# This is the preprint version of the contribution published as:

Chautrand, T., Souak, D., Kondakova, T., Depayras, S., Merlet-Machour, N., **Heipieper, H.J.**, Feuilloley, M., Orange, N., Duclairoir-Poc, C. (2020): Air pollution and other environmental stresses: gaseous NO<sub>2</sub> exposure leads to specific alterations of *Pseudomonas fluorescens* In: Casares, J., Longhurst, J., Barnes, J. (eds.) *Air pollution XXVIII* WIT Transactions on Ecology and the Environment 244 WIT Press, Southampton, p. 53 – 63

# The publisher's version is available at:

http://dx.doi.org/10.2495/AIR200051

## AIR POLLUTION AND OTHER ENVIRONMENTAL STRESSES: GASEOUS NO<sub>2</sub> EXPOSURE LEADS TO SPECIFIC ALTERATIONS OF PSEUDOMONAS FLUORESCENS.

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#### ABSTRACT

In the environment, microorganisms are subjected to a wide range of stresses. These stresses can be of natural origin, like temperature variations and ultraviolet exposure, but can also originate from humans like air pollution. The effects of air pollution on humans are more and more studied and reveal increasing concerns for human health, including augmentations of respiratory infections. However, the microbial responses to atmospheric pollution are still largely unknown. In a similar fashion, few studies investigate the effects of UV radiation on microorganisms. As NO<sub>2</sub> is an air pollutant causing nitrosative stress in biological organisms by reacting with various biological molecules, solar UV radiations are also an important environmental source of cell damage. UVB can directly damage DNA and cause erythema, but only represent 6% of the total UV reaching the earth surface. The 94% others are UVA, that cause oxidative stress in the cells. Since oxidative and nitrosative stresses are interlinked, the exposition of airborne bacteria to these two stresses could have synergistic consequences. In this study, the airborne Pseudomonas fluorescens strain MFAF76a was exposed successively to gaseous NO2 and UV light to assess whether these two environmental stresses have synergistic effects on bacterial physiology. Bacterial growth was assessed by optical density and membrane permeability by flow cytometry. Exposures to successively gaseous NO2 and UVB light lead to a non-synergistic decrease of bacterial viability. Furthermore, only NO<sub>2</sub> seems to damage the membrane and induces membrane permeabilization. Lipidomic analysis reveals similarities between the lipidic profile of bacteria in their exponential growth phase or for the exposed ones to NO<sub>2</sub> during their stationary growth phase. Furthermore, lipidic alterations show that mechanisms induced by NO<sub>2</sub> differ from those implemented by temperature. In conclusion, this study reveals that bacterial alterations caused by  $NO_2$  are specific, with a strong emphasis on membrane damage.

Keywords: NO2, Ultraviolet, Pseudomonas fluorescens, air pollution, health, bacterial membranes

## **1 INTRODUCTION**

Air pollution is a world-wide problem for both human health and environment. A major part of current air pollution is the result of human activities such as transport, industry and power plants [1]. Nitrogen dioxide  $(NO_2)$  is an air pollutant released by fuel combustion processes. High concentrations of  $NO_2$  are associated with both environmental problems such as photochemical smog and acid rains, as well as health issues such as asthma and bronchitis [1], [2]. NO<sub>2</sub> is a reactive nitrogen species (RNS), along other related molecules such as nitric oxide (NO) and peroxynitrite (ONOO<sup>-</sup>). RNS can damage various biomolecules such as DNA [3], lipids [4] and proteins [5] by reacting with them and are toxic for biological organisms at high concentrations.

In nature, microorganisms are regularly exposed to a wide range of stresses, such as temperature changes, ultraviolet (UV) exposure, and low water availability. Solar UV radiations are an important cause of cell damage for biological organisms [6]. Ultraviolet B (UVB) have a wavelength comprised between 280 and 315nm. Exposure to UVB can directly damage DNA, and cause various skin pathologies in human, such as erythema, dermatitis and skin cancer [6]. However, UVB only represent 6% of the total UV reaching the surface, the other 94% being mostly ultraviolet A (UVA), which possess a wavelength comprised between 315 and 400nm. UVA are less energetic than UVB and do not directly affect DNA, but they can cause more oxidative stress in cells. Atmospheric pollution can add to this stress, notably by the nitrosative stress induced by anthropogenic RNS. The effects of air pollution on bacteria is not well studied, and its combination to other environmental stresses even less. Indeed, the addition of multiple environmental stresses could have synergistic effects on bacterial physiology. Since oxidative stress and nitrosative stress are strongly interlinked, the combination of these two stresses could lead to synergistic effects not suspected when they are studied separately. Comprehension of the interactions between air pollution and other environmental stresses on bacteria could help to understand the anthropogenic alterations on environmental microbial communities and their effect on human health.

This study aims to elucidate the relationship between air pollution and other environmental stresses regarding their effect on the physiology of the airborne strain of Pseudomonas fluorescens MFAF76a. First, bacteria were successively exposed to gaseous NO<sub>2</sub> and UV light to assess their viability and membrane permeability. The effects of NO<sub>2</sub> on bacterial permeability prompted us to assess the potential impacts of NO<sub>2</sub> on the lipid composition of the strain MFAF76a. To appreciate the specificity of the cell response to NO<sub>2</sub> as an environmental stress, these results were compared to the effects of temperature and growth phase on bacterial lipidic composition. As MFAF76a presents several virulence factors [7] and could potentially adapt to a human host, the temperature studied were the optimal growth temperature of the strain of 28°C and the human body temperature of 37°C.

#### 2 MATERIALS AND METHODS

#### 2.1 Bacterial strain and growth conditions

The bacteria used in this study is the strain *Pseudomonas fluorescens* MFAF76a. This strain was isolated in the Rouen harbor (Normandy, France) from dust clouds generated during crop ship loading [7]. Strain MFAF76a was precultured 24h at 28°C or 37°C in Lysogeny Broth (LB) (yeast extract 5g/L, peptone 10g/L, NaCl 5g/L). The preculture was used to prepare the culture at a final optical density at 580nm (OD<sub>580</sub>) of 0.08 in Davis Minimal Broth (DMB) [8]. Cultures were incubated at 28°C or 37°C under agitation at 180rpm.

#### 2.2 Exposition to NO<sub>2</sub>

After 16h of incubation at 28°C, bacterial cultures were centrifuged at 13,000g for 10min at 4°C and were resuspended in DMB at a final optical density at an OD<sub>580</sub> of 4. 125 $\mu$ L of bacteria were deposited on cellulose acetate filters (pore size 0.2 $\mu$ m, Sartorius Stedim Biotech GmbH, Göttingen, Germany). The filters were then deposited on Petri dishes containing DMB with 15g/L agar. Bacteria were incubated 4h at 28°C after which the filters were transferred on 15g/L agar plates. The agar plates were placed into the gaseous exposition system developed in the laboratory [9]. Bacteria were exposed to 45ppm of

gaseous NO<sub>2</sub> with a N<sub>2</sub>/O<sub>2</sub> ratio of 8/2 or synthetic air for 2h with a constant gas stream of 2L/min.

## 2.3 UV exposure

After exposure to gaseous  $NO_2$ , all the agar plates were put to obscurity. One agar plate exposed to  $NO_2$  and one agar plate exposed to synthetic air were placed at the centre of the UV exposition chamber (Irradiation chamber BS-03, Opsytech Dr. Gröbel GmbH, Ettlingen, Germany). Bacteria were exposed to either 33mJ/cm<sup>2</sup> of UVA, 15mJ/cm<sup>2</sup> of UVB or both. Control bacteria were not exposed to any radiation. Bacteria were then resuspended in sterile physiological water (NaCl 9g/L).

## 2.4 Membrane permeability assays

The bacterial suspension after exposure was stained for 15 min with propidium iodine 30nM (PI) and SYTO9 5.01nM from the kit Live/Dead Backlight (L-7012, Thermo Fisher Scientific, Saint-Herblain, France). Bacteria were observed using flow cytometry (CytoFLEX S, Beckman Coulter, Villepinte, France) and the Cytexpert v1.2 software (Beckman Coulter, Villepinte, France). PI and SYTO9 were excited at a wavelength of 488nm and their fluorescence emission were detected at 690±50nm and 525±40nm respectively. 10,000 events were collected for each measurement at medium flow ( $30\mu$ I/min). Positive control for membrane permeabilization was performed on bacteria by treating them with 50% ethanol for 10 minutes before staining. The software was calibrated using the positive permeabilization control. Significativity of differences between mean values were assessed using t-test.

## 2.5 HPTLC-MALDI TOF MSI analysis of glycerophospholipids

The bacterial GP analysis was performed as previously described [10]. After exposure and resuspension at  $OD_{580}=2$  in sterile saline solution, bacteria were centrifuged at  $4^{\circ}C$ (13,000xg) and rinsed three times with sterile saline solution. Aliquots were resuspended in deionized water and lyophilized using a Freeze Dryer Heto PowerDry PL9000-50/ HSC500 (Thermo Fisher Scientific, Saint-Herblain, France). Lipid extraction was performed according to the method of Bligh and Dyer [11]. Bacterial lipid extracts (100  $\mu$ L) were separated in triplicates on High Performance Thin-Layer Chromatography (HPTLC) plates using CHCl<sub>3</sub>/CH<sub>3</sub>CH<sub>2</sub>OH/H<sub>2</sub>O/N(CH<sub>2</sub>CH<sub>3</sub>)<sub>3</sub> (35/35/7/35, v/v/v/v) as mobile phase. The GP spots were visualized by UV fluorescence (365nm) after labeling with 0.05% primulin dye in acetone/water, (8/2, v/v). Retention factors ( $R_f$ ) are calculated using the Sweday JustTLC software (v. 4.0.3, Lund, Sweden). Moreover, the following staining was performed in order to identify the lipid classes present in the HPTLC bands: (i) 2% azure A in 1 mM sulfuric acid for sulfatides and sulfoglycolipids, and (ii) 0.25% ninhydrin prepared in acetone-lutidine (9:1, v/v) for phosphatides and lipids having a free amino group. GPs were identified directly on HPTLC plates by Matrix-Assisted Laser Desorption Ionization mass spectrometry - Time of Flight (MALDI-TOF) using an Autoflex III mass spectrometer equipped with a laser Optibeam TM Nd/YAG (355 nm) with a 200-Hz tripled-frequency (Bruker Daltonics, Bremen, Germany). The DHB (200 g/L in acetonitrile/0.1% TFA, 90/10, v/v) was chosen as MALDI matrix. Autoflex III mass spectrometer was run in the reflector positive ion mode using the TLC-MALDI software (v. 1.1.7.0, Bruker Daltonics, Bremen, Germany). The Post Source Decay (PSD) spectra were acquired using FlexControl software (Bruker Daltonics).

The Mass Spectrometry Imaging (MSI) was performed using the FlexImaging software (v. 2.1., Bruker Daltonics, Bremen, Germany). The GP distribution was reconstructed according to the m/z of GPs, identified by MS and PSD spectra in combination with LIPID MAPS database and previously obtained data [10]. Individual lipid spots were labeled by a specific color code according to their m/z.

## 2.6 Fatty acids analysis

Fatty acid methyl esters were prepared by incubation for 15 min at 95°C in a boron trifluoride  $(BF_3)$ /methanol (140 g BF<sub>3</sub> per liter of methanol) and extracted with hexane as described by Morrison and Smith [12]. Fatty acid methyl esters were separated and analyzed by gas chromatography (GC) coupled to flame ionization detection using an Agilent Technology, 6890 Network GC System, 7683 Series Injector equipped with a split/splitless injector. The apparatus was equipped with a CP-Sil 88 capillary column (Chrompack, Middelburg, the Netherlands; length, 50m; inner diameter, 0.25 mm; 0.25 mm film). FAs were identified by coinjection as internal standards of reference compounds obtained from Supelco (Bellefonte, Pennsylvania, USA) and were quantified on the basis of their peak areas in total ion chromatograms. All experiments were performed in triplicate. The degree of FA saturation was determined as the ratio between the saturated FAs and the unsaturated FAs [13].

## **3 RESULTS AND DISCUSSION**

In the environment, bacteria are submitted to a wide array of stresses, such as variations of temperatures, UV radiation and nitrosative stress through atmospheric pollution. In particular, the effects of the atmospheric pollutant  $NO_2$  on bacteria is poorly understood, and its interaction with other types of environmental stresses even less. In order to study the effects of atmospheric  $NO_2$  on bacteria and its potential modulation by other environmental conditions, the airborne Pseudomonas fluorescens strain MFAF76a was exposed to  $NO_2$  or synthetic air for 2 hours, through an exposition system described previously [9], as well as different temperatures and different UV radiations.

3.1 NO<sub>2</sub> and UV exposure induce bacterial mortality through different processes

 $NO_2$  is a RNS, which can cause important damage to biological molecules in a similar fashion to ROS. UV radiation can harm biological organisms through different modes of actions depending mostly on the wavelength of the radiation. UVA-induced damages result mostly from their capacity to induce the formation of ROS in the cell through photochemical reactions with proteins and other cell components [14]. Meanwhile, UVB are more energetic and can mutate DNA by forming pyrimidine dimers at sequences containing 5methylcytosine, but they induce less oxidative stress in the cell. Since oxidative and nitrosative stresses are highly intertwined, we hypothesized that UV radiation could potentiate the nitrosative stress induced by NO2. To assess whether this hypothesis was true, P. fluorescens MFAF76a growth was followed after exposure to both NO<sub>2</sub> and different UV wavelengths (Fig. 1). NO<sub>2</sub> concentration of 45 ppm correspond to the concentration of NO<sub>2</sub> capable of causing irreversible consequences to human health [15]. Since UVB and UVA have drastically different ways to affect the cell, it is important to differentiate them to understand the bacterial response to this stress. However, most of the information available on the doses received at the surface does not specify the nature of the UV measured. The UVB dose used in this study was 15mJ/cm<sup>2</sup>, close to the ones used in phototherapy to treat patients with dermatitis [16]. To maintain an equivalent exposure time for both types of UV



Figure 1: Growth curve of P. fluorescens MFAF76a strain after exposure to UV radiations and gaseous NO<sub>2</sub>. Bacteria were exposed to either 45ppm of NO<sub>2</sub> (Exposed to NO<sub>2</sub>) or synthetic air (Control). After exposure to NO<sub>2</sub>, bacteria were exposed to either 33mJ/cm<sup>2</sup> of UVA (+UVA), 15mJ/cm<sup>2</sup> of UVB (+UVB), both 33mJ/cm<sup>2</sup> of UVA and 15mJ/cm<sup>2</sup> of UVB (+UVAB) or not exposed to light. Bacterial growth was then monitored through OD<sub>580</sub> for 24h. Data represents the means ±SD of three experiments.

taking in account the limits of the UV exposition chamber, the dose of 33mJ/cm<sup>2</sup> was used for UVA.

While none of the conditions seems to affect the growth rate of the bacteria on the long term, the shift of the growth curves in bacteria exposed to  $NO_2$  and UVB indicates an important mortality rates among bacteria exposed to these conditions. Intriguingly, despite their ability to induce oxidative stress, UVA does not increase mortality even when combined with  $NO_2$ . It could be argued that the doses used in this study were not important enough, but similar experiments were done with doses as high as  $4J/cm^2$  with no difference (data not shown). Furthermore, even though  $NO_2$  and UVB both induce mortality in MFAF76a strain, the combination of these two stresses does not further increase mortality.

To better understand the mechanisms underlying these independent mortality rates, membrane permeability of bacteria exposed to NO<sub>2</sub> and UV radiation was assessed (Fig. 2).



Figure 2: Membrane permeability of MFAF76a after exposure to UV radiations and gaseous NO<sub>2</sub>. Bacteria were exposed to either 45ppm of NO<sub>2</sub> (Exposed to NO<sub>2</sub>) or synthetic air (Control). After exposure to NO<sub>2</sub>, bacteria were exposed to either 33mJ/cm<sup>2</sup> of UVA (+UVA), 15mJ/cm<sup>2</sup> of UVB (+UVB), both 33mJ/cm<sup>2</sup> of UVA and 15mJ/cm<sup>2</sup> of UVB (+UVAB) or not exposed to light. Membrane permeability was assessed by cytometry using the Live/Dead Backlight kit (L-7012, Thermofisher) according to the manufacturer recommendations. Data represents the means ±SD of three experiments. Statistical significance of the results was assessed by t-test. \*\*\*\*: p-value <0.0001.</p>

Permeability assays were realized with the green-fluorescing membrane-permeant DNA probe Syto9 and the red-fluorescing membrane impermeable DNA probe propidium iodide (PI), and their fluorescence was measured by flow cytometry. This labelling reveals three populations, the cells labelled by Syto9 only, not permeabilized, the cells labelled by IP (Syto9- IP+), totally permeabilized, and the cells labelled by both Syto9 and IP (Syto9+ IP+), partially permeabilized. Membrane permeability assays show that only less than 30% of bacteria not exposed to NO<sub>2</sub> are labelled with IP or IP and Syto9, compared to around 70% of bacteria exposed to NO<sub>2</sub>, regardless of UV exposure. Therefore, NO<sub>2</sub> exposure causes an important increase in bacterial membrane permeability, which is usually associated with a reduction of cell viability. UV exposure however does not affect membrane permeability at the concentrations studied. Lethality induced by UVB is therefore probably linked to DNA alterations and not oxidative stress. This suggests that the mechanisms leading to cell mortality between NO<sub>2</sub> and UVB exposure are unrelated.

3.2 High NO<sub>2</sub> concentrations lead to a specific alteration of membrane composition.

To assess whether the membrane permeability alterations caused by  $NO_2$  are linked to a reorganization of the bacterial membrane, the lipid composition of MFAF76a was assessed after exposure to  $NO_2$  at different concentrations (Table 1). The impact of  $NO_2$  on bacterial lipidome was assessed at 0.1, 5 and 45ppm. 0.1ppm mimics the maximum average exposure to  $NO_2$  recommended by the World Health Organization. 5ppm mimics the threshold causing reversible effects on human health, while 45ppm mimics concentrations causing irreversible effects on human health [2], [15].

Table 1: Effect of NO<sub>2</sub> exposure on glycerophospholipid inventory in *P. fluorescens* MFAF76a strain. The lipid identification was performed on the basis of MS and PSD spectra using LIPID MAPS database and the analysis of lipid standard. The presence of assigned m/z in lipid extract in triplicate is shown by grey squares. The colourless squares show the absence of assigned m/z in lipid extract. The lipid extractions, MS and PSD analysis in all studied conditions were performed in three replicates. EGP: exponential growth phase; SGP: stationary growth phase; GP: glycerophospholipid; PC: phosphatidylcholine; UGP: unknown glycerophospholipid; PE: phosphatidylethanolamine; PG: phosphatidylglycerol

Glycerophospholipid identification						NO <sub>2</sub> level, ppm			
Retention factor	GP class	Species	Molar mass (M), g/mol	m/z	Assignement	0	0.1	5	45
0.24±0.01		PC [16:0 + 16:1]	731.5	732.6	$[PC_1\!\!+\!\!H]^{\scriptscriptstyle +}$				
	PC	PC [16:0 + 16:1]	731.5	754.5	[PC1+Na]+				
		PC [16:0 + 18:1]	759.6	782.6	[PC <sub>2</sub> +Na] <sup>+</sup>				
0.26±0.02	UGP	UGP [16:0 + 17:0]	904.5	927.5	[UGP <sub>1</sub> +Na] <sup>+</sup>				
		UGP [16:0 + 18:0]	918.5	941.5	[UGP <sub>2</sub> +Na] <sup>+</sup>				
		UGP [17:1 + 18:0]	930.5	953.5	[UGP <sub>3</sub> +Na] <sup>+</sup>				
	PE	PE [16:0 + 16:1]	689.5	712.5	[PE1+Na]+				
0.43±0.02				734.5	[PE1+2Na- H] <sup>+</sup>				
		PE [16:0 + 17:1]	703.5	726.5	[PE <sub>2</sub> +Na] <sup>+</sup>				
				748.5	$\begin{array}{l} [PE_2\!\!+\!\!2Na\!\!-\!\!H]^+ \end{array}$				
		PE [16:0 + 18:1]	715.5	738.5	[PE <sub>3</sub> +Na] <sup>+</sup>				
				760.5	[PE <sub>3</sub> +2Na- H] <sup>+</sup>				
0.62±0.03	PG	PG [16:0 + 16:1]	720.5	743.5	[PG1+Na]+				
				765.5	[PG1+2Na- H] <sup>+</sup>				
		PG [16:0 + 18:1]	746.5	769.5	[PG <sub>2</sub> +Na] <sup>+</sup>				
				791.6	[PG <sub>2</sub> +2Na- H] <sup>+</sup>				

The glycerophospholipid (GP) composition of MFAF76a, exposed to three concentrations of NO<sub>2</sub> was comparable to that of control sample, exposed to synthetic air (Table 1). PC, PE and PG were found in all experimental conditions. However, one noticeable difference was that an unknown glycerophospholipid (UGP) was not detected in bacteria exposed to 45 ppm of NO<sub>2</sub>. This disappearance indicates that this GP is affected by this free radical, as was previously described [17]. An important concentration of NO<sub>2</sub> is necessary to notice the complete disappearance of UGP however, as UGP was still detected after exposure to 0.1ppm and 5ppm of NO<sub>2</sub>.

The literature reports, that the nitrogen reactive species could mediate the oxidation and/or nitration of membrane GPs, leading to the formation of lipid hydroperoxides, alcohols, and aldehydes [18]. However, no modification of GP m/z was found, indicating that NO<sub>2</sub> does not react with the GPs of MFAF76a in these conditions. However, RNS are described to react with polyunsaturated fatty acids, which are absent from the lipidome of MFAF76a strain. In consequence, the absence of GP reaction with NO<sub>2</sub> appears to be logical. Therefore, the membrane permeabilization caused by NO<sub>2</sub> (Fig. 2) does not come from alterations of their lipid composition, with maybe the exception of UGP.

To better understand the significance of this lipidic composition shift after bacterial exposure to NO<sub>2</sub>, it is necessary to determine the adaptation capabilities of this strain. Therefore, the lipidomic profile of the MFAF76a strain was also followed in two conditions (Table 2) that are known to affect membrane structuration, temperature and growth phase [19], [20]. Bacteria were cultivated at 28°C, the optimal growth temperature of the strain, and 37°C, the temperature of the human body, to observe the modifications that would be necessary for this potential pathogen to infect humans. Lipidomic analysis of the bacteria was then realized either during their exponential growth phase (EGP) or their stationary growth phase (SGP).

Interestingly, the augmentation of temperature did not lead to major differences in GP composition inside the bacterial membrane. However, results show that UGP is not detected in bacteria during the EGP. The lipidic profile of bacteria exposed to  $NO_2$  during the stationary growth phase (Table 1) is therefore very similar to the lipidic profile of bacteria in exponential growth phase (Table 2). During the EGP, bacterial membranes are usually more fluid and disorganized than during the SGP. The shift caused by  $NO_2$  toward an EGP-like lipidic profile could therefore be partially responsible for the constated membrane permeabilization after exposure. This transition could improve bacterial fitness in an environment causing important nitrosative stress.

Table 2:Effect of temperature and growth phase on glycerophospholipid inventory *in P. fluorescens* MFAF76a strain. The lipid identification was performed on the basis of MS and PSD spectra using LIPID MAPS database and the analysis of lipid standard. The presence of assigned m/z in lipid extract in triplicate is shown by grey squares. The colourless squares show the absence of assigned m/z in lipid extract. The lipid extractions, MS and PSD analysis in all studied conditions were performed in three replicates. EGP: exponential growth phase; SGP: stationary growth phase; GP: glycerophospholipid; PC: phosphatidylcholine; UGP: unknown glycerophospholipid; PE: phosphatidylethanolamine; PG: phosphatidylglycerol.

Glycerophospholipid identification						Temperature, °C / growth phase				
								37°C		
Retention factor	GP class	Species	Molar mass (M), g/mol	m/z	Assignement	EGP	SGP	EGP	SGP	
		PC [16:0 + 16:1]	731.5	732.6	$[PC_1+H]^+$					
0.24±0.01	PC	PC [16:0 + 16:1]	731.5	754.5	[PC <sub>1</sub> +Na] <sup>+</sup>					
		PC [16:0 + 18:1]	759.6	782.6	[PC₂+Na]⁺					
0.26±0.02	UGP	UGP [16:0 + 17:0]	904.5	927.5	[UGP <sub>1</sub> +Na] <sup>+</sup>					
		UGP [16:0 + 18:0]	918.5	941.5	[UGP₂+Na]⁺					
		UGP [17:1 + 18:0]	930.5	953.5	[UGP₃+Na]⁺					
0.43±0.02	PE	PE [16:0 + 16:1]	689.5	712.5	[PE1+Na]+					
				734.5	[PE₁+2Na-H] <sup>+</sup>					
		PE [16:0 + 17:1]	703.5	726.5	$[PE_2+Na]^+$					
				748.5	$[PE_2+2Na-H]^+$					
		PE [16:0 + 18:1]	715.5	738.5	[PE₃+Na]⁺					
				760.5	[PE₃+2Na-H]⁺					
0.62±0.03	PG	PG [16:0 + 16:1]	720.5	743.5	[PG₁+Na]⁺					
				765.5	[PG₁+2Na-H]⁺					
		PG [16:0 + 18:1]	746.5	769.5	[PG <sub>2</sub> +Na] <sup>+</sup>					
				791.6	[PG₂+2Na-H]⁺					

To respond to environmental stresses, bacteria are not limited to changing the GP makeup of their membranes, but can also modify the degree of saturation of these GP's fatty acids [19], [20]. The global degree of saturation of MFAF76a's fatty acids was determined after exposure to  $NO_2$  or temperature variations at different growth phases (Fig. 3).



Figure 3: Effects of growth phase, temperature and exposition to NO<sub>2</sub> on fatty acid saturation in MFAF76a strain. Bacterial fatty acids were analysed and quantified using the GC from at least three independent lipid extracts. NO<sub>2</sub> exposure was done at 28°C. SGP: stationary growth phase, EGP: exponential growth phase. The degree of fatty acid saturation was calculated as the ratio between saturated fatty acids and unsaturated fatty acids, and presented as average value  $\pm$  SD. Statistical significance was calculated by one way ANOVA. \*P < 0.05; \*\*\*P < 0.001; n.s. non-significant

Interestingly, the augmentation of temperature did not lead to major differences in GP composition inside the bacterial membrane. However, results show that UGP is not detected in bacteria during the EGP. The lipidic profile of bacteria exposed to NO<sub>2</sub> during the stationary growth phase (Table 1) is therefore very similar to the lipidic profile of bacteria in exponential growth phase (Table 2). During the EGP, bacterial membranes are usually more fluid and disorganized than during the SGP. The shift caused by NO<sub>2</sub> toward an EGP-like lipidic profile could therefore be partially responsible for the constated membrane permeabilization after exposure. This transition could improve bacterial fitness in an environment causing important nitrosative stress. To respond to environmental stresses, bacteria are not limited to changing the GP makeup of their membranes, but can also modify the degree of saturation of these GP's fatty acids [19], [20]. The global degree of saturations at different growth phases (Fig. 3).

## **4** CONCLUSION

Here, for the first time, the use of two original exposition systems allowed us to highlight the combined impacts of gaseous NO<sub>2</sub> and UV on bacteria. Firstly, atmospheric NO<sub>2</sub> pollution induces specific effects on bacterial physiology. Exposure to NO<sub>2</sub> or UVB affects bacterial viability. But these two combined stresses do not seem to synergize. This suggests that UV-induced oxidative stress is not important enough to affect bacterial viability or synergize with NO<sub>2</sub> at doses bacteria are susceptible to encounter in the environment. NO<sub>2</sub> alteration of biological membranes seems to be an important component of its effects on bacterial viability, which is not the case for UVB. UVB radiations cause DNA damages, which could lead to bacterial mortality without affecting their membrane permeability. Furthermore, at the membrane level, NO<sub>2</sub> seems to induce responses different from other factors known to alter membrane stability such as temperature and growth phases. Overall, these results demonstrate that air pollution constitutes a unique challenge for bacteria to overcome. Indeed, NO<sub>2</sub> damage biological membranes notably, in ways that other environmental stresses do not.

## ACKNOWLEDGEMENTS

This work was conceived by T.C., D.S., S.D., T.K. and C.D.P. We would like to thank N.M.M and H.H. for their help with lipidic studies. We would also like to thank H.H., M.F. and N.O. for critical reading of the manuscript. This study was supported by PhD grants from the GRR SESA (Region Normandie) and was financially supported by the European Union (FEDER), "Région Normandie" and "Evreux Porte de Normandie" (France).

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