production. Finally, the effect was specific to PI5P, as ABL loaded with PI3P, a second lipid messenger involved in membrane trafficking and autophagy (12), did not result in any modulation of intracellular *P. aeruginosa* killing (18).

The *ex vivo* analysis of MDM from CF patients indicated the presence of two groups of patients that we classified as “impaired” or “controller,” based on their different capability to control *in vitro* *P. aeruginosa* infection (bacterial replication index >1 or <1, respectively). It has been reported that host-genotypic traits have a critical role in the outcome of *P. aeruginosa* infection (37). In particular, the host susceptibility and the severity of infections caused by *P. aeruginosa* also depend upon a wide complex arrangement of genes, which is highly variable among immunocompromised individuals, including CF patients (38). Changes in clinical disease signs are mostly dependent on secondary gene variants that affect the outcome of the infection. These genes are identified as “modifier genes,” some of which play a role in innate immune response (39–41). Importantly, we observed that ABL/PI5P *ex vivo* treatment of macrophages induced a significant intracellular bacterial killing in the “impaired” group, highlighting the immunostimulant properties of ABL/PI5P, which
restores the dysfunctional CF bactericidal response. On the contrary, the same treatment did not further increase the intracellular *P. aeruginosa* killing of macrophages from the “controller” group or from functional MDM by healthy donors. In agreement with these observations *in vitro*, we did not observe variations in terms of pulmonary bacterial burden by healthy donors. Together, these data support the hypothesis that ABL/PI5P treatment has no general and broad-spectrum immunoenhancing effect, but it is endowed with the potential to rescue impaired microbicidal innate immune function.

Airway inflammation is a hallmark of CF disease that leads to the decline in lung function (26) and is characterized by elevated levels of NF-κB activation and proinflammatory cytokine and chemokine production (30), resulting in chronic inflammation, neutrophil recruitment, and progressive airway destruction. It is still a matter of debate on whether excessive inflammation in CF is the result of either underlying chronic bacterial infection(s) in the lungs or of exaggerated NF-κB signaling (42). Results reported herein show that the levels of NF-κB activation increase in macrophages following *P. aeruginosa* infection, and such an increase is significantly higher following pharmacological inhibition of CFTR, both in uninfected and in infected macrophages in comparison with the control cells. However, despite higher basal NF-κB activation in the cells with pharmacologically inhibited CFTR, differences in TNF-α, IL-1β, IL-6 levels were observed in *P. aeruginosa*-infected macrophages only, suggesting that the presence of the pathogen is necessary to NF-κB-dependent proinflammatory cytokine production. These results support the hypothesis of a higher, NF-κB dependent, predisposition to a hyperinflammatory response by the macrophages with dysfunctional CFTR, which requires the presence of bacterial pathogens to over-express proinflammatory cytokines (30).

PS exposure at the outer surface of the cell membrane is a physiologically relevant signal for phagocytic cells, for which it represents the “eat me” signal provided by apoptotic bodies generated by cells undergoing apoptosis (43). This process is an anti-inflammatory/tolerogenic signal with immunomodulatory properties (44), which have been previously exploited for the treatment of autoimmune diseases (45). Furthermore, PI5P is involved in the activation of PI3K/Akt pathway that is crucial in restricting proinflammatory and promoting anti-inflammatory response (32, 46). The results reported herein support the anti-inflammatory and protolerogenic role of PS and PI5P even when they are delivered as a single liposome formulation. Based on these *in vitro* experimental results, we moved to the *in vivo* murine model of chronic *P. aeruginosa* infection and assessed the effects of ABL/PI5P treatment in terms of lung KC, MIP-2, JE and TNF-α production, leukocyte infiltrates, and pulmonary bacterial burden. Results show that in ABL-/PI5P-treated mice, the number of BALF neutrophils was significantly reduced, and such reduction paralleled with KC and MIP-2 levels, whereas any reduction of TNF-α and JE levels was not observed. The different results obtained following *in vitro* and *in vivo* infection, in terms of TNF-α production, may be explained by the activation of different cell types, such as antigen-specific Th1, Th17,
and Th22 cells that may be involved and recruited to the lung during in vivo infections (47). Anti-inflammatory therapies, such as corticosteroids or biotechnologicals, may cause immunosuppression, which in turn is associated with the emergence of latent or opportunistic infections, and for this reason, they are often administered in combination with antibiotics (48). A clinical study to investigate the leukotriene B(4) (LTB(4)-receptor antagonist BIIL284 in CF patients was prematurely terminated due to a significant increased risk of adverse pulmonary events (49). Subsequent in vivo models showed that decreased airway neutrophils induced lung proliferation and severe bacteremia in a murine model of P. aeruginosa lung infection (50), indicating that strategies that interfere with neutrophil mechanisms have to be implemented with great caution. Of note, the reduction in inflammatory reactions in the lung of infected mice treated with ABL/PI5P was not associated with a significant increase in bacterial burden, suggesting that the in vivo administration of ABL/PI5P, by activating the macrophage component, may compensate for the reduction in neutrophil response and may have a therapeutic value also in critical conditions such as neutropenia.

Altogether, our data support the possibility that PI5P conveyed by ABL represents a novel therapeutic strategy devoid of immunosuppressive side effects, aimed at improving the efficiency of phagocytosis of mononuclear phagocytes and at reducing the

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**FIGURE 5** | ABL/PI5P stimulation modulates NF-κB and cytokine production in MDM with pharmacologically inhibited CFTR. MDMs were treated or not with INH172, infected or not with MDR P. aeruginosa (Pa, BAA-2113 strain), and then stimulated or not with ABL/PI5P for 2 h. Thereafter, cells were lysed (A) or supernatants were collected (B–F), and both were stored at −20°C until analysis. (A) Cell lysates were analyzed by NF-κB p65 (Total/Phospho) Human InstantOne™ ELISA kit, and results are shown as the ratio between phosphorylated and total NF-κB p65. The production of TNF-α (B), IL-1β (C), IL-6 (D), IL-10 (E), and TGF-β (F) was analyzed by ELISA. The results are shown as mean ± standard deviation of the values obtained from triplicate of each conditions and are representative of experiments with cells from at least three different donors. *p < 0.05; **p < 0.01; ***p < 0.001 one-sided t test.
FIGURE 6 | ABL/PISP treatment modulates KC and MIP-2 production in a murine model of MDR P. aeruginosa chronic lung infection. C57Bl/6NcrBR mice were infected with MDR P. aeruginosa (RP73 strain) and then treated with ABL/PISP (n = 16) or vehicle (n = 16), as described in the Material and Methods section. At day 6 post-infection, mice were sacrificed. Levels of KC (A), MIP-2 (B), TNF-α (C), and JE (D) upon treatment with ABL/PISP or vehicle in the supernatants of lung homogenates were measured by ELISA. Data are shown as mean values ± standard error. The data are pooled from two independent experiments. Statistical analysis was performed by using the two-sided Mann-Whitney test. Statistical significance is indicated: *p < 0.05. Outlier data, identified by Grubbs’ test, were excluded by the analysis.

FIGURE 7 | ABL/PISP treatment reduces neutrophilic recruitment in a murine model of MDR P. aeruginosa chronic lung infection. C57Bl/6NcrBR mice were infected with MDR P. aeruginosa (RP73 strain) and then treated with ABL/PISP (n = 16) or vehicle (n = 16), as described in the Materials and Methods section. At day 6 postinfection, mice were sacrificed, BALF was collected, and lungs were excised, homogenized, and plated onto TSA to determine bacterial burden. Counts of total number of cells (A), neutrophils (B), and macrophages (C) were performed in BALF. (D) The bacterial burden and was assessed by CFU assay. Data are shown as mean values ± standard error. The data are pooled from two independent experiments. Statistical analysis was performed by using two-sided Mann-Whitney test. Statistical significance is indicated: *p < 0.05. Outlier data, identified by Grubbs’ test, were excluded by the analysis.
damage of chronic inflammation. In conclusion, the ABL/-PI5P-based immunomodulatory strategy may represent an additional therapeutic tool in the fight against MDR opportunistic pathogens, such as P. aeruginosa, with the added value of the capacity to reduce the hyperinflammatory reactions in chronic lung infections that are particularly invalidating in CF patients.

**DATA AVAILABILITY STATEMENT**

All datasets generated for this study are included in the article/Supplementary Material.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by Ethics Committee of “Bambino Gesù” Children’s Hospital, Rome, Italy. Ethics approval #738/2017. Written informed consent to participate in this study was provided by the participants’ legal guardian/next of kin. The animal study was reviewed and approved by Institutional Animal Care and Use Committee (IACUC) #733.

**AUTHOR CONTRIBUTIONS**

VL, RN, AB, and MF contributed to the conception and design of the study. NP, FDS, AR, SR, IDF, AH, and FC contributed to data acquisition. NP, MMDA, RN, AB, and MF participated in data analysis and manuscript writing. All authors contributed to the article and approved the submitted version.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: [https://www.frontiersin.org/articles/10.3389/fimmu.2020.532225/full#supplementary-material](https://www.frontiersin.org/articles/10.3389/fimmu.2020.532225/full#supplementary-material)
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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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