Response to Gaseous NO\textsubscript{2} Air Pollutant of *P. fluorescens* Airborne Strain MFAF76a and Clinical Strain MFN1032

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Human exposure to nitrogen dioxide (NO\textsubscript{2}), an air pollutant of increasing interest in biology, results in several toxic effects to human health and also to the air microbiota. The aim of this study was to investigate the bacterial response to gaseous NO\textsubscript{2}. Two *Pseudomonas fluorescens* strains, namely the airborne strain MFAF76a and the clinical strain MFN1032 were exposed to 0.1, 5, or 45 ppm concentrations of NO\textsubscript{2}, and their effects on bacteria were evaluated in terms of motility, biofilm formation, antibiotic resistance, as well as expression of several chosen target genes. While 0.1 and 5 ppm of NO\textsubscript{2} did not lead to any detectable modification in the studied phenotypes of the two bacteria, several alterations were observed when the bacteria were exposed to 45 ppm of gaseous NO\textsubscript{2}. We thus chose to focus on this high concentration. NO\textsubscript{2}-exposed *P. fluorescens* strains showed reduced swimming motility, and decreased swarming in case of the strain MFN1032. Biofilm formed by NO\textsubscript{2}-treated airborne strain MFAF76a showed increased maximum thickness compared to non-treated cells, while NO\textsubscript{2} had no apparent effect on the clinical MFN1032 biofilm structure. It is well known that biofilm and motility are inversely regulated by intracellular c-di-GMP level. The c-di-GMP level was however not affected in response to NO\textsubscript{2} treatment. Finally, NO\textsubscript{2}-exposed *P. fluorescens* strains were found to be more resistant to ciprofloxacin and chloramphenicol. Accordingly, the resistance nodulation cell division (RND) MexEF-OprN efflux pump encoding genes were highly upregulated in the two *P. fluorescens* strains. Noticeably, similar phenotypes had been previously observed following a NO treatment. Interestingly, an *hmp*-homolog gene in *P. fluorescens* strains MFAF76a and MFN1032 encodes a NO dioxygenase that is involved in NO detoxification into nitrites. Its expression was upregulated in response to NO\textsubscript{2}, suggesting a possible common pathway between NO and NO\textsubscript{2} detoxification. Taken together, our study provides evidences for the bacterial response to NO\textsubscript{2} toxicity.

Keywords: airborne, *Pseudomonas fluorescens*, nitrogen dioxide, biofilm, antibiotic sensitivity, motility, air pollution
INTRODUCTION

Most world-wide cities have serious air-quality problems, which have attracted attention in the past decade. One of the most common source of air pollution is engine emissions, which include, among other toxic molecules, the nitrogen oxides (NO\(_2\); reviewed in Sher, 1998; Skalska et al., 2010). The general term NO\(_x\) includes nitric oxide (NO) and nitrogen dioxide (NO\(_2\)). NO in turn is able to damage bacterial cells interacting with bacterial proteins (McLean et al., 2010; Laver et al., 2013) and DNA (Tamir et al., 1996; Burney et al., 1999) either directly, or via formation of reactive nitrogen species (RNS), causing alterations in bacterial metabolism, among which respiration, and homeostasis. As a result, bacteria have developed specific NO detoxification pathways and defense mechanisms (Cruz-Ramos et al., 2002; Flatley et al., 2005; Spiro, 2007). In order to counteract the NO-mediated respiratory arrest (Husain et al., 2008), the detoxification processes are completed in several bacteria by metabolism reprogramming (Auger et al., 2011; Auger and Appanna, 2015). NO was furthermore identified as a signaling molecule, which promotes the biofilm dispersion in various bacterial strains, including Pseudomonas aeruginosa (Barraud et al., 2009; Cutruzzola and Frankenberger-Dinkel, 2015) and P. putida (Liu et al., 2012). This molecule is also known to modulate bacterial antibiotic sensitivity, protecting bacteria from a wide range of antibacterial agents (Gusarov et al., 2009; McCollister et al., 2011; van Sorge et al., 2013), such as vancomycin and daptomycin (van Sorge et al., 2013). Contrary to NO, NO\(_2\) has a low solubility in water (Augusto et al., 2002). Thence NO\(_2\) in aqueous media concerned a few reports in the microbiological context. However, in natural environments NO is unstable and quickly oxidized to form NO\(_2\) (Skalska et al., 2010), considered as a major air pollutant. Its atmospheric level is ruled by European environmental commission and World Health Organization (INERIS, 2011; Reduction of pollutant emissions from light vehicles, 2015; WHO |Ambient (outdoor) air quality health, 2015). NO\(_2\) toxicity to human health is well documented and is known to increase cardiovascular diseases (Chaloulakou et al., 2008), or to aggravate respiratory symptoms especially in children (Pershagen et al., 1995; Chauhan et al., 1998). On the opposite, the stress promoted by NO\(_2\) was poorly evaluated on bacteria.

It is increasingly evident that the air is a biotic environment, containing bacteria as one of the major compounds of primary atmosphere aerosol particles (Burrows et al., 2009b; Després et al., 2012). Mean airborne bacterial concentrations can indeed be greater than 1 \(10^4\) cells m\(^{-3}\) (Bauer et al., 2002; Burrows et al., 2009a). Although unstable, the air microbiota is frequently constituted with members of Pseudomonas genus (Fang et al., 2007; Pearce et al., 2010; Després et al., 2012; Dybwad et al., 2012; Šantl-Temkiv et al., 2015). Among these highly versatile elements, the P. fluorescens strains are widely adaptable and distributed (Bodilis et al., 2004) in all major natural environments, including water (Bodilis et al., 2004), soil (Varivarn et al., 2013) and clouds (Ahern et al., 2007). Several P. fluorescens strains were also found to promote humans acute infections and were reported in clinical samples of immuno-compromised patients (Chapalain et al., 2008; Scales et al., 2014). All these properties make P. fluorescens a good model for further investigations of airborne bacteria.

We have investigated in previous studies the microbiota (bacteria, yeasts and fungi) of Rouen harbor terminal (France) (Morin et al., 2013). Thus, several P. fluorescens strains were isolated. Among them, the airborne P. fluorescens strain MFAF76a was characterized as a virulent strain, particularly its exoproducts against human epithelial pulmonary cells (Duclairoir Poc et al., 2014). The aim of this study is to investigate the physiological response of airborne P. fluorescens MFAF76a to NO\(_2\) as a marker of air pollution in terms of motility, biofilm formation and antibiotic resistance. This response was compared to that of the clinical strain P. fluorescens MFN1032 isolated from the sputum of a pneumonia-suffering patient (Chapalain et al., 2008). The parameters of bacterial NO\(_2\) exposure were adapted to mimic real-life air conditions. Thus, the two strains were exposed to gaseous NO\(_2\) at three concentrations: 0.1 ppm as an annual guideline value (WHO |Air quality guidelines - global update, 2005) 5 ppm as the threshold causing reversible effects on human health, and 45 ppm as a high NO\(_2\) concentration provoking irreversible effects (INERIS, 2011).

MATERIAL AND METHODS

Strains and Growth Conditions

Cyan Fluorescent Protein (CFP)-labeled P. fluorescens MFN1032 and MFAF76a were used in this study. The strains and plasmids are listed in Table S1. The 729-bp cfp gene, encoding the CFP, was extracted from pTetONCFPopt plasmid (Sastalla et al., 2009) using PstI and Xmal enzymes (NEB, Ipswich, USA). Then CFP cassette was separated by 1% agarose gel electrophoresis and purified with QIAquick PCR Purification Kit (Qiagen, Hilden, Allemagne). The pPSV35 vector (Rietsch et al., 2005) was digested using PstI and Xmal and purified using QIAquick PCR Purification Kit (Qiagen, Hilden, Allemagne). The CFP cassette was then cloned into the PstI and Xmal sites of the pPSV35 vector. The resulting pCFP vector was introduced into One Shot® TOP10 Chemically Competent E. coli (LMSM collection) by heat shock. After antibiotic selection of the clones (gentamycin 15 \(\mu\)g/mL), the transformation was confirmed by confocal laser scanning microscope (CLSM 710, ZEISS). The obtained plasmid was then extracted from E. coli using QIAprep Spin Miniprep Kit (Qiagen, Hilden, Allemagne) and introduced into P. fluorescens strains by electroporation. The transformants were selected in LB containing 15 \(\mu\)g/mL of gentamycin and fluorescence was assayed using CLSM.

Bacteria were grown at 28°C under limited agitation (180 rpm) in DMB (Davis Medium Broth) minimal medium with 2.16 g/L glucose as carbon source (Duclairoir-Poc et al., 2011). Overnight cultures were diluted (\(A_{580} = 0.08\)) in fresh DMB and grown to the end of exponential phase (\(A_{580} = 2, 13 \times 10^8\) CFU/mL). Bacterial cultures at the end of exponential growth phase (about 3 \(\times\) 10\(^7\) bacteria per filter) were transferred on cellulose nitrate membrane filter (0.45 \(\mu\)m, pore size 0.2 \(\mu\)m, diameter 47 mm, Sartorius Biolab Products,
Gottingen, Germany) and grown on DMB agar plates at 28°C for 4 h to obtain a single layer’s bacterial population. After 4 h of incubation, the cellulose membranes containing bacteria were placed on agar one-well dishes (size 127.8 × 85.5 mm, Thermoscientific Nunc, Roskilde, USA), which were directly transferred into the gas delivery device (Figure 1).

**Exposition to Nitrogen Dioxide**

In order to mimic the atmospheric conditions, bacterial NO₂ exposure was achieved in gas phase for 2 h, according to Ghaffari et al. (2005). The gas delivery device consisted of two sterile cylindrical Plexiglas exposure chambers (one for the NO₂ exposure, the second one for the control—exposure to synthetic air). The exposure chambers were deposited in a drying oven at 28°C (Figure 1). The NO₂, N₂, and O₂ obtained from Air Liquide GMP Europe (Mitry-Mory, France) were mixed together using digital mass flow regulators (Alicat Scientific, Inc., Tucson, USA) in order to get pre-calculated concentrations of NO₂ and maintain the O₂/N₂ ratio at 2/8 (v/v). The resulting gas mixture and the synthetic air were routed independently to each of the exposure chamber at a constant flow rate of 2 L/min, allowing parallel treatment of bacteria originating from the same bacterial culture. After passing through the exposure chamber, the NO₂ concentrations were monitored by AC32M nitrogen oxides analyzer (Environnement S.A, Poissy, France) and safely vented to a chemical hood. Temperature and relative humidity data were monitored to control reliable steady-state environmental conditions inside the exposure chambers. Three concentrations of NO₂ (0.1 ppm; 5 ppm and 45 ppm) were used in this study. After exposure, bacteria were diluted to A₅₈₀ = 2 in sterile saline solution and used for the subsequent experiments.

**Antibiotic Sensitivity Assays**

After NO₂ exposure, bacterial sensitivity to ciprofloxacin, chloramphenicol, tobramycin and kanamycin (Sigma-Aldrich, St. Quentin Fallavier, France) was assayed. The minimum inhibitory concentration (MIC) was determined by the broth microdilution method achieved in DMB. Briefly, NO₂-exposed bacteria were diluted to A₅₈₀ = 0.08 and added to a 96-well test plate (Nunc™, Roskilde, Denmark) containing different concentrations of antibiotics in triplicate. The test plates were incubated at 28°C for 24 h. Synthetic air-exposed bacteria were used as control. MIC was defined as the lowest antibiotic concentration that inhibited bacteria growth as determined by turbidimetry at A₅₈₀.

Growth inhibition assays were achieved as previously described (van Sorge et al., 2013). Exposed bacteria were diluted in DMB supplemented by the indicated antibiotics in subinhibitory concentrations (the last antibiotic concentrations allowing bacterial growth). Bacteria were added to Bioscreen Honeycomb plates (Oy Growth Curves Ab Ltd., Helsinki, Finland) in a total volume of 200 µL of DMB (A₅₈₀ = 0.08). Growth was measured every 15 min (A₅₈₀) for 24 h. The NO₂ effect on the bacterial antibiotic sensitivity was calculated as the percentage of bacterial growth with antibiotics after NO₂ exposure on the bacterial growth with antibiotics after exposure to synthetic air, using the following formula: 100×A₅₈₀ NO₂ exposed bacteria/A₅₈₀ synthetic air exposed bacteria (%).

**Motility Assays**

Swimming and swarming motility assays were performed on agar plates using DMB containing 0.2% (wt/vol) and 0.5% (wt/vol) agar, respectively, as previously described (Déziel et al., 2001), Briefly, 5 µL of NO₂ or synthetic air-exposed bacteria were spotted on the surface of agar plates. The resultant diameters of swim and swarm zones were measured after 24 h of incubation at 28°C. Motilities were assayed in three independent experiments with three replicates for each experimental condition.

**Biofilm Monitoring By Confocal Laser Scanning Microscopy**

NO₂ or synthetic air-exposed bacteria were diluted in sterile saline solution to A₅₈₀ = 1 to avoid bacterial multiplication, and added to glass-bottom dishes (SensoPlate™, VWR, Fontenay-sous-Bois, France). After 2 h of incubation at 28°C, planktonic bacteria were removed and bacterial adhesion on glass-bottom dishes was observed using a confocal laser scanning microscope (CLSM 710, ZEISS) with an immersion objective 63×. After addition of DMB, the samples were incubated at 28°C for 24 h. Biofilms were rinsed with saline solution and observed using CLSM. All biofilm assays were performed in three independent experiments with two replicates for each experimental condition. The biofilm thickness and related biomass (bacterial volume, µm³/µm²) were estimated from 6 fields on 3 independent experiments using COMSTAT software (Heydorn et al., 2000).

**Gene Sequences Identification**

The non-annotated genome drafts of MFN1032 and MFAF76a were used to identify the corresponding nucleotide sequences (data not shown). Homologous sequences search in P. fluorescens annotated genomes was performed using pseudomonas genome database (http://pseudomonas.com/). The conserved nucleotide sequences were identified in P. fluorescens MFN1032 and MFAF76a using Blast+ (Stand-alone) software (v. 2.2.30, NCBI) according to Altschul et al., 1997, and are listed in Table S2.

**Extraction and Quantification of Bis-(3′, 5′)-Cyclic Dimeric Guanosine Monophosphate (c-di-GMP)**

Extraction and quantification of intracellular c-di-GMP level were performed in NO₂ or synthetic air-exposed bacteria as previously described (Spangler et al., 2010; Strehmel et al., 2015). Identification and quantification of c-di-GMP was performed using three specific mass transitions from molecule ion m/z 691 to the product ions: m/z 152, m/z 135, and m/z 540. The external calibration was carried out at c-di-GMP concentrations ranging from 10 ng to 200 ng in 500 µL H₂O using the internal standard cXMP (50 ng). The resulting concentrations of c-di-GMP were normalized against total protein contents of respective cultures, which was determined by the bicinchoninic acid assay (Smith et al., 1985). All experiments were performed in three replicates for each experimental condition.
P. fluorescens Response to Gaseous NO₂

FIGURE 1 | Schematic representation of NO₂ gas delivery system. Bacterial NO₂ exposure was done in gas phase for 2 h. Two exposure chambers (one for the NO₂ exposure, the second one for the control—synthetic air exposure) were used. The gases, including NO₂, N₂, and O₂ were mixed together to obtain pre-calculated concentrations of NO₂ and maintain the O₂/N₂ ratio at 2/8 (v/v). NO₂ concentrations, temperature and relative humidity were controlled.

Quantitative RT-PCR
Total RNA was prepared by the hot acid-phenol method (Bouffartigues et al., 2012) from NO₂-exposed and not bacteria. Residual DNAs were eliminated by acid phenol treatment. The absence of DNA was confirmed by showing that PCR reactions failed without prior cDNA synthesis. RNAs were nonspecifically converted to single stranded cDNAs using the High Capacity cDNA Archive Kit (Applied Biosystems). Synthesis of cDNAs and real time PCR, allowing the quantification of mRNAs of interest were performed as previously described (Gicquel et al., 2013) using primers listed in Table S3.

Statistical Analysis
All experiments were carried out several times. To assess the significance of differences between the obtained data, Mann-Whitney test or pairwise strain comparisons (t-test) were applied and quantified the significance as (*) for \( p < 0.05 \), (**) for \( p < 0.01 \) and (***) for \( p < 0.001 \).

RESULTS AND DISCUSSION
NO₂ is one of the most common air pollutants, but its effects on the air microbiota is poorly studied. In order to assess the bacterial response to NO₂, airborne P. fluorescens MFAF76a and clinical control MFN1032 strains were exposed to gaseous NO₂ (as shown Figure 1) at 0.1, 5, or 45 ppm concentrations, and their effects on bacteria were evaluated in terms of motility, biofilm formation, antibiotic resistance, as well as expression of several chosen target genes. While 0.1 and 5 ppm of NO₂ did not lead to any significant modification of the studied parameters in both the bacteria (data not shown), several alterations were observed when the bacteria were exposed to 45 ppm of gaseous NO₂. We thus chose to focus on this concentration.

NO₂-Mediated Modifications of Bacterial Biofilm
In order to test the NO₂ effect on P. fluorescens biofilm, both airborne MFAF76a and clinical MFN1032 were exposed to gaseous NO₂ and synthetic air and grown for 4 h in static conditions. In the control condition, the airborne strain MFAF76a produced only a poorly structured biofilm with low biomass and thickness (Figure 2A). To the best of the authors' knowledge this is the first time that the biofilm of airborne P. fluorescens strain was investigated. On the opposite, the clinical strain MFN1032 was able to form a structured mushroom-like biofilm, with about 2 and 3 fold more biomass and thickness than the airborne strain MFAF76a, respectively (Figure 2B, control). These data are consistent with previous studies showing that clinical strains can strongly adhere and form structured biofilms (Rossignol et al., 2008; Ma et al., 2009). After NO₂ exposition, the airborne strain MFAF76a produced biofilms with about 3 fold increase of the maximal thickness, while the biomass was similar (Figure 2A), when compared to synthetic air treatment. These data suggest that NO₂ led to induce biofilm formation in this strain. Accordingly, similar NO concentrations were previously found to promote an increase of biofilm formation in P. aeruginosa (Barraud et al., 2006), suggesting a common effect between NO and NO₂ treatment. On the other hand, NO₂...
exposure of the clinical strain MFN1032 led to a 1.7 fold increase in biofilm production in terms of biomass, while the maximal thickness was unchanged (Figure 2B). Taken together these data suggest that the NO$_2$-mediated biofilm modifications are strain-dependent. Since we have shown previously that airborne MFAF76a expresses a virulence activity toward A549 epithelial pulmonary cells (Duclairoir Poc et al., 2014), these data suggest that elevated concentrations of NO$_2$ increases biofilm formation in potentially virulent airborne strain and may represent a sanitary risk.

Since biofilm formation is related to increased c-di-GMP production (Ha and O’Toole, 2015), we next quantified the c-di-GMP levels after NO$_2$ or synthetic air exposure. As shown in Figure 2C, both NO$_2$-exposed _P. fluorescens_ strains did not exhibit statistically significant variations of intracellular c-di-GMP concentrations. This was quite surprising since observed in our study NO$_2$ mediated biofilm induction. NO$_2$-mediated reduction of the intracellular c-di-GMP level leading to dispersion of _P. aeruginosa_ biofilms has been related to increase phosphodiesterases (PDEs) activity, and as a consequence to promote the switch between the biofilm and the planktonic ways of life (Petrova and Sauer, 2012; Roy et al., 2012; Li et al., 2013; Petrova et al., 2014). In this bacterium, the following PDEs, including DipA, MucR, NdbA and BdlA, are enzymes that are involved in c-di-GMP catabolism (Petrova and Sauer, 2012; Roy et al., 2012). The mRNA levels of _dipA, mucR, ndbA_ and _bdlA_ genes (KT186437, KT186445, KT186444 and KT186436 respectively, Table S2) were quantified by qRT-PCR experiments, in the two strains, that were both previously exposed to NO$_2$ or synthetic air. For the two strains, NO$_2$ exposure did not lead to any modification in gene expression (data not shown). Altogether, these data suggest that (i) NO$_2$ may have an effect on the structure or on the biomass of the biofilm, in case of the studied airborne or clinical strains, respectively, (ii) these phenotypes would not be related to variations of the intracellular c-di-GMP levels, and (iii) NO and gaseous NO$_2$ may have a common and concentration-dependent effect on biofilm formation.

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**FIGURE 2 | NO$_2$ effect on _P. fluorescens_ biofilm and intracellular c-di-GMP level.** (A) Airborne MFAF76a and (B) clinical MFN1032 _P. fluorescens_ strains were exposed in triplicate to 45 ppm of NO$_2$. Biofilm formation was analyzed in static conditions after 24 h development using confocal laser scanning microscope. The biofilm biomass and the maximum thickness were estimated from 6 fields on 3 independent experiments using COMSTAT software. Intracellular c-di-GMP concentrations (C) were measured in triplicate by LC-MS/MS for control and 45 ppm of NO$_2$ treated MFAF76a and MFN1032. Obtained results are presented as average values ± SEM. Statistical significance was calculated by the non-parametric Mann-Whitney-Test. n.s., non-significant.
NO₂ Reduced Bacterial Motility

Since biofilm structure and production were not implemented by c-di-GMP level in our conditions, we next assayed the effects of NO₂ on bacterial motility, since appendices like flagella and type IV pilus are also involved in the first step of biofilm formation, i.e., adhesion (Caiazza et al., 2007; Guttenplan and Kearns, 2013).

Swarming concerns motility in a liquid medium, mediated by production and activity of flagella. As shown on Figure 3A, gaseous NO₂ exposition significantly decreased the swimming motility of both strains, suggesting an impairment of the flagellum production and/or activity.

Swarming is a complex motility that has been related to functional flagella, type IV pilus and production of biosurfactants like cyclic lipopeptides (Ducloir-Poc et al., 2011) for some P. fluorescens strains, or rhamnolipids for P. aeruginosa strains (Caiazza et al., 2005). The airborne strain MFAF76a was unable to swarm in tested experimental conditions. On the opposite, MFN1032 is a swarmer clinical strain (Rossignol et al., 2008). As shown in Figure 3B, exposition to NO₂ but not to synthetic air led to fully inhibit the swarming motility of this strain.

Taken together, our data show that gaseous NO₂ treatment results in a decreased motility in both of the studied strains. This decrease in motility could be a consequence of a lower production of the required appendices. Alternatively it could also be due to lower appendices activity, suggesting that they could increase the attachment of the bacterium on the glass slide. This phenotype would then be consistent with the increase in biofilm maximal thickness in case of the airborne strain, and biomass in case of the clinical strain. However, to date, the switch between motility and biofilm had frequently been associated to variations in the c-di-GMP level (Ha and O’Toole, 2015), but, herein, the gaseous NO₂-mediated differences in terms of biofilm structure could not be related to any c-di-GMP level variations.

Effect of NO₂ on MexEF-OprN Efflux Pump Expression and Antibiotic Resistance

To further characterize the effects of gaseous NO₂ on bacterial physiology, we next assayed antibiotic resistance. Since NO₂, a member of RNS, was found to induce the expression of mexEF-oprN genes (Fetar et al., 2011) and modulate bacterial resistance to fluoroquinolones, chloramphenicol and aminoglycosides (Gusarov et al., 2009; McCollister et al., 2011; van Sorge et al., 2013), we investigated the effect of gaseous NO₂ on these phenotypes.

In order to study the effect of NO₂ on MexEF-OprN efflux pump, the transcription levels of mexE, mexF and oprN genes (KT070324, KT070321 and KT070325 for MFAF76a; KT070323, KT070322 and KT186432 for MFN1032, respectively) were compared using qRT-PCR in two P. fluorescens strains exposed or not to 45 ppm of NO₂. In airborne and clinical strains, the mexE mRNA level was increased by almost 14- and 100-fold respectively; that of mexF almost 3.5- and 47-fold respectively and that of oprN almost 4.6- and 73-fold respectively (Figure 4). These data show that NO₂ promoted mexEF-oprN expression, potentially causing modifications in P. fluorescens antibiotic resistance. We next tested the functionality of this pump. Since the MexEF-OprN RND efflux pump is involved in fluoroquinolone resistance, we next assayed bacterial sensitivity to ciprofloxacin by evaluating their MICs. As shown in Table 1, both the P. fluorescens strains were more resistant to this antibiotic after exposure to NO₂ than to synthetic air. Chloramphenicol is a nitroaromatic antimicrobial that is a substrate for MexEF-OprN (Köhler et al., 1997; Sobel et al., 2005). Accordingly, NO₂-exposed P. fluorescens strains MFAF76a and MFN1032 were about 2 fold more resistant to this antibiotic than synthetic air-treated bacteria (Figure 5). Taken together, these data suggest a possible higher activity of this efflux pump in response to NO₂ exposure. We next followed the growth of the NO₂-exposed P. fluorescens strains in DMB medium containing ciprofloxacin or chloramphenicol at the higher antibiotic concentration leading to bacterial growth (Figure 5). Data were standardized with the control, the synthetic air treated cells growth. While ciprofloxacin had no effect on NO₂-exposed bacteria, chloramphenicol at a concentration of 25 and 100 µg/mL for strain MFAF76a and MFN1032, respectively, led to an increase in growth for the two NO₂-exposed P. fluorescens strains (Figure 5). Remarkably, the statistically significant increase of bacterial growth was maintained from 2 to 10 h, suggesting a possible NO₂ protective effect that would be conserved for 8 h after exposure. Taken together, our data show that NO₂ induced mexEF-oprN gene expression, and consequently increased the resistance to ciprofloxacin and chloramphenicol.

MexEF-OprN-overproducing mutants with enhanced fluoroquinolone resistance often increase bacterial susceptibility to aminoglycosides apparently owing to impairment of the MexXY system (Sobel et al., 2005; Morita et al., 2015). The effect of NO₂ on tobramycin and kanamycin sensitivity was then assayed by performing MICs. As shown in Table 2, NO₂ treatment led to reduce the MICs of the two tested antibiotics, suggesting that NO₂ increases P. fluorescens sensitivity to aminoglycosides. Tobramycin and kanamycin, at subinhibitory concentration of 1.55 and 3.1 µg/mL, respectively, were found to decrease the growth of NO₂-exposed bacteria (Figure 6). This effect was observed only from 6 to 10 h of growth for MFN1032 and from 6 to 18 h of growth for MFAF76a, highlighting the time-limited NO₂ effect on bacterial antibiotic sensitivity. Altogether, our data show that NO₂ increases P. fluorescens sensitivity to tobramycin and kanamycin, accordingly its homolog NO is also found to increase P. aeruginosa sensitivity to tobramycin.

| Strain       | NO₂ concentration (ppm) | Ciprofloxacin MIC (µg/mL) | Chloramphenicol MIC (µg/mL) |
|--------------|-------------------------|---------------------------|-----------------------------|
| MFAF76a     | 0                       | 6.25                      | 50                          |
|             | 45                      | 12.5                      | > 100                       |
| MFN1032     | 0                       | 3.125                     | 150                         |
|             | 45                      | 6.25                      | 200                         |
FIGURE 3 | NO$_2$ decreases *P. fluorescens* motility. Airborne MFAF76a and clinical MFN1032 *P. fluorescens* strains were exposed in triplicate to 45 ppm of NO$_2$. Swimming (A) and swarming (B) motilities were assayed on DMB-swim/swarm plates after 24 h incubation. The motile bacterial movement was evaluated in three independent experiments with three replicates. The data were compared with control exposed to synthetic air. Obtained results are presented as average values ± SEM. Statistical significance was calculated by the non-parametric Mann-Whitney-Test $p < 0.05$ (*) and $< 0.001$ (**).

FIGURE 4 | NO$_2$ effect on MexEF-OprN and MexXY efflux pump gene transcription. The nucleotide sequences of the mexEF-, oprN- and mexXY-homolog genes were obtained using the non-annotated genome drafts of airborne MFAF76a and clinical MFN1032 *P. fluorescens*. The GenBank accession numbers of nucleotide sequences are listed in Table S2. Quantification of mRNA level was assayed using qRT-PCR on RNAs extracted from NO$_2$- and synthetic air-exposed *P. fluorescens*. The PCR reactions were performed in triplicate and the standard deviations were lower than 0.15 Ct. Statistical analysis used pairwise strain comparisons (t-test) $p < 0.01$ (**) and $< 0.001$ (**). Dotted line shows the gene expression in synthetic air-exposed control.

Remarkably, we have shown herein a link between gaseous NO$_2$ and soluble NO treatment. Indeed, NO is found to induce the expression of *mexEF-oprN* genes (Fetar et al., 2011) and modulates bacterial resistance to several antibiotics (Gusarov et al., 2009; McCollister et al., 2011; van Sorge et al., 2013). Since NO$_2$ and NO are related chemical toxic compounds, and since NO detoxification pathways have been deeply investigated, the NO$_2$ effects on several chosen target genes were tested.
FIGURE 5 | NO\textsubscript{2} protects \textit{Pseudomonas fluorescens} from chloramphenicol toxicity. After 2 h exposure to 45 ppm of NO\textsubscript{2}, growth of airborne MFAF76a (A) and clinical MFN1032 (B) \textit{P. fluorescens} with ciprofloxacin (\textbullet) and chloramphenicol (\square) was assayed. Growth curves were performed with ciprofloxacin (3.125 \textmu g/mL for MFAF76a and 1.156 \textmu g/mL for MFN1032) and chloramphenicol (25 and 100 \textmu g/mL respectively), and A\textsubscript{580} was recorded at the indicated time points. The control sample was bacteria exposed to synthetic air, and grown in presence of antibiotics in indicated concentrations. The data are shown as percentages of growth relative to synthetic air-exposed control. Pooled data from three independent experiments in duplicate ± SEM are reported. Statistical significance was calculated by the non-parametric Mann-Whitney-Test (*); n.s., non-significant. Dotted line shows the control (100%).

The most well-studied pathway for NO detoxification is based on flavohemoglobin (FlavoHb) (Hmp for \textit{E. coli} and Fhp for \textit{P. aeruginosa}), which acts as an NO dioxygenase to transform NO to NO\textsuperscript{−}\textsubscript{3} (Figure 7A) (Corker and Poole, 2003; Arai et al., 2005). After exposure to 45 ppm of NO\textsubscript{2}, the \textit{hmp} mRNA levels were increased almost 25- and 23-fold in MFAF76a and in MFN1032 (respectively KR818822 and KR818823 in Table S2 and Figure 8), indicating that NO\textsubscript{2} induces \textit{hmp} expression in both \textit{P. fluorescens} and suggesting a possible involvement of Hmp in NO\textsubscript{2} detoxification. The NO\textsubscript{2} effect on the \textit{hmp} synthesis was observed in other studies, where, to activate the Hmp-dependent detoxification pathway, NO\textsubscript{2} was proposed to be reduced to NO (Poole et al., 1996). In \textit{Pseudomonas} spp., NO\textsubscript{2} reduction can be performed by nitrite reductase (NIR) enzymes (Figures 7B,C), including the well-studied respiratory cytochrome cd\textit{l} nitrite reductase (Figure 7B) of the denitrification pathway (Arai et al., 2005; Shiro, 2012). According to the genome draft analysis (data not shown), both MFAF76a and MFN1032, like the majority of \textit{P. fluorescens} strains (Redondo-Nieto et al., 2013), do not possess denitrifying genes, but harbor the genes encoding for the assimilatory nitrite reductase NirBD (Figure 7C). The latter is part of the Nas assimilatory pathway (from nitrate assimilation), where nitrate is reduced to nitrite, which is then reduced to ammonia (Jeter et al., 1984; Moreno-Vivián et al., 1999). In order to test the NO\textsubscript{2} effect on the expression of \textit{nirBD} operon, the \textit{nirB} mRNA level (\textit{Pfl76a_nirB} - KT186428 - and \textit{Pfl1032_nirB} - KT070320 -, Table S2) was compared in the NO\textsubscript{2}-exposed or non-exposed \textit{P. fluorescens} strains. In both strains, the mRNA level of \textit{nirB} was not modified compared to the control condition (data not shown), indicating the absence of NO\textsubscript{2} effect on the expression of genes coding for assimilatory NIR. To the best of
our knowledge, the involvement of Nas pathway in NO/NO$_2$
detoxification was not demonstrated. Given the presence of
ammonium in DMB medium (Duclairoir-Poc et al., 2011), we
think that the production of supplementary ammonium through
the nitrite reduction is not appropriate. However, in order to
better understand the mechanism of the NO$_2$ detoxification,
the Hmp-, Nir- and Nas-mediated mechanisms should be
investigated in more details.

In this study, the response of airborne $P$. fluorescens MFA76a
to gaseous NO$_2$, as a marker of air pollution, was for the first
time investigated and compared to the response of the clinical
$P$. fluorescens MN1032 strain. We show that NO$_2$ leads to
increased biofilm formation through a c-di-GMP independent
mechanism, reduced motility, as well as increasing ciprofloxacin,
chloramphenicol resistance and aminoglycosides susceptibility.
The question is now to understand how the NO$_2$ leads to the
observed phenotypes. NO$_2$ has some similarities with its relative
NO. NO$_2$, like NO, induced the expression of mexEF-oprN genes,
coding the RND efflux pump MexEF-OprN. Its overexpression
could, among others, be involved in the observed increase of $P$. fluorescens resistance to ciprofloxacin and chloramphenicol. NO$_2$
duces also bacterial biofilm formation by strain-dependent
mode, without c-di-GMP production variation. Thus, the high $P$. fluorescens adaptability to many environments, and a
possible NO$_2$ propensity to increase some bacterial antibiotic resistance and biofilm formation may diminish the effectiveness
of antibiotic therapies in highly polluted area. In addition,
we show the NO$_2$-mediated upregulation of the hmp-homolog
gene in $P$. fluorescens, suggesting a possible common pathway
between NO and NO$_2$ detoxification. Taken together, our data
show that gaseous NO$_2$ can be perceived by airborne bacteria,
leading to physiological modifications that may be relevant
for human health (biofilm formation, antibiotic resistance).
In the context of the worrying increase of atmospheric NO$_2$
concentrations (Bernagaud et al., 2014), these findings are
of ecological relevance, especially because of the high NO$_2$
concentrations, found in the close vicinity of any vehicle.

**AUTHOR CONTRIBUTIONS**

TK contributed to the design of project, experiments, acquisition,
analysis, interpretation of data, and wrote the manuscript. CC
contributed in genes identification and qRT-PCR analysis. MB
contributed to the transformation of $P$. fluorescens strains. MN
and GB participated in c-di-GMP quantification. FD encouraged
the study on the airborne bacteria. SC and NO participated in
the design and drafted the manuscript. CDP led and coordinated
the global project by conceiving the study, and participated in
manuscript writing. All authors have read and approved the final
manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2016.00379
REFERENCES

Ahern, H. E., Walsh, K. A., Hill, T. C. J., and Moffett, B. F. (2007). Fluorescent pseudomonads isolated from Hebridean cloud and rain water produce biosurfactants but do not cause ice nucleation. Biogeosciences 4, 115–124. doi: 10.5194/bg-4-115-2007

Altschul, S. F., Madden, T. L., Schäffer, A. A., Zhang, J., Zhang, Z., Miller, W., et al. (1997). Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res. 25, 3389–3402. doi: 10.1093/nar/25.17.3389

Arai, H., Hayashi, M., Kuroi, A., Ishii, M., and Igarashi, Y. (2005). Transcriptional regulation of the flavohemoglobin gene for aerobic nitric oxide detoxification by the second nitric oxide–responsive regulator of Pseudomonas aeruginosa. J. Bacteriol. 187, 3960–3968. doi: 10.1128/JB.187.12.3960-3968.2005

Auger, C., and Appanna, V. D. (2015). A novel ATP-generating machinery to counter nitrosative stress is mediated by substrate-level phosphorylation. Biochim. Biophys. Acta 1850, 43–50. doi: 10.1016/j.bbagen.2014.09.028

Auger, C., Lemire, J., Cecchini, D., Bignucolo, A., and Appanna, V. D. (2011). The metabolic reprogramming evoked by nitrosative stress triggers the anaerobic utilization of citrate in Pseudomonas fluorescens. PLoS ONE 6:e28469. doi: 10.1371/journal.pone.0028469

Augusto, O., Bonini, M. G., Amano, A. M., Linares, E., Santos, C. C. X., and De Menezes, S. L. (2002). Nitrogen dioxide and carbonate radical anion: two emerging radicals in biology. Free Radic. Biol. Med. 32, 841–859. doi: 10.1016/S0891-5849(02)00786-4

Barraud, N., Haslett, D. J., Hwang, S.-H., Rice, S. A., Kjelleberg, S., and Webb, J. S. (2006). Involvement of nitrite in biofilm dispersal of Pseudomonas aeruginosa. J. Bacteriol. 188, 7344–7353. doi: 10.1128/JB.00779-06

Barraud, N., Storey, M. V., Moore, Z. P., Webb, J. S., Rice, S. A., and Kjelleberg, S. (2009). Nitric oxide–mediated dispersal in single—and multi—species biofilms of clinically and industrially relevant microorganisms. Microbiot. Biotechnol. 2, 370–378. doi: 10.1111/j.1751-7915.2009.00098.x

Bauer, H., Kasper-Giebl, A., Löflund, M., Giebl, H., Hitzenberger, R., Zibuschka, N., Barraud, N., Hassett, D. J., Hwang, S.-H., Rice, S. A., Kjelleberg, S., and Webb, J. S. (2008). Comparative study of 7 fluorescent pseudomonad clinical isolates. Can. J. Microbiol. 54, 19–27. doi: 10.1139/W07-110

Chaplain, A., Rossignol, G., Lesouhaitier, O., Merieu, A., Gruffat, C., Guerrillon, J., et al. (2008). Exposure to nitrogen dioxide (NO2) and respiratory disease risk. Rev. Environ. Health 13, 73–90.

Coker, H., and Poole, R. K. (2003). Nitric oxide formation by Escherichia coli deendence on nitrite reductase, the NO-sensing regulator Fnr, and flavohemoglobin Hmp. J. Biol. Chem. 278, 31584–31592. doi: 10.1074/jbc.M303282200

Cutruzzola, F., and Frankenberg-Dinkel, N. (2015). Origin and impact of nitric oxide in Pseudomonas aeruginosa biofilms. J. Bacteriol. 198, 55–65. doi: 10.1128/JB.00371-15

De Menezes, S. L. (2002). Nitrogen dioxide and carbonate radical anion: two emerging radicals in biology. Free Radic. Biol. Med. 32, 841–859. doi: 10.1016/S0891-5849(02)00786-4

Després, V. R., Alex Huffman, J., Burrows, S. M., Hoce, C., Salatoff, A. S., Buryak, G., et al. (2012). Primary biological aerosol particles in the atmosphere: a review. Tellus B 64, 262010 (10.3402/tellusb.v64i6.15398)

Déziel, E., Comeau, Y., and Villemure, R. (2001). Initiation of biofilm formation by Pseudomonas aeruginosa 57RP correlates with emergence of hyperpiliated and highly adherent phenotypic variants deficient in swimming, swarming, and twitching motilities. J. Bacteriol. 183, 1195–1204. doi: 10.1128/JB.183.4.1195-1204.2001

Duclairoir-Poc, C., Meylheuc, T., Ngoya, S., Groboillot, A., Bodulis, J., Taupin, L., et al. (2011). Influence of growth temperature on cyclolipopeptides production and on adhesion behaviour in environmental strains of Pseudomonas fluorescens. J. Bacteriol. Parasitol. S1-002. doi: 10.4172/2155-9597.S1-002

Duclairoir Poc, C., Verdon, J., Groboillot, A., Barreau, M., Toucourou, H., Mijouin, L., et al. (2014). Airborne fluorescent pseudomonads: what potential for virulence? Int. J. Curr. Microbiol. Appl. Sci. 3, 708–722.

Dybwad, M., Granum, P. E., Bruheim, P., and Blatny, J. M. (2012). Characterization of airborne bacteria at an underground subway station. Appl. Environ. Microbiol. 78, 1917–1929. doi: 10.1128/AEM.07212-11

Fang, Z., Ouyang, Z., Zheng, H., Wang, X., and Hu, L. (2007). Culturable airborne bacteria in outdoor environments in Beijing, China. Microb. Ecol. 54, 487–496. doi: 10.1007/s00248-006-0526-4

Flatley, J., Barrett, J., Pullan, S. T., Hughes, M. N., Green, J., and P oole, R. K. (2007). The chemistry of DNA damage from nitric oxide and peroxynitrite. Mutat. Res. Mol. Mech. Mutagen. 424, 37–49. doi: 10.1016/S0168-2723(07)00253-4

Glafarri, A., Neil, D. H., Ardakani, A., Road, J., Ghahary, A., and Miller, C. C. (2005). A direct nitric oxide gas delivery system for bacterial and mammalian cell cultures. Nitric Oxide 12, 129–140. doi: 10.1016/j.niox.2005.01.006

Gicquel, G., Bouffartigues, E., Bains, M., Oxaran, V., Rosay, T., Lesouhaitier, O., et al. (2013). The extra-ctytoplasmic function sigma factor SigX modulates biofilm and virulence-related properties in Pseudomonas aeruginosa. PLoS ONE 8e80407. doi: 10.1371/journal.pone.0080407

Guttenplan, S. B., and Kearns, D. B. (2013). Regulation of flagellar motility during biofilm formation. FEMS Microbiol. Rev. 37, 849–871. doi: 10.1111/1574-6976.12018

Kondakova et al. P. fluorescens Response to Gaseous NO2
van Sorge, N. M., Beasley, F. C., Gusarov, I., Gonzalez, D. J., Köckritz-Blickwede, M., von, Anik, S., et al. (2013). Methicillin-resistant *Staphylococcus aureus* bacterial nitric oxide synthase affects antibiotic sensitivity and skin abscess development. *J. Biol. Chem.* 288, 6417–6426. doi: 10.1074/jbc.M112.448738

Varivarn, K., Champa, L. A., Silby, M. W., and Robleto, E. A. (2013). Colonization strategies of *Pseudomonas fluorescens* Pf0-1: activation of soil-specific genes important for diverse and specific environments. *BMC Microbiol.* 13:92. doi: 10.1186/1471-2180-13-92

WHO |Air quality guidelines - global update (2005). WHO. Available online at: http://www.who.int/phe/health_topics/outdoorair/outdoorair_aqg/en/ (Accessed September 17, 2015).

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