Optimization of nitric oxide donors for investigating biofilm dispersal response in \textit{Pseudomonas aeruginosa} clinical isolates

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Received: 2 July 2020 / Revised: 13 August 2020 / Accepted: 23 August 2020 / Published online: 31 August 2020
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\textbf{Abstract}
\textit{Pseudomonas aeruginosa} biofilms contribute heavily to chronic lung infection in cystic fibrosis patients, leading to morbidity and mortality. Nitric oxide (NO) has been shown to disperse \textit{P. aeruginosa} biofilms in vitro, ex vivo and in clinical trials as a promising anti-biofilm agent. Traditional NO donors such as sodium nitroprusside (SNP) have been extensively employed in different studies. However, the dosage of SNP in different studies was not consistent, ranging from 500 nM to 500 μM. SNP is light sensitive and produces cyanide, which may lead to data misinterpretation and inaccurate predictions of dispersal responses in clinical settings. New NO donors and NO delivery methods have therefore been explored. Here we assessed 7 NO donors using \textit{P. aeruginosa} PAO1 and determined that SNP and Spermine NONOate (S150) successfully reduced > 60% biomass within 24 and 2 h, respectively. While neither dosage posed toxicity towards bacterial cells, chemiluminescence assays showed that SNP only released NO upon light exposure in M9 media and S150 delivered much higher performance spontaneously. S150 was then tested on 13 different cystic fibrosis \textit{P. aeruginosa} (CF-PA) isolates; most CF-PA biofilms were significantly dispersed by 250 μM S150. Our work therefore discovered a commercially available NO donor S150, which disperses CF-PA biofilms efficiently within a short period of time and without releasing cyanide, as an alternative of SNP in clinical trials in the future.

\textbf{Key points}
- S150 performs the best in dispersing \textit{P. aeruginosa} biofilms among 7 NO donors.
- SNP only releases NO in the presence of light, while S150 releases NO spontaneously.
- S150 successfully disperses biofilms formed by \textit{P. aeruginosa} cystic fibrosis clinical isolates.

\textbf{Keywords} Nitric oxide · \textit{Pseudomonas aeruginosa} · Biofilm · Cystic fibrosis · Chemiluminescence

\section*{Introduction}

The major cause of morbidity and mortality in cystic fibrosis (CF) patients is the chronic bacterial colonization of patients’ lungs and airways leading to pulmonary dysfunction and infection (Gilligan 1991; Govan and Deretic 1996). When bacteria invade healthy individuals, opportunistic pathogens that overcome mucociliary clearance can be targeted by phagocytic cells and specific opsonizing antibodies (Govan and Deretic 1996). However, in CF patients, the dehydrated surface liquid on respiratory epithelium results in defective mucociliary clearance and frustrated phagocytosis due to the impaired opsonisation process, hence contributing to the chronic colonization (Govan and Deretic 1996). \textit{Pseudomonas aeruginosa} has been well recognized as the most commonly found and important pathogen in progressive and severe CF lung disease. CF \textit{P. aeruginosa} clinical isolates frequently show higher levels of persister cells and antibiotic resistance (Saiman et al. 1996; Mulcahy et al. 2010). Furthermore, \textit{P. aeruginosa} tend to form aggregates/biofilms in vivo, leading to a much higher tolerance to treatments, driven by both the protective extracellular polymeric...
substances (EPS) produced by bacteria and local host environment (Ciofu and Tolker-Nielsen 2019). Once biofilms are established, they are almost impossible to eradicate (Hoiby et al. 2005).

One promising strategy is to combine biofilm dispersal agents and conventional antibiotics, where bacterial cells become more susceptible once reversed back to planktonic form. Nitric oxide (NO) was discovered to be an anti-biofilm signaling molecule and the mechanisms are still under investigation (Barraud et al. 2006). In P. aeruginosa, biofilm phenotypes have been associated with an important secondary messenger, cyclic-di-GMP (c-di-GMP), which contributes to a myriad of physiological changes and the establishment of biofilms. A decrease in intracellular c-di-GMP level promotes motile mode of growth of bacteria and triggers biofilm dispersal (Ute Römling et al. 2013). NO was demonstrated to reduce c-di-GMP level by stimulating the activities of phosphodiesterases responsible for the hydrolysis of c-di-GMP, upregulate genes involved in motility and downregulate those related to the expression of adhesins and virulence factors (Rinaldo et al. 2018; Barraud et al. 2009b). Although NO can be endogenously generated by P. aeruginosa, which facilitates biofilm dispersal at the late stage of biofilm life cycle, exogenously added NO significantly accelerates the procedure and can also prevent the early attachment. Previous studies reported that the efficacies of conventional antibiotics towards established biofilms were significantly enhanced when NO was applied as a dispersal agent (Barraud et al. 2006; Howlin et al. 2011; Soren et al. 2019). NO gas has also been applied in clinical trials, where it increased the efficacy of conventional antibiotics in CF lung infection treatment (Howlin et al. 2017; Cathie et al. 2014). Therefore, NO has been regarded as a putative anti-biofilm adjunctive therapy. Conventional NO donors such as sodium nitroprusside (SNP), S-Nitrosothiols (RSNOs) and Diazoniumdiolates (NONOates) have been applied as dispersal agents against biofilms formed by different clinically relevant bacterial species such as P. aeruginosa, Escherichia coli, Neisseria gonorrhoeae, Staphylococcus aureus and Staphylococcus epidermidis (Barraud et al. 2006; Barraud et al. 2009a; De La Fuente-Núñez et al. 2013; Barnes et al. 2013; Sulemankhil et al. 2012; Falsetta et al. 2009; Jardeleza et al. 2011). Along with these, many studies were also carried out to explore novel methods for NO delivery, such as incorporating NO donors into nanoparticles and polymer coating (Sadrearhami et al. 2017; Duong et al. 2014a; Nablo and Schoenfisch 2003; Nablo et al. 2005). However, more often than not these studies tested different concentrations of NO donors towards early stage biofilms formed by type strains, or did not specify optimal treatment time which is crucial for the interpretation of data from young biofilms to distinguish between prevention and dispersal (Barraud et al. 2009a; Barnes et al. 2013; Sadrearhami et al. 2017; Duong et al. 2014a; Shen et al. 2019; Duong et al. 2014b; Zhu et al. 2018; Marvasi et al. 2014). To this end, we systematically compared the optimal concentrations and treatment time of 7 NO donors for triggering biofilm dispersal using P. aeruginosa PAO1, including SNP, S-nitroso-glutathione (GSNO), S-nitroso-N-acetyl-DL-penicillamine (SNAP), 1-(hydroxy-NO-azoxy)-L-proline (PROLI NONOate), 6-(2-hydroxy-1-methyl-2-nitrosohydrazino)-N-methyl-1-hexanamine (MAHMA NONOate), (Z)-1-[N-[3-aminopropyl]-N-[4-(3-aminopropylammonio)butyl]-amino]diazene-1-ium-1,2-diolate (Spermine NONOate) and diethylammonium (Z)-1-(N,N-diethylamino)diazen-1-ium-1,2-diolate (DEA-NONOate). The selection was based on different categories of NO donors previously reported to be used in laboratories, animal experiments or clinical settings with different NO release mechanisms. While SNP has been traditionally and widely applied, the mechanism for NO release is very complicated depending on conditions and involves three stages (Smith and Dasgupta 2002). GSNO and SNAP belong to S-nitrosothiols, a class of NO donor releasing NO and nitrosonium (NO⁺) spontaneously from the moiety (-SNÖ) (Napoli and Ignarro 2003). Both GSNO and SNAP were initially investigated for their role as antiplatelet agents in cardiovascular system, as S-nitrosothiols do not appear to engender vascular tolerance (Belcastro et al. 2017; Brisbois et al. 2013). In contrast, NONOates, i.e. diazeniumdiolates, consist of a diolate group [N(O-)N=O] bound to a nucleophile adduct via a nitrogen atom (Maragos et al. 1991). Although not yet approved for clinical use so far, NONOate is one of the most investigated NO donors due to its capability to release two moles of NO per mole of donor at physiological conditions, showing outstanding clinical application prospects (Yang et al. 2018; Li et al. 2020). As such, four different NONOates previously investigated in other biofilm studies (Barnes et al. 2013; Barnes et al. 2015; Zhu et al. 2018; Marvasi et al. 2014) were chosen in this study. After a high-throughput screening, both 250 μM SNP and Spermine NONOate (S150) showed great dispersal efficacies after 24 h and 2 h, respectively. However, by employing a highly sensitive chemiluminescence detection method, we confirmed that S150 exhibited better performance in releasing NO and is more suitable for testing biofilm dispersal response to NO.

As CF-PAs can undergo genetic adaption catalysed by hypermutation in chronic infection lungs (Mena et al. 2008; Bianconi et al. 2019; Caçador et al. 2018; Winstanley et al. 2016), it is suspected that some CF-PA isolates may contain mutations that lead to higher tolerance to NO. Using 250 μM S150, we tested the NO response of 72-h biofilms in microtiter plates formed by 13 different CF-PA isolates and most strains were successfully dispersed. In summary, our study showed that (1) S150 is superior to SNP, which can be applied in wider settings as it does not require light to release NO; (2) S150 can disperse biofilms formed by genotypically different CF-PA strains, indicating its potential clinical applications.
Materials and methods

Ethics for cystic fibrosis patient sputum collection

Sputum samples from 72 patients with CF were obtained by CF physiotherapist-assisted sample expectoration following Good Clinical Practice guidelines (ICH) (Blau et al. 2014; Aaron et al. 2004). All sampling protocols and procedures were approved by UK NHS Research Ethics Committee (South Central – Hampshire A Research Ethics Committee, Reference 08/H0502/126, Mechanisms of lung infection and inflammation in respiratory disease). Informed consent was obtained from all subjects or, if subjects were under 18, from a parent and/or legal guardian.

Bacterial strains and culture conditions

Bacterial strains used in this study are listed in Supplementary Table S1. All bacterial overnight cultures were grown in lysogeny broth (LB) medium for 15 h at 37 °C. For CF-PA isolation, sputa samples were digested using Mucolyse™ Sputum Digestant (Pro-Lab Diagnostics, UK) for 15 mins at 37 °C, followed by culture on *P. aeruginosa*-specific cetrimide agar (Sigma-Aldrich, UK). Multiplex PCR was used to confirm *P. aeruginosa* as previously described by De Vos et al. (1997).

Preparation of NO donor/NO scavenger solution

NO donors with different half-lives in pH 7.4 buffer tested in this study are listed in Table 1. Sodium nitroprusside (SNP), S-nitroso-N-acetyl-DL-penicillamine (SNAP), S-nitrosoglutathione (GSNO), PROLI NONOate and Spermine NONOate (S150), diethylamine NONOate sodium salt hydrate (DEA NONOate) and carboxy-PTIO potassium salt (PTIO) were purchased from Sigma Aldrich, UK. SNP and RSNOs stock solutions were prepared in phosphate saline buffer (pH 7.4), with SNP prepared and kept in dark. NONOates stock solutions were prepared in 0.01 M NaOH. All stock solutions were filter sterilized and diluted into fresh M9 media on ice. During preparation procedures, all solutions were kept on ice before use and SNP was kept in the dark. All treatments were conducted at 37°C, with SNP exposed to light.

Batch cultured biofilms

For microtiter plate–based biofilm assays, 100 μl of each culture in M9 medium (OD600nm ~ 0.01) was inoculated into each well. Microtiter plates were incubated statically for 24 h, and biofilms stained with 0.1% (v/v) crystal violet after two washes, dissolved in 30% (v/v) acetic acid. Crystal violet staining was quantified at a wavelength of 584 nm.

For microscopic biofilm examination, 3 ml of culture in M9 medium (OD600nm ~ 0.01) was inoculated into a MatTek plates (P35G-1.5-14-C). Plates were shaken at 50 rpm and biofilms stained with LIVE/DEAD® BacLight (Invitrogen, UK) and examined by confocal laser scanning microscopy (CLSM). A wavelength of 488 nm was used for SYTO-9 and 561 nm was used for propidium iodide excitation. At least 3 image stacks were taken from random locations in each MatTek plate. Total biomass for biofilms in each micrograph was calculated by software COMSTAT (Heydorn et al. 2000).

For NO donor screening assay using PAO1 biofilms, NO donors were added to the 24-h pre-established biofilms in microtiter plates and then incubated for a further 1, 2, 4, 6, 8, 12 or 24 h as treatment. For NO-induced dispersal in PAO1 and CF-PA biofilms, S150 was added to 72-h pre-established biofilms in both microtiter plates and MatTek plates and then incubated at 37 °C for 2 h to trigger dispersal.

Bactericidal test

Bactericidal test method was modified from Barnes et al. (2013). Briefly, 1 ml overnight cultures were centrifuged at 4000 × g for 10 mins to harvest the cells in Eppendorf tubes, washed twice in sterile PBS and re-suspended in 1 ml M9. Serial dilutions of the cells were made to approximately 10^6 CFU/ml in M9. SNP and S150 donor stock solutions were added into the culture making a desired final concentration. Cells were then incubated at 37 °C for 2 h or 24 h in M9 before determining the final CFU following a modified Miles et al. (1938).

NO release quantification using chemiluminescence

NO gas released by SNP and S150 in M9 medium was detected in a chemiluminescence CLD 88Y NO analyser (EcoPhysics, Durnten, Switzerland) with synthetic air (99.99999% BOC) as an inert carrier. The photomultiplier detects emissions above 600 nm, and the NO concentration was calculated from the emitted intensity against a calibration standard using NaNO2 as previously reported (standard curve at 250, 375, 500, 750 and 1000 pmol NO, Supplementary Fig. S1) (Piknova and Schechter 2011). Tracings were recorded at 4-Hz frequency using PowerChrom® (eDAQ Pty LtD).

| Table 1 | NO donors tested in this study and their half-lives at 37°C, pH ~ 7.4 |
|-----------------|-----------------|
| SNP             | <2 mins         |
| SNAP            | 6 h             |
| GSNO            | 1–3 h           |
| PROLI NONOate   | 1.8 s           |
| MAHMA NONOate   | 1 min           |
| Spermine NONOate (S150) | 39 min         |
| DEA-NONOate     | 2 min           |
Australia). Data were plotted using Origin 9, calculating areas to quantify total NO release within 2 h, comparing NO donor samples against standards.

**Statistical analyses**

All assays were assessed using the two-tailed Student’s T test. Statistical significances were $P < 0.05$ for all measurements reported unless otherwise stated. Statistics and graphs were produced using GraphPad Prism.

**Results**

**SNP and S150 successfully remove PAO1 biofilms**

The efficacies of 7 donors in M9 medium at 9 concentrations (1 μM, 2.5 μM, 5 μM, 10 μM, 25 μM, 50 μM, 100 μM, 250 μM and 500 μM) and 7 treatment time periods (1 h, 2 h, 4 h, 6 h, 8 h, 12 h, 24 h) were systematically tested at 37°C in microtiter plates. All data were shown in Supplementary Fig. S2–8. Among all conditions, 24-h 250 μM SNP treatment and 2-h 250 μM S150 treatment were determined as optimal (> 60%) for triggering *P. aeruginosa* dispersal (Fig. 1). 500 μM dosage for both compounds performed comparably with 250 μM. To avoid overexposure of NO, 250 μM was selected for further tests.

**S150 exhibits higher NO release efficiency than SNP**

Gas phase chemiluminescence is a sensitive method to record and quantify the precise NO release, which was therefore employed in this study to compare the difference between SNP and S150. Due to the 1 ppm maximum detection limit of CLD 88Y, NO released from 5 μM S150 and SNP was quantified in M9 media at 37°C across ~ 1.5 h timeframe as shown in Fig. 2a and b. Results showed that at 37°C, S150 spontaneously released NO upon contact with the medium, while SNP steadily released low amount of NO. As 1 molecule of S150 releases 2 molecules of NO and SNP only releases 1, the predicted NO release from S150 is twice more than the same amount of SNP. However, the efficiency of S150 within 1.5 h was 68.4 ± 8.1%, while SNP was only 29.2 ± 2.2% (Fig. 2b). Therefore, it can be concluded that at the same concentration, S150 is more effective in releasing NO within a relatively short time frame. As SNP degradation is associated with light exposure, a PHOTONIC PL3000 device (maximum light intensity 26 Mlx, colour temperature 3250 K) was applied for a constant cold light source without disturbing the incubation temperature. 500μM SNP was chosen for more obvious releasing curves, and foil paper was used to wrap the whole system when light was forbidden in the tests. From Fig. 2c, it can be concluded that SNP effectively released NO when light was present. However, SNP stopped releasing NO immediately after the light exposure was withdrawn, confirming the necessity of light for SNP as a NO donor. Even at a concentration of 500 μM, the peak NO concentration released from NO only reached 0.8 ppm, while S150 immediately released 0.35 ppm at 5 μM. Therefore, S150 was chosen for its spontaneous, more consistent and efficient performance without the production of cyanide in further assays.

**S150 triggers biofilm dispersal through NO, not cytotoxicity**

Previous studies have shown that NO scavenger (PTIO) can abolish the dispersal effect from SNP (Barraud et al. 2006; Howlin et al. 2017). Here, we also tested whether biofilm dispersal triggered by S150 was due to side effects or NO released from the donor using NO scavenger PTIO. Usually a higher concentration of PTIO than that of NO donor was added to ensure all NO could be scavenged (Barraud et al. 2006). As shown in Fig. 3a, 250 μM S150 triggered dispersal in 2 h, while PTIO abolished biofilm dispersal totally (biomass increased by 33.4 ± 9.7%, $P < 0.001$). For control + PTIO, a higher increase occurred (47.2 ± 11.3%). The colour of combination of S150 and PTIO turned yellow while control added in PTIO remained purple in Fig. 3b, indicating S150 was reacting with PTIO. Thus, PTIO prevented NO release from S150, and the increase of biofilm might be due to
excessive PTIO inhibited NO release from nitrite reductase (Barraud et al. 2006).

After confirming it was NO triggering biofilm dispersal, we next determined if the removal of biofilms was due to the bactericidal effect. The toxicity of NO was tested for planktonic PAO1 cells in M9 media at 37°C using an initial inoculum of CFU $\sim 1 \times 10^4$/ml. Planktonic cultures were treated for 2 h using S150, and 24 h using SNP according to their optimal dispersal time. Figure 4 showed that both SNP and S150 surprisingly enhanced planktonic cells’ number at higher concentrations. Theoretically 250 $\mu$M S150 and 500 $\mu$M SNP should release the same amount of NO, and they increased CFU by 2.4 ± 0.6-fold ($P < 0.001$) and 46 ± 8-fold ($P < 0.001$), respectively. Whether it was NO per se or the breakdown products from donors that enhanced the growth remains unknown. Nevertheless, the long treatment time (24 h) needed for SNP contributed to much larger errors in...
S150 was therefore again proven to be superior. S150 successfully disperses biofilms formed by CF-PAs. As CF-PA strains often go through genetic adaptation in chronically infected lungs, we tested whether the biofilms formed by different clinical isolates can also be dispersed by optimized NO donor. However, most clinical isolates exhibit much slower growth rates and biofilm establishment compared with type strain PAO1. Previous microtiter plate screening in the lab showed that most CF-PA isolates formed the thickest biofilms at 72 h, after which the automatic dispersal stage began (data not shown). S150 treatment was also carried out on 24-h, 48-h, 72-h, 96-h, and 120-h PAO1 biofilms, and the results showed that the biomass reduction of 72-h PAO1 biofilms after S150 treatment was around 40–45%. As such, to test S150 on the most robust CF-PA biofilms, a 72-h incubation time was chosen. As shown in Fig. 5, 72-h biofilms formed by 12 out of 13 CF-PA strains can be successfully dispersed by S150 within 2 h despite the different response compared with PAO1, with only PA58 being an exception showing tolerance to NO. Microtiter plate results were re-enforced by selected CLSM micrographs in Fig. 6. PAO1 WT, PA10, PA21, PA26, PA30, PA58, and PA68 were selected due to their substantial multilayer biofilm formation in MatTek plates. However, PA58 biofilms formed in MatTek plate also showed significant dispersal (67.5 ± 21% biomass reduction) here. We speculate that the discrepancy was due to the fact that CV staining is a relatively rough method, staining biofilm rings formed at both air-liquid surface and the bottom of each well containing cells and EPS. In contrast, confocal microscopic method measured just total cell mass attached to the bottom of each well. Nevertheless, S150 successfully dispersed biofilms formed by most CF-PA strains tested in this study and thus may also be efficient towards a variety of P. aeruginosa strains with different sources and genetic backgrounds.

Data interpretation due to much higher increase in CFU, and S150 was therefore again proven to be superior.

Discussion

Low-dose nitric oxide has been repeatedly reported to disperse biofilms formed by different species. Traditionally used SNP was reported to reduce biofilms formed by P. aeruginosa, Vibrio cholerae, Serratia marcescens, Escherichia coli, Bacillus licheniformis and Neisseria gonorrhoeae, with a working concentration ranging from 25 to above 500 nM. (Barraud et al. 2009a; De La Fuente-Núñez et al. 2013; Barnes et al. 2013; Sulemankhil et al. 2012; Falsetta et al.
Fig. 6  a Selective confocal laser scanning microscopic micrographs of 72-h PAO1, PA10, PA21, PA26, PA30, PA58 and PA68 biofilms grown at 37 °C with/without S150 treatment. Scale bar = 50 μm. b 72-h CF-PA biofilms with 250 μM S150 treatment. Biomass reduction was analysed by COMSTAT.
A higher concentration of SNP (1–10 μM) was required to disperse biofilms formed by *Fusobacterium nucleatum* and *Staphylococcus epidermidis* (Barraud et al. 2009a). The maximum removal percentage within 24 h ranged from 38.1% in *E. coli* to 92.8% in *B. licheniformis*, and the average removal rate for different species was 63%. Therefore, SNP is effective in dispersing biofilms formed by different bacteria, although each species shows distinct responses. However, when comparing data on SNP-induced *P. aeruginosa* biofilms from various studies, different groups reported using different concentrations of SNP ranging from 500 nM to 500 μM (Barraud et al. 2006; Howlin et al. 2011; Barraud et al. 2009a; Chua et al. 2014; Barraud et al. 2009b; Roy et al. 2012). The discrepancy might be due to the fact that SNP does not release NO unless after photolysis or the addition of reducing agents (Kowaluk et al. 1992), resulting in the differences of its efficacy under different laboratory light sources. Furthermore, by-products such as cyanide released from SNP may cause side effects to cells and misinterpretation of experimental results, especially when *P. aeruginosa* is a cyanogenic bacterium (Arnold et al. 1984; Cipollone et al. 2007).

Alternative NO donors with more consistent performances have also been explored. For instance, 200 μM and 1000 μM DETA-NONOate induced the biofilm dispersal of *Shewanella woodyi* and *Staphylococcus aureus*, respectively (Liu et al. 2012; Jardeleza et al. 2011). 50 μM and 100 μM DPTA NONOate reduced *Shewanella oneidensis* and *Vibrio harveyi* biofilms (Arora et al. 2015; Henares et al. 2013). Direct usage of gaseous NO also led to the dispersal of *Staphylococcus aureus*, *Acinetobacter baumannii* and *Nitrosomonas europaea* (Sulemankhil et al. 2012; Schmidt et al. 2004). SNAP and GSNO were reported to decrease *P. aeruginosa* biofilms although less effectively than SNP (Barraud et al. 2006), while MAHMA NONOate and PROLI NONOate and Spermine NONOate exhibited higher performance against *P. aeruginosa* biofilms although less effectively than SNP (Barraud et al. 2006), while MAHMA NONOate, PROLI NONOate and SNAP and GSNO were reported to decrease *P. aeruginosa* biofilms (Barraud et al. 2006; Howlin et al. 2011; Barraud et al. 2009a; Chua et al. 2014; Barraud et al. 2009b; Roy et al. 2012). The discrepancy might be due to the fact that SNP does not release NO unless after photolysis or the addition of reducing agents (Kowaluk et al. 1992), resulting in the differences of its efficacy under different laboratory light sources. Furthermore, by-products such as cyanide released from SNP may cause side effects to cells and misinterpretation of experimental results, especially when *P. aeruginosa* is a cyanogenic bacterium (Arnold et al. 1984; Cipollone et al. 2007).

Gas phase chemiluminescence (CLD) results have shown that while the same molarity of SNP is expected to release 50% of NO less than S150, it actually released 78.7% less NO within 1.5 h (Fig. 2a and b). Furthermore, the decomposition of S150 started as soon as the compound was in contact with the solution, corresponding to its spontaneous NO release following the first-order kinetics (Ramamurthy and Lewis 1997). In contrast, SNP remained a constant but slow release state under normal room lighting. The lower efficacy of SNP was further confirmed by results in Fig. 2c, where SNP can only generate NO in the presence of intense light. Even so, 500 μM SNP only reached the maximum 0.8 ppm NO release peak compared with 0.35 ppm peak from 5 μM S150. As S150 can efficiently release NO with or without light, it is more suitable for different applications such as in patients.

While it was shown that S150 efficiently releases NO, Figs. 3 and 4 confirmed that the biofilm dispersal effect came from NO rather than any by-product or side effect. When NO scavenger was added, biofilm dispersal was abolished. Neither SNP nor S150 posed toxic effects towards planktonic cells, indicating the biofilm removal was not due to cytotoxicity. Interestingly, both NO donors significantly enhanced bacteria numbers without additional nutrients added into M9 media. As NO radical can be readily converted to nitrite and...
nitrate, and previous studies have shown that nitrate can support the growth of *P. aeruginosa* in anaerobic/anoxic environment (Fang et al. 2013; Yoon et al. 2011; Line et al. 2014), the additional nitrite or nitrate might have contributed to the increased growth of planktonic *P. aeruginosa* cells. Alternatively, as the backbone of S150 is rich in carbon and nitrogen, while cyanide can be degraded and utilized as nutrients (Cipollone et al. 2007; Knowles 1988), the chemical breakdown products from SNP and S150 may have acted as additional nutrients for cell growth during the treatment. However, the precise mechanism of S150/SNP-induced growth increase is yet to be elucidated. Some previous studies used ‘increased CFU/OD/turbidity in the effluents’ after SNP treatment as the indicator of biofilm dispersal into planktonic forms (Barraud et al. 2006; Howlin et al. 2017; Chua et al. 2014). Depending on the treatment period, this measurement may not be accurate enough to reflect the dispersal rate, as the donors may have enhanced the planktonic growth at the same time. In summary, 250 μM S150 is the optimal NO donor dosage among all compounds tested here for triggering *P. aeruginosa* biofilms. Additionally, due to its desirable decomposition half-life (39 mins at pH 7.4, 37°C) compared with other short half-life NONOates such as MAHMA NONOates and PROLI NONOates, it is easier to prepare and control under different conditions.

Darling and Evans (2003) reported that NO production in vivo reduced *P. aeruginosa* adherence to human bronchial epithelial cells and enhanced the killing of internalized bacteria. However, various publications indicated that exhaled NO from CF patients is reduced compared with that produced by normal patients (Elphick et al. 1999; Jöbsis et al. 2000; Mhanna et al. 2001). In healthy individuals, inducible NOS (iNOS) is expressed maximally following an inflammatory stimulus and produces large, micromolar scale of NO (Darling and Evans 2003). However, in CF patients’ airways with chronic severe inflammation, the amount of exhaled NO is not increased and the expression of epithelial iNOS is reduced (Darling and Evans 2003). Therefore, CF-PA biofilms developed in chronic lung infection may be constantly exposed to sublethal dosage of NO. A recent study stated that pre-treated biofilms with non-dispersing concentrations of NO showed much increased tolerance to NO (Zhu et al. 2018). As CF-PAs usually exhibit much genetic variation compared with type strains due to in vivo adaption, the repeated NO exposure may lead to mutations resulting in higher tolerance to NO. Therefore, we suspected that the biofilms formed by CF-PAs isolated from sputum samples could develop tolerant to NO. From Figs. 5 and 6, it can be concluded that biofilms formed by different CF-PAs sampled from non-familiar patients were successfully dispersed by S150, regardless of their total biomass. Our data suggested that S150 can potentially prevent biofilm formation or disperse pre-established biofilms in clinical settings by direct application or incorporation into medical devices such as bone cement, dermal fillers and wound dressing. It may also be applied in combination with antibiotics to increase the susceptibility of cells encased in biofilms.

Acknowledgements The authors would like to thank Prof. Martin Feelisch for the generous offer of CLD facilitates in the Southampton General Hospital and Dr. Diogo Silva for setting up CLD and detailed explanations. We would also like to thank Dr. Robert Howlin and the nurse team for the provision of all the CF-PA clinical isolates, Dr. Odel Soren for the fruitful discussion about experimental designs and data interpretations, as well as Dr. Catherine Bryant for proofreading the manuscript.

Author contributions JSW and YMC designed the study. YMC conducted experimental work. YMC drafted the manuscript. JSW proofread and corrected the manuscript. Both authors approved the final version.

Funding information This work was supported by BBSRC and Innovate UK grant to National Biofilms Innovation Centre (NBIC) and JSW.

Compliance with ethical standards
Conflict of interest The authors declare that they have no competing interests.

Ethical approval All sampling protocols and procedures were approved by UK NHS Research Ethics Committee (South Central – Hampshire A Research Ethics Committee, Reference 08/H0502/126, Mechanisms of lung infection and inflammation in respiratory disease). Informed consent was obtained from all subjects or, if subjects were under 18, from a parent and/or legal guardian.

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