

Skin biological responses to urban pollution in an ex vivo model

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1	Skin biological responses to urban pollution in an ex vivo model
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ABSTRACT

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2 The skin epidermis is continuously exposed to external aggressions, including environmental 3 pollution. The cosmetic industry must be able to offer dedicated products to fight the effects 4 of pollutants on the skin. We set up an experimental model that exposed skin explants maintained in culture to a pollutant mixture. This mixture representing urban pollution was 5 6 designed on the basis of the French organization 'Air Parif' database. A chamber, called Pollubox®, was built to allow a controlled nebulization of P on the cultured human skin 7 explants. We investigated ultrastructural morphology by transmission electron microscopy of 8 high pressure frozen skin explants. At first, we detected by transmission electron microscopy 9 some matters smaller than 300 nm similar to diesel particles in the granular layer of the 10 epidermis. A global transcriptomic analysis indicated that the pollutant mixture was able to 11 12 induce relevant xenobiotic and antioxidant responses. Modulated detoxifying genes were further investigated by laser micro-dissection coupled to qPCR, and immunochemistry. Both 13 approaches showed that P exposure correlated with overexpression of detoxifying genes and 14 15 provoked skin physiological alterations down to the *stratum basale*. The model developed herein might be an efficient tool to study the effects of pollutants on skin as well as a 16 powerful testing method to evaluate the efficacy of cosmetic products against pollution. 17 **Keywords:** environmental pollution, ex vivo skin, transcriptomic, xenobiotic response, laser 18 19 capture micro-dissection, transmission electron microscopy

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1. INTRODUCTION

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2 Nowadays, urban air pollution is known to be a major threat to health and environment. The World Health Organization (WHO) reported that in 2016, 91% of the world population was 3 living in places where the WHO air quality guidelines were not met. Moreover, outdoor air 4 pollution in both cities and rural areas was estimated to cause 4.2 million premature deaths 5 6 worldwide in 2016. This mortality is due to exposure to small particulate matter of 2.5 µm or 7 less in diameter which causes cardiovascular and respiratory diseases and cancer. Increasing urbanization enhances particulate matter content in the atmosphere as wells as the level of 8 several other pollutants such as polycyclic aromatic hydrocarbons and heavy metals. Effect of 9 10 urban air pollutant exposure on the lung physiology has been largely described but little is known about those impacts on the skin (Estrella et al. 2019). 11 Epidermis, as the outermost line of defense of our organism is continuously exposed to a vast 12 13 range of stressors, including ultraviolet radiations and atmospheric pollutants. Skin ageing related to pollutant or UV radiation exposures has been clearly established (Vierkötter et al. 14 15 2010). Repeated stimulations of protective processes in skin seem to induce skin ageing (Estrella et al. 2019). The negative impact of environmental pollution on skin microbiota 16 composition has been also described (Jo et al. 2017). Moreover, pore obstruction by 17 18 particulate matter is able to enhance the formation of an anaerobic environment, which leads to the proliferation of *Propionibacterium acnes*, contributing to acne development. At a 19 molecular level, pollutants trigger the activation of Aryl hydrocarbon Receptor (AhR) that is 20 21 involved in the detoxification process (Denison and Nagy 2003; Iyanagi 2007). This chemosensor molecule is found in various types of tissues including the skin where it 22 mediates the xenobiotic response and modulates cell proliferation, inflammation and 23 melanogenesis (Abel and Haarmann-Stemmann 2010). Following stimulation, AhR 24 translocates to the nucleus of the cell where it interacts with AhR nuclear translocator 25

(ARNT) and binds the Xenobiotic Response Element (XRE) DNA domain to initiate the 1 2 transcription of genes involved in cell detoxification (Sonoda et al. 2003). Although this process is considered as a protective response against toxic compounds, its sustained 3 activation upon repeated exposures can be deleterious and might be responsible for skin 4 alterations (Mancebo and Wang 2015). For instance, among AhR target genes, cytochrome 5 P450 is responsible for the production of reactive oxygen species (ROS) (Denison and Nagy 6 2003; Hirabayashi 2005). The control of ROS is crucial for cellular homeostasis. An 7 excessive ROS production can cause DNA and protein damages as well as lipid peroxidation 8 that alters the barrier function and consequently skin hydration (Rhee 2006; Addor 2017). The 9 10 resulting redox imbalance leads to an antioxidant response involving the NRF2 pathway. As a transcription factor, NRF2 binds the Antioxidant Response Element DNA domain (ARE) to 11 promote the transcription of genes involved in antioxidant processes (Jackson et al. 2015). 12 13 NRF2 activation may be one way to avoid chronic inflammation through the decrease of oxidative stress (Kurutas 2016). 14 15 Most of the investigations launched to study the impact of pollutants on skin physiology were based mainly on in vitro studies using primary cells such as keratinocytes, fibroblasts or 16 immortalized human cell lines (HaCaT) (Min-Duk Seo et al. 2012; Binelli et al. 2018; Zhang 17 18 et al. 2017). These cellular models lack the high biological complexity that characterizes the human's biggest organ which is the skin. Cell culture cannot take into account the cross-talk 19 between distinct cell types and skin appendages such as hair follicles, apocrine and eccrine 20 21 sweat glands required to reflect the activities of the cutaneous tissue. Most previous studies in this field were conducted using a single pollutant rather than a mix of different pollutants like 22 it is the case under real conditions (Kazi et al. 2008; Philips et al. 2010). These limitations can 23 be considered as a brake to study the impact of pollutants on skin, and for the development by 24 the cosmetic industries of protective or regenerative products. Herein we present an 25

experimental model that relies on the use of human skin explants. In contrast to cultured cells and reconstructed epidermis, explants contain all resident cell types of the epidermis and dermis as well as skin appendages and can be cultured under air-liquid interface conditions for up to 10-12 days (Gasser et al. 2008). Thus, this ex vivo model constitutes an interesting alternative although skin explants stem from surgical tissues of donors and might differ in some biological processes or responses due to their genetic variability. Therefore, explants were used to see whether or not they could respond to pollutant exposure and serve as the basis of a test to evaluate anti-pollutant activities of cosmetic ingredients or products. To achieve experimental conditions as close as possible to reality, a pollutant mixture was designed according to the French non-profit organization 'Air Parif' and nebulized on the cultured human skin explants (HSE) within a dedicated chamber, the Pollubox®. Using this tool, only the *stratum corneum* (SC) was directly exposed to pollutants. Transmission electron microscopy (TEM) observations of high-pressure frozen HSE after pollutant exposure allowed the visualization of some matter similar to diesel particles in the epidermis and of an increase in the number of extracellular vesicles (EV) like structures, which are known to be involved in cell communication (Carrasco et al., 2019). Furthermore, gene expression analyses using whole genome microarrays showed that our experimental conditions were able to elicit a xenobiotic and antioxidant responses as well as to modulate genes involved in skin barrier homeostasis. Although the extent of such a modulation varied according to the donors, the expression of skin barrier-related genes was consistently found most affected. RT-qPCR performed on RNA extracted after laser capture micro dissection demonstrated that the level of induction was higher in the basal layers than in the granular layers. Altogether, these results supported that our experimental device can be efficient to test cosmetic ingredients or end products for their ability to counteract the effects of pollutants.

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MATERIALS & METHODS

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2 2.1 Preparation of human skin explants

- 3 BIO-EC Laboratory possesses an authorization from the Bioethics group of the general
- 4 director services of the French research and innovation ministry (registered under n°DC-
- 5 2008-542) to use human skin from surgical waste since 5th May 2010.
- 6 The study was performed in accordance with the Declaration of Helsinki after the patients had
- 7 given informed consent to use their skin samples by BIO-EC Laboratory.
- 8 Full-thickness human skin biopsies were obtained from abdomen of healthy female donors
- 9 who had undergone plastic surgery. The hypodermis was removed from the skin and circular
- explants (~1 cm diameter, 0.2 cm thickness and ~200 mg weight) were excised using a
- sample punch. Samples were placed immediately in BIO-EC's Explant Medium (BEM). From
- day 1 they were cultured under classical cell culture conditions (37°C in 5% CO₂). Table S1
- summarizes the different experiments completed during our study on the HSE from various
- volunteers in the context of the URBASKIN project supported by the French FUI (Fonds
- 15 Unique Interministériel).

16 2.2 Air pollutant exposure

- In order to develop the experimental conditions, several concentrations of pollutant mix (P)
- were evaluated. The pollutants were dissolved in nitric acid (Merck, 1.00441.0250), ethanol
- 19 (VWR, 20281.467), and DMSO (Sigma-Aldrich, 47231), as indicated in Table S2.
- 20 The dose of P, which can induce physiological modifications at the molecular level after
- 21 nebulization at 5 different doses (expressed in mg/m³ or in µg/ml) on HSE, was evaluated by
- 22 RT-qPCR quantification of stress marker gene expression (CYP1A1, GPX2, HMOX1, and
- 23 SQSTM1) comparing P versus organic solvent (OS) (Figure S1B). Induction of all evaluated
- 24 genes was observed after 24h of exposure to the highest dose (P1) among those tested,
- corresponding to 100 000 mg/m³ (Figure S1A). Therefore this dose was chosen for all further

- 1 investigations. The explants from the same donor were placed in BEM for 5 days before being
- 2 transferred to the Pollubox® and nebulized with 3 ml of P for 1h30 at the different selected
- 3 doses.
- 4 The Pollubox® device (Figure S2) is an exposure chamber designed by BIO-EC laboratory.
- 5 The system is composed of a chamber and a basis both made of poly (methyl
- 6 methacrylate) resin.
- 7 The basis contains 12 holes with a diameter of 8 mm restricting the exposure to the skin
- 8 explant surface alone. For the exposure to pollutants, skin explants are placed in a classical
- 9 12-well cell culture plate with 1 ml of BEM (BIO-EC culture medium) per well. The culture
- plate is then positioned under the basis of the Pollubox® in order to align skin explants at the
- levels of the holes of the basis. A nebulizer (Aerogen Pro®), placed on the top of the
- 12 chamber, allows to nebulize the liquid solution containing the pollutants. The generated
- aerosol precipitates uniformly on the surface of skin explants placed at the basis of the
- Pollubox®, avoiding any systemic contamination of the samples.
- 15 HSE controls received the mix of OS used to solubilize the pollutants. The final
- 16 concentrations of OS were: Nitric acid 1.7% (v/v), DMSO 8.5% (v/v) and ethanol 4% (v/v).
- 17 For transcriptomic study (microarray or qPCR), HSE were harvested 24h after P or OS
- 18 exposure. For immunostaining studies and general morphology evaluation by optical and
- transmission electron microscopies, HSE were harvested 48h after P and OS exposure. For
- 20 diesel particle penetration analysis by tape-stripping assay (see § 2.8), skin explants were
- 21 treated in the Pollubox® with a solution containing diesel particles at 0.1% dissolved in OS
- and sampled 24 hours later.

23 *2.3 Sampling*

- On day 0, 3 explants from the batch T0 were collected and cut in 3 parts: one third was frozen
- at -80°C for immunostaining, another third was fixed in formol solution for evaluation of the

- skin morphology, and the third part was preserved in RNAlater (Qiagen, 76106) for the
- transcriptomic study. On day 6 and day 7, respectively, 24h and 48h after pollutant exposure,
- 3 skin explants were processed in the same way as on day 0.

4 2.4 Optical microscopy analysis

- 5 The observation of the general morphology was evaluated after staining of formol-fixed
- 6 paraffin-embedded (FFPE) skin sections according to Masson's trichrome protocol, Goldner
- variant. The primary antibodies used in immunohistochemistry are listed in Table S3.
- 8 For all the primary antibodies, a pre-diluted horse serum (Vector laboratories, ref. PK7200)
- 9 and a universal horse secondary antibody were used (Vector laboratories, ref. PK7200).
- All the microscopical observations were performed using a Leica DMLB or a BX43 Olympus
- microscope. Pictures were digitized with an Olympus DP72 camera and the Cell^D data
- storing software.

13 2.5 Gene Expression Profile

- 14 Total RNAs were extracted from skin explants using the ReliaPrep Tissue Miniprep kit
- 15 (Promega Z6111) after mechanical disruption and homogenization by TissueLyser (Qiagen).
- 16 RNAs (70ng) of each explant from 4 donors (V1 to V4 in Table S1) were used for reverse
- transcription, amplification and Cy3 labeling, using the Low Input Labeling kit, one-color
- 18 (Agilent Technologies). All cRNAs were hybridized to human whole genome oligo
- 19 microarrays (Agilent Technologies V3 AMADID072363), which contains 60.000 probes,
- derived from the National Center for Biotechnology Information Reference Sequence (NCBI)
- 21 RefSeq. Microarray data were quantified (GeneExtraction Feature V10.7) and normalized
- 22 with R tools (Bioconductor) and deposited in the public domain (GEO Submission
- GSE126440). Subsequently, fold-changes (FC) (Table S4) were deduced from the calculated
- ratios: gene intensity of treated samples by P versus gene intensity of control samples treated
- by OS. 79 upregulated genes were selected for a FC≥1.45 and 68 downregulated genes for a

- FC \leq 0.65. Only annotated genes presenting intensity values \geq 50 (that included those higher
- than background) for treated conditions were conserved in our analysis.
- 3 Selected genes were subjected to functional analysis by PredictSearch® in order to identify
- 4 the induced biological effects. PredictSearch® is a powerful text mining software that
- 5 identifies correlations between genes and biological processes/diseases across all scientific
- 6 publications cited in the PubMed database (Benech and Patatian 2014; Eyles et al. 2007;
- 7 Michel et al. 2017).

8 2.6 Term enrichment analysis

- 9 Terms related to the set of genes modulated by P were issued from PredictSearch® analysis
- according to their p-values. For each term, the ratio (C=A/B) represents the number of term-
- related genes (A) among the number of selected genes (B). This calculation was compared to
- the ratio (F=D/E) representing the number of term-related genes (D) among all the 20 080
- 13 genes (E) contained in the PredictSearch® Database. The ratio (I=C/F) provides an
- enrichment value that was further normalized through the calculation of a ratio (K=J/F) based
- on the ratio (J=G/H) representing the number of term-related genes (G) among a set of
- randomly chosen genes (H) of the same size than the set of our gene selection (B) compared
- again to the F ratio. This new ratio (K) is then compared to the I ratio to give a normalized
- 18 enrichment score (NES) defined by the I/K ratio.

19 2.7 Laser capture micro-dissection (LCM), RNA extraction from micro-dissected explants

20 and real-time qPCR

- 21 According to Percoco et al. (2012), the *stratum granulosum* (SG) and the *stratum basale* (SB)
- of the epidermis were micro-dissected by LCM and total RNAs were extracted using an
- 23 RNeasy Micro kit (Qiagen) following the manufacturer's instructions. The quality and
- quantity of the RNAs were assessed by microfluidic capillary electrophoresis (Agilent 2100
- 25 Bioanalyzer, Agilent Technologies). Amplification reactions were performed using a

- 1 QuantStudio 12K Flex Real-time PCR System (Applied Biosystems). When possible, intron-
- 2 spanning primers were designed using Primer Express 3.0.1 software (Applied Biosystems;
- 3 Table S6).
- 4 **2.8 Tape-stripping experiment**
- 5 A D-Squame® disc with a 14 mm diameter (D101, CuDerm), was used (24h after Pollubox®
- 6 exposure) to remove corneccytes from 3 skin explants (from donor V10; Table S1) exposed
- 7 or not to diesel particle-enriched solution (diluted at 0.1% in OS) in the Pollubox®. The disc
- 8 covering the entire surface of the skin explants was successively applied 10 times and was
- 9 then fixed for 30 min at 4°C in a solution of 2% glutaraldehyde. After post-fixation in 1%
- 10 OsO₄ for 1h at 4°C, it was embedded in 3% low melting point agarose. Samples were
- dehydrated as follows: 35% ethanol for 10 min; 70% ethanol for 3 min; 100 % ethanol for 1
- min. Infiltration was carried out at room temperature in London Resin White (LRW, EMS)
- for 30 min. Polymerization was performed at 60°C for 24h. Ultrathin sections (70 nm) were
- 14 collected on formvar carbon-coated nickel grids, stained with UranyLess solution and lead
- citrate solution (Delta microscopies). Observations were performed with a Philips Tecnai 12
- 16 Biotwin FEI transmission electron microscope (Philips).
- 2.9 High pressure freezing (HPF) and freeze substitution (FS)
- 18 For ultrastructural and immunogold analyses, 48h after Pollubox® exposure, triplicate skin
- explants from a healthy 39-year-old Caucasian women (donor V4, table S1), exposed to OS or
- 20 P in the Pollubox®, were cryofixed by HPF as previously described (Percoco et al. 2013).
- 21 Briefly, specimens were cut into small discs (≈3 mm in diameter and 300 μm thick) and
- 22 cryofixed in a high-pressure freezer (HPM100, Leica). After high pressure freezing, samples
- were cryosubstituted with the automate freeze substitution (AFS-FSP, Leica) and embedded
- 24 in LRW resin as described by Percoco et al. (2013). Ultrathin sections (70 nm) were
- processed and observed as mentioned in § 2.8.

2.10 Immunogold labeling

- 2 Ultrathin sections of skin explants were treated with 0.1M glycine for 10 min and incubated
- 3 for 30 min with 10% normal goat serum (NGS) in PBS + 0.1% Tween 20 + 1% BSA.
- 4 Samples were incubated with primary and secondary antibodies as follows: 1h at 25°C,
- 5 mouse-anti-AhR (1/5; MA1-514, Thermo Fischer; Table S3) in saturation buffer (0.1%
- 6 Tween 20 + 1% BSA + 1% NGS in PBS); goat anti-mouse antibody coupled to 25 nm gold
- 7 particles (1/20; Aurion) in PBS + 0.005% Tween 20.

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2.11 Statistical analysis

- 10 A Two-way ANOVA (R, Mkmisc package) was performed for p-values (Pval) evaluation
- using a design based on each probe including treatment factor (P vs OS) and volunteer factors
- 12 (V1-V2-V3-V4) in order to take into account the heterogeneity between volunteers. For fold-
- change computing (P vs OS) we applied the same design using the removeBatchEffect
- 14 function from the Limma package. Only p-values (P) less than 0.05 were considered
- 15 significant.
- The ratio of gene expression between OS and P treatment in SB and SG enriched fractions
- was determined with the $\Delta\Delta$ Ct method and were calculated using the OS expression profile as
- a reference. First, the Shapiro-Wilk normality test allowed to determine which data followed a
- 19 normal distribution. In case of a Gaussian distribution, the data were analyzed statistically by
- 20 unpaired t-test with Welch's correction if variances were non-equal (GraphPad PRISM v7.0,
- 21 GraphPad Software Inc). In case of a non-Gaussian distribution, the data were analyzed using
- 22 the Mann-Whitney test (*P <0.1; ** P <0.05). All data were shown as mean \pm SEM
- 23 calculated from three independent experiments.

2. RESULTS AND DISCUSSION

To investigate the skin alterations that might be caused by a pollution mixture, we combined different molecules according to the French non-profit organization 'Air Parif' (Table S2). The resulting pollutant mixture (P) was nebulized on the cultured explants into a dedicated chamber, the Pollubox® (Figure S2). A mixture of the different OS used to solubilize the pollutants served as control (Table S2). The impacts of P either on skin morphology, or on gene and protein expression were investigated on skin explants originating from several donors (Table S1). The optimal dose allowing the detection of an effective cutaneous response to pollutant stress was determined through the evaluation of the induced expression of genes well known to be involved in the detoxification process such as CYP1A1, GPX2, *HMOX1*, and *SQSTM1* (Figure S1).

3.1 Effect of pollutant exposure on skin morphology

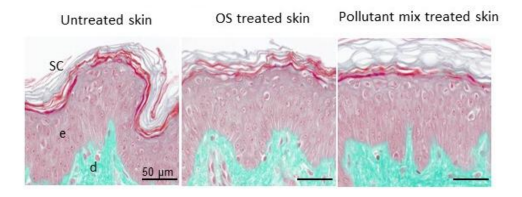


Figure 1: Skin morphology 48 h after pollutant mix exposure.

General morphology was analyzed on FFPE skin sections stained using Masson's trichrome protocol. e, epidermis; d, dermis; OS, organic solvent; SC, *stratum corneum*. Scale bars: 50 um.

P exposure did not lead to global skin morphology alterations (Figure 1). Nevertheless, TEM analysis on explants treated with P revealed an increase of extracellular vesicle (EV)-like structures in the epidermis (Figure 2).

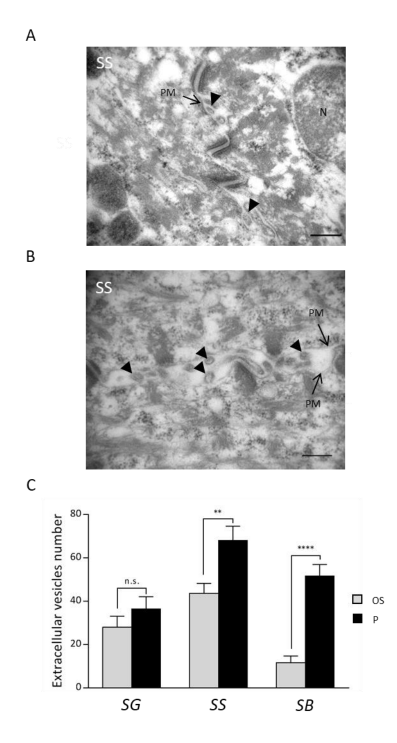


Figure 2: Observation of extracellular vesicle-like structures by transmission electron

3 microscopy (TEM).

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- 4 Ultrathin sections of skin explants 48h after treatment with OS or P were observed by TEM.
- 5 Electron micrographs of keratinocytes from *stratum spinosum* after OS (**A**) or P exposure (**B**).
- 6 Extracellular vesicle-like structures are indicated by black arrowheads. C, Quantification of
- 7 extracellular vesicle-like structures observed by TEM was performed on 10 cells per
- 8 epidermal layer. **** P < 0.0001; **P < 0.01; n.s.: not significant. N: nucleus; OS: organic
- optioning rayor. I volot, its inot significant. It indicates, ob. organic
- 9 solvent; P: pollutant mix; PM: plasma membrane; SB: stratum basale; SG: stratum
- 10 granulosum; SS: stratum spinosum. Scale bars: 250 nm.

Quantification revealed a significant increase of EV-like structures in *stratum spinosum* (SS) 1 2 and in SB but not in SG after P exposure (Figure 2C). These observations were in line with a study showing that particulate matter exposure can trigger an increase of EV release (Bonzini 3 et al. 2017). Interestingly enough, EVs were found to be involved in tissue remodeling and 4 inflammation modulation in lung after pollutant exposure (Benedikter et al. 2018). Cells 5 6 release into the extracellular environment diverse types of membrane vesicles that originate 7 from endosomal and plasma membranes called exosomes (30 to 150 nm) and microvesicles/microparticles (100 to 1000 nm), respectively (Raposo and Stoorvogel, 2013). 8 The size of EV-like structures observed in our study was less than 100 nm, suggesting these 9 10 structures were exosomes, structures discovered in 1985 by Pan et al. (Pan et al. 1985). However, these EV-like structures may also correspond to microvesicles budding from the 11 keratinocyte surface and, in part, to transversally cut cell surface villosities. To define more 12 13 precisely such EV-like structures, immunolabelling with specific markers should be further performed. The EVs are considered as critical mediators of cell communication (Pleet et al. 14 15 2018) regulating several genes in various cell types including dermal fibroblasts (Huang et al. 2015). Recent studies demonstrate that changes of exosomal cargo depend on external stimuli 16 17 with consequences for the receiving cells (Rokad et al. 2019). Because of their ability to 18 transfer biologically active molecules such as proteins, nucleic acids, and lipids, exosomes play a major role in influencing numerous physiological as well as pathological functions in 19 20 response to environmental toxicants. These toxicants can be heavy metals (Rokad et al. 2019) 21 as those contained in our P mix. TEM analysis was also performed on corneccytes previously exposed to diesel particle-22 enriched solution and collected by tape-stripping (Figure S4 A and B). 23 TEM analysis was also used to observe diesel particles within the explants. First, the upper 24 epidermal layers of explants exposed to diesel particle enriched solution, and TEM analysis 25

- 1 was performed on cornecetes collected by tape-stripping (Figure S4). Some dense electron
- element smaller than 300nm were seen in the cornecytes of exposed explants, while they
- were absent in cornecytes from explants of the same donor exposed to diesel-free particles
- 4 solution. Measurements of these particles showed a size of 2.5 nm or smaller. Similar
- 5 particles elements were observed even down to the SG in ultrathin sections of other explants
- 6 from the same donor 48h after the P nebulization in the Pollubox® (Figure S4 C-F). We
- 7 hypothesize that these electron dense elements could come from diesel particles. More
- 8 investigations such as energy dispersive X ray analysis will be necessary to identify them.
- 9 demonstrating that diesel particles could penetrate the skin explants.

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3.2 Effect of P on gene expression in cutaneous cells

3.2.1 Gene selection and the functional analysis by PredictSearch®

13 To evaluate whether our device was indeed efficient to observe the effect of P on HSE at a

gene expression level on P-induced relevant targets, a transcriptomic analysis using whole

human genome microarrays was performed. Explants generated in triplicate from 4 donors

were either left untreated or exposed to P or OS for 1h30 and then maintained in culture for

24h before RNA extraction. Only intensity values higher than background according to

Agilent calculations were considered. Fold changes (FCs) were calculated from the ratios

between P and OS exposed explants for all triplicates. For each donor and each triplicate,

FC \geq 1.45 and \leq 0.65 were used to determine up-regulation and down-regulation, respectively.

The mean of these FCs among triplicates was then calculated. This selection led to 70 unique

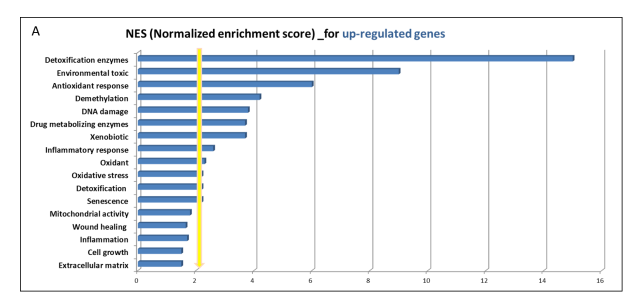
up-regulated and 68 unique down-regulated genes (see supplemented data Table S5) common

to all 4 donors. The different biological processes and pathways significantly associated with

the differentially expressed genes common to all donors were identified using PredictSearch®

(Figure 3A and 4B). For each of the terms associated with these processes and pathways, the

- 1 enrichment values were calculated (see material and methods) and classified from the highest
- 2 to the lowest score. Only scores greater than or equal to 2 were considered significant.



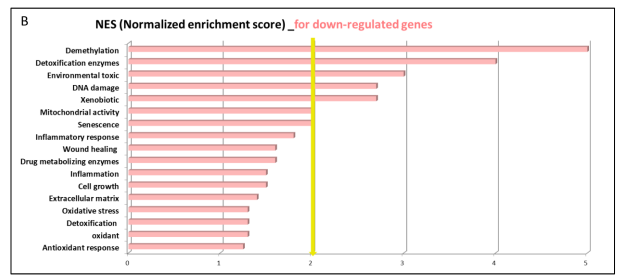


Figure 3: Enrichment of biological terms

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- 6 Terms related to the selected genes were identified using PredictSearch® software.
- 7 Classification was achieved according to the normalized enrichment score (NES) as defined
- 8 in materials and methods. NES exhibiting a value ≥ 2 are indicated by the yellow arrow. A:
- 9 up-related genes; **B**: Down-regulated genes.
- 10 As shown in Figure 3, the term "detoxification enzymes" appeared at the first position for the
- up-regulated genes (Figure 3A) and at the second position for the down-regulated genes
- 12 (Figure. 3B). This term was followed by other terms such as "environmental toxic", "DNA
- damage", "antioxidant response", "xenobiotics" and "demethylation" confirming some of the
- 14 known effects described upon treatment with pollutants. These results were a confirmation not

only of our experimental design but also of the robustness of our transcriptomic data. Of note, 1 2 the observation that down-regulated genes shared some terms with up-regulated genes might suggest that at 24h a negative feedback occurred for these specific genes to decrease their 3 4 expression. For instance, the term "demethylation" might refer, at least in part, to the ability of enzymes such as cytochrome P450 monooxygenases (CYPs) to metabolize drug via N-5 6 demethylation (Zanger and Schwab 2012). In addition, epigenetic activities such as DNA 7 methylation are known to be involved in the transcriptional activity of genes encoding enzymes that participate in detoxification and drug metabolism (Zhang et al., 2010). On the 8 other hand, ALDH5A and ALDH7A1 encode enzymes involved in the detoxification of 4-9 10 hydroxy-trans-2-nonenal (HNE) implicated in the pathogenesis of numerous neurodegenerative disorders (Murphy et al. 2003), and aldehydes generated by alcohol 11 metabolism and lipid peroxidation (Brocker et al. 2010; Chan et al. 2011), respectively. 12 13 Mammals have developed oxidative systems that eliminate endogenous and foreign toxic compounds. In humans, this oxidative detoxification depends on the activity of CYPs such as 14 15 CYP2J2 and CYP4B1. Thus, a repressed expression of these genes might prevent an 16 exacerbation of the response and/or a loss of protection against potential harmful metabolites produced by detoxification. Nevertheless, the functional terms correlated to these genes 17 18 supported that our device and cultured explants were efficient to elicit a response to pollutants. 19

3.2.2 Effect of P on the AhR pathway: Xenobiotic responses

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Based on the functional correlations found by Predictsearch®, the selected genes were integrated within specific pathways. One of them was related to the AhR-target gene activation. A significant number of genes found modulated by P encoded phase I and II detoxifying enzymes (Figure 4 and Table S4).

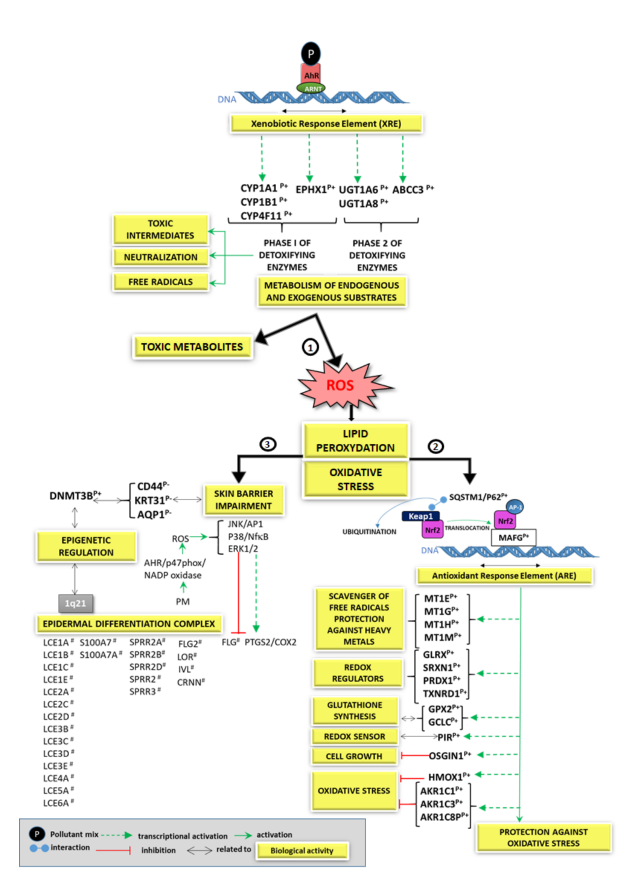


Figure 4: Schematic representation of functional networks deduced from PredictSearch® analysis of the set of genes modulated in HSE 24h after P exposure.

Induced genes or repressed genes are noted by P+ or P- in superscript, respectively. Opposite modulation of skin barrier function-associated genes among volunteers are indicated by # in

superscript. 1) Genes of the xenobiotic response; 2) genes of the antioxidant response; 3) genes related 1 2 to the skin barrier function. 3 The AhR is an evolutionarily conserved receptor that is widely expressed in many organs 4 5 including brain, liver, lung, and skin (Baker et al. 2018; Guastella et al. 2018; Napolitano and 6 Patruno 2018). Once activated by xenobiotic ligands, including dioxin and polycyclic 7 aromatic hydrocarbons, AhR translocates to the nucleus and dimerizes with co-factors 8 including the aryl hydrocarbon receptor nuclear translocator (ARNT), and binds to XREs present in AhR-responsive genes thus increasing their expression (Denison et al. 2011). 9 10 Interestingly, immunogold analysis on ultrathin sections of skin explants exposed to P compared to OS (Figure S3) showed a significant increase of the nuclear AhR location in 11 granular keratinocytes. 12 13 Among the up-regulated genes following P exposure and AhR activation identified by microarrays (Figure 4), the proteins encoded by CYP1A1 and CYP1B1 metabolize and 14 15 detoxify carcinogens, drugs, environmental pollutants and ROS (Başak et al. 2017). CYP1A1 is expressed by epidermal keratinocytes, dermal fibroblasts, sebaceous glands, hair follicles 16 and subcutaneous striated muscles in normal skin. Other members of the cytochrome family 17 18 like CYP4F11 whose gene was induced by P, metabolize compounds into irreversible inhibitors of stearoyl CoA desaturase (SCD). SCD is essential to sebocyte development and 19 consequently SCD inhibitors cause skin toxicity (Theodoropoulos et al. 2016). Similarly, 20 21 induction of genes encoding proteins of phase II (UGT1A6-8 and ABCC3) was also observed in response to P (Table S4). Unlike for CYPs, few studies have demonstrated the expression 22 of UGT1A6 and especially UGT1A8 in human skin cells (Diawara et al. 1999; Sumida et al. 23 2013). Our transcriptomic results showed that the expression of both genes can be induced in 24 HSE after exposure to pollutants and the presence of the corresponding proteins was 25 26 confirmed by immunostaining in skin explants 48h after P exposure (Figure 5A).

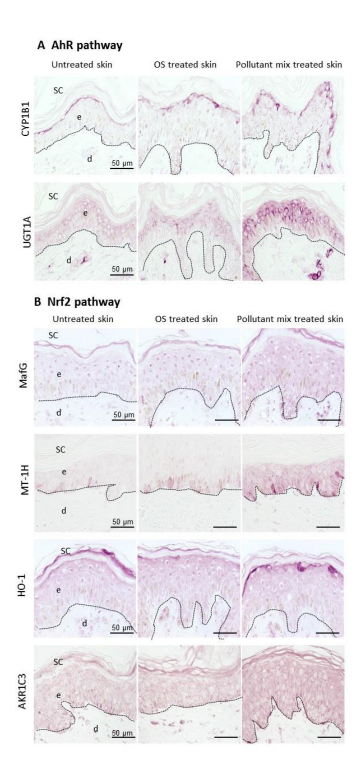


Figure 5: Skin morphology and protein immunolocalization after 48 h following pollutant mix exposure.

Left column: untreated skin samples; central column: skin explants treated with OS; right column: skin explants treated with pollutant mix. For each protein used for immunolabelling, the picture shown in this figure is one representative out of at least 9 images. e: epidermis; d: dermis; OS: organic solvent; SC: *stratum corneum*. Scale bars: 50 µm.

In carcinogenic cells or in liver (Bock and Bock-Hennig 2010; Bock and Köhle 2005), UGTs 1 2 attenuates the generation of mutagenic benzopyrene metabolites, thus facilitating their detoxification. Accordingly, an induced expression of some of the UGT genes (Figure 4) was 3 4 observed in our present study. Similarly, the overexpression of one member of the ABCC gene family, ABCC3, which was described in skin (Osman-Ponchet et al. 2014; Takenaka et 5 al. 2013) was also detected (Figure 4). Genes coding for glutathione/glucuronide sulfate 6 transporters such as ABCC1, ABCC3, and ABCC4 encoding multidrug resistance proteins 7 have been reported to be strongly upregulated during keratinocyte and HaCaT cell 8 differentiation. These proteins are also involved in the translocation of sulfated lipids during 9 10 SC formation (Kielar et al. 2003). In addition to their role in drug resistance and epidermal lipid layer reorganization, there is substantial evidence that these efflux pumps have 11 overlapping functions in tissue defense and are able to transport a vast and chemically diverse 12 13 array of toxicants. These toxicants include bulky lipophilic, cationic, anionic, and uncharged drugs and toxins as well as conjugated organic anions that encompass dietary and 14 15 environmental carcinogens, pesticides, metals, metalloids, and lipid peroxidation products (Leslie et al. 2005). 16 Related to detoxification processes, EPHX1 coding for the epoxide hydrolase was also 17 significantly induced by P (Figure 4). EPHX1 is a critical biotransformation enzyme that 18 19 converts epoxides from the degradation of aromatic compounds to trans-dihydrodiols, which can be conjugated and excreted from the body. Lipid mediators such as arachidonic acid-20 21 derived epoxyeicosatrienoic acids, produced by cytochrome P450 epoxygenases, are hydrolyzed by EPHX1 and contribute to tissue growth and wound epithelialization (Edin et al. 22 2018; Panigrahy et al. 2013). 23

24 3.2.3 Effect of P on the NRF2 pathway: Antioxidant responses

The second group of modulated genes illustrated the activation of a NRF2-dependent 1 2 antioxidant response (Figure 5) that follows the production of ROS likely in part generated by CYPs (Table S4). Exposure of cells to xenobiotics, drugs or ionizing radiation is known to 3 generate ROS and electrophiles that lead to oxidative and electrophilic stresses, and has a 4 profound impact on the survival of all living organisms (Kasai 1989; Meneghini 1997). The 5 6 cells respond to oxidative/electrophilic stresses by activating defense mechanisms that result 7 from the coordinated induction of a battery of genes to protect cells (Dhakshinamoorthy et al. 2000). 8 The activation of the NRF2 pathway in response to pollutants can be positively regulated by 9 10 SQSTM1/p62 (Figure 4), which mediates sequestration of Keap1 and promotes translocation of NRF2 to the nucleus (Lau et al. 2013). After translocation, NRF2 forms heterodimers with 11 other transcription factors such as those encoded by genes of the MAF family (MAFB, 12 13 MAFG), which bind to ARE, found in promoters of various detoxifying/defensive genes including those described above (Hirotsu et al. 2012; Katsuoka et al. 2005). It should be noted 14 that MAFG immunostaining is significantly enhanced in epidermal cells after P exposure 15 (Figure 5B). SQSTM1 is also a target gene for NRF2 and creates a positive feedback loop by 16 inducing ARE-driven gene transcription (Jain et al. 2010; Mildenberger et al. 2017). 17 18 Among NRF2 target genes, the induction of several members of the metallothionein gene family (MT1E, MT1H, MT1G, MT1M), which elicited the highest modulation as well as genes 19 involved in antioxidant response (GPX2, GCLC, HMOX1) was detected (Figure 4 and Table 20 S4). Among them, the positive modulation in response to P of MAFG, MT-1H and HO-1 was 21 22 confirmed at the protein level by immunohistology (Figure 5B). Metallothioneins are cysteine-rich low molecular weight metal-binding proteins with multiple functions such as 23 cell protection against oxidative stress and heavy metal toxicity as well as the regulated 24 balance of essential metals (Cu and Zn) (Ochiai et al. 2008; Sato and Kondoh 2002). 25

- 1 Glutathione peroxidase 2 (GPX2) catalyzes the reduction of organic hydroperoxides and
- 2 hydrogen peroxide by glutathione, and thereby protects cells against oxidative damage. The
- 3 reduction of organic compounds by glutathione can be potentiated through overexpression of
- 4 a key enzyme encoded by GCLC which contributes to glutathione synthesis.
- 5 HMOX1 (HO-1) decreases lipid peroxidation and inhibits the induction of ROS scavenging
- 6 proteins (Zhang et al. 2012) after an oxidative stress. HMOX1 may also exert anti-
- 7 inflammatory activities by suppressing the TNF or INF-induced ICAM1/CD54 expression and
- 8 subsequent monocyte-keratinocyte adhesion (Seo et al. 2011; Seo et al. 2010).
- 9 Aldo-ketoreductase 1Cs (AKR1Cs) enzymes catalyze the NADPH-dependent reduction of
- 10 ketosteroids to hydroxysteroids and are members of Phase I metabolizing enzymes involved
- in the metabolism of steroids (C1-3), prostaglandins (C3), polyaromatic hydrocarbons (C1-3)
- and xenobiotics (C1-2-4) (Penning and Byrns 2009). They are related to tobacco-
- carcinogenesis since they activate polycyclic aromatic trans-dihydrodiols to yield reactive and
- 14 redox active o-quinones. They also detoxify reactive aldehydes derived from exogenous
- toxicants, e.g., aflatoxin, endogenous toxicants, and those formed from the breakdown of lipid
- peroxides. AKRs are stress-regulated genes and play a central role in the cellular response to
- osmotic, electrophilic and oxidative stress (Jin and Penning 2007; Palackal et al. 2002). In line
- with an induced expression of AKR1C1 and AKR1C3 genes detected by microarray (Figure 4,
- Table S4), AKR1C3 was increased at the protein level after P exposure (Figure 5B). In
- addition to their role in xenobiotic detoxification, AKR1C1 and AKR1C3 are known to elicit
- an inhibitory effect on oxidative stress (Matsunaga et al. 2013).

22 3.2.4 Impact of the pollutant mix on the epidermal terminal differentiation complex

- Unlike genes of the xenobiotic response, the modulation of the genes located within the
- 24 epidermal differentiation complex (EDC) locus (Marenholz 2001) was much more
- 25 heterogeneous among donors (Figure 4, Table S4). Expression of the majority of these genes

was induced in donor V1 and V2, while their expression was mostly unchanged in donor V3 1 2 and even reduced in donor V4. It has been reported that particulate matter induces expression of PTGS2/COX2 and represses FLG expression (Lee et al. 2016). These opposite 3 modulations, dependent on AhR signaling and ERK1/2, p38/NF-KB and JNK/AP1 activation, 4 were associated with skin barrier alteration. In our study, while a modulation of PTGS2/COX2 5 6 was not observed, FLG expression was effectively suppressed or unchanged in donors V3 and 7 V4, respectively, but induced in donors V1 and V2 (Table S4). In addition to FLG, a similar differential pattern among donors was observed with genes encoding FLG2 (filaggrin family 8 member 2), several members of the late cornified envelop protein family (LCE1A, LCE1B, 9 10 LCE1C, LCE2C, LCE2D, LCE3D, LCE4A, LCE5A and LCE6A), and of the small proline rich protein family (SPRR3, SPRR2A, SPRR2B, SPRR2D, SPRR2G). These genes together 11 with IVL (involucin), LOR (loricrin), CRNN (cornulin), and S100A7/PSOR1 (S100 calcium 12 13 binding protein A7/psoriasin) induced only in donors V1 and V2 belong to the EDC locus on chromosome 1q21.3 (Kypriotou et al. 2012). Genes of this locus encode proteins involved in 14 15 terminal differentiation and cornification of keratinocytes. To explain such differences between volunteers, it can be suggested that the differentiation follows different kinetics 16 depending on their genetic background or an epigenetic control of gene expression. Indeed, 17 18 this type of control plays an essential role in regulating stem cell maintenance by repressing the expression of differentiation genes while allowing cell-cycle progression and cell renewal 19 (Goldberg et al. 2007; Spivakov and Fisher 2007). 20 21 Local deposition and removal of DNA methylation are tightly coupled with transcription factor binding, although the relationship varies with the specific differentiation process. 22 A recent study in skin has confirmed a critical role of epigenetic modification for EDC gene 23 transcription (Perdigoto et al. 2014) Therefore, it is possible that the reduced expression of the 24 EDC genes as seen in donors V3 and V4 was donor-dependent under epigenetic control and 25

- contributed to an alteration of the differentiation process and consequently of the skin barrier.
- 2 It can also be postulated that even in the presence of an efficient activation of a xenobiotic
- 3 response (equally activated in all 4 donors) to the pollutants, the keratinocyte differentiation
- 4 process could restore the homeostasis of the cutaneous barrier only later on in donors V3 and
- 5 V4. Such a difference may rely on a reduced anti-oxidant response illustrated in donors V3
- 6 and V4 eliciting a lower expression of GPX2, AKR1C, AKR1C3, AKR1C8P, HMOX1, SRXN1,
- 7 PRDX1, TXNRD1, MT1E, MT1H, MT1G and MT1M (Table S4) when compared to the
- 8 expression of these genes in donors V1 and V2.
- However, repression of some skin integrity key genes such as AQP1 (aquaporin 1), CD44 9 10 (receptor for hyaluronic acid), and KRT31 (keratin 31) was observed in all donors (Figure 4, 11 Table S4). This repression might indicate that P treatment compromised cutaneous barrier 12 homeostasis although the recovery to homeostasis might be different among donors. Indeed, it was shown that defective channels like those encoded by aquaporin genes lead to an impaired 13 skin barrier (Blaydon et al. 2014). Moreover, it has been reported that the invalidation of 14 CD44 in mouse skin leads to a delay in the early barrier recovery following barrier disruption 15 (Bourguignon et al. 2006). In addition, CD44 downregulation has also been related to 16 inhibited hyaluronan-mediated keratinocyte differentiation. Moreover, the repression of AQP1 17 and CD44 genes was found to be subjected to epigenetic modification as a result of increased 18 19 methylation by DNMT3 (Smith et al. 2019; Woodson et al. 2006). To investigate the involvement of this process, a methylation profile in HSE upon P treatment will have to be 20 21 further performed to correlate gene expression and their location within

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hypo/hypermethylated regions.

- 1 Nevertheless, altogether these transcriptomic results supported that our experimental
- 2 conditions and testing device can evaluate the effects of pollutants on HSE through the
- 3 modulation of genes of the xenobiotic and anti-oxidant responses as well of EDC genes.

3.2.5 Skin layer-specific response

- To gain more insight in the specific response of the different cell layers following P exposure,
- 6 we isolated by laser capture micro-dissection SB- and SG-enriched fractions from full-
- thickness explants (Percoco et al., 2012). After RNA extraction from each fraction, RT-qPCR
- 8 was performed to evaluate the expression of xenobiotic and anti-oxidant response genes. In
- 9 accordance with microarray and immunohistochemical data, both AhR and NRF2 pathways
- were modulated (Figure 6). As shown in Figure 6A, P exposure led to a significant increase of
- these genes in the SB-enriched fraction.
- In the SG-enriched fraction (Figure 6B) a significant upregulation for the same genes but to a
- lesser extent than in the SB-enriched fraction was observed. These data suggested that 24h
- after P exposure a more efficient transcriptional activity was detectable in the deepest layer of
- the epidermis than in the granular layer. The response within the deep layers of the epidermis
- indicated that either some of the pollutant compounds passed through the tissue and/or that
- 17 the presence of P on the epidermal surface induced a cascade of signaling events from the
- surface to the deeper layers. Using ¹⁴C-labelled benzo[a]pyrene applied on the skin of mice,
- 19 Yang et al. (1986) showed that hydrocarbons can be recovered in urine, feces and tissues
- suggesting that these types of pollutants can cross the dermo-epidermal barrier. In contrast,
- 21 following skin explant exposure in the Pollubox®, we did not detect diesel particles beyond
- 22 the SG cells. Furthermore, the number of EV-like structures in SS and in SB but not in the SG
- 23 (Figure 2) suggested also that the deepest layers of the epidermis were able to react to P
- exposure. While the AhR and NRF2 pathway genes are significantly overexpressed in the SB,
- 25 the AhR gene was weakly overexpressed or even downregulated in the SG. Interestingly,

detection of AhR receptors in the nucleus of SG cells increased significantly 48h after

2 exposure to P, showing that SG cells were also involved in the P response. It would be

interesting to perform these two types of transcriptomic and microscopic analyses at other

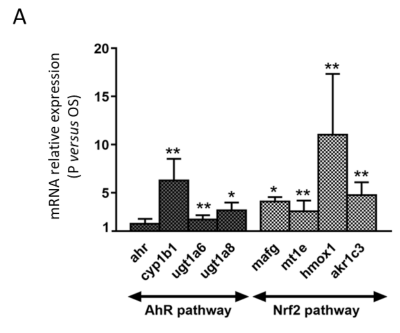
kinetic points in order to investigate the spatial temporal response of keratinocytes to

5 pollutants.

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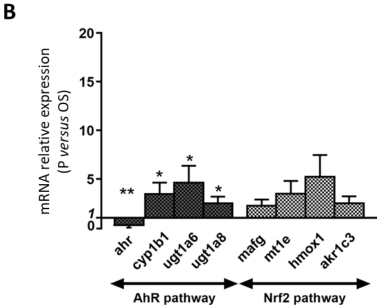


Figure 6: Ratio (P versus OS) of gene expression using RNAs extracted from microdissected layers of skin explants 24h after P exposure. The ratios of gene expression between P versus OS treatment in SB (A) and SG (B) enriched fractions were determined

with the ΔΔCt method. SB: *stratum basale*; SG: *stratum granulosum*; OS: organic solvents; P:

pollutant mix. Mean \pm SEM are represented. *P < 0.1; **P < 0.05 (3 explants each from 4

donors were analyzed).

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3. CONCLUSION

6 Our results demonstrated that efficient xenobiotic and antioxidant responses were triggered in human skin explants exposed in a dedicated chamber to a mix of urban pollutants. Although 7 8 the dose of each compound contained in the pollutant mixture was far higher than those encountered in a real environment, no morphological alteration of the treated explants (until 9 48h after exposure) was observed. From our results based on the cutaneous explant model, it 10 can be postulated that diesel particles, smaller than 300 nm, can penetrate the skin down to the 11 granular layer. In contrast, The other Pollutants were able to reach the deepest epidermal 12 layers to modify expression levels of several genes and to increase extracellular vesicle-like 13 14 secretion. The presence of these vesicles might indicate intensive cross-talk between cells. Even though no definitive conclusion can be drawn from 4 donors, we found some differences 15 among them between the transcription profiles of genes related to either the 16 xenobiotic/antioxidant responses or the EDC. Most of the genes of the xenobiotic and 17 antioxidant responses were induced in all donors although two donors exhibited a higher 18 19 induction of these genes than the two others. Interestingly, for these last two donors, no induction (even a repression for one of them) was seen nearly for all the EDC genes while 20 21 these genes were significantly induced in response to P in the two other donors. Since 22 expression of EDC genes is important for maintaining the skin barrier integrity, the absence of 23 induction or repression of these genes after pollutant exposure might possibly lead to a 24 deleterious effect on terminal differentiation. Such a hypothesis should be investigated further 25 as well as the involvement of individual traits influencing a tissue response to P. Indeed, it can be postulated that the impact of pollutants on terminal differentiation generated by a signal in 26 the deepest layer of the epidermis can vary depending on individual specificities. However, in 27

order to confirm whether or not gene profiles can be used to assess the skin's ability to resist 1 pollutants and possibly skin aging, explants of more donors should be tested. The comparison 2 3 of gene expression data with results from microscopic and physico-chemical analyses among donors will have to be systematically performed. Nevertheless, our results demonstrate that 4 5 these experimental conditions (the P mixture), the tissue model (human skin explants), and the 6 device (Pollubox®) provide a suitable approach to study ex vivo the effect of pollutants on skin. This approach can be considered as a new tool to test the potential preventive and/or 7 8 curative effects of dermo-cosmetic ingredients or end products.

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13 AUTHORS CONTRIBUTION

- 14 Patatian A, Delestre-Delacour C, Percoco G, Ramdani Y, Di Giovanni M, Bader Th and
- 15 Bénard M performed the research;
- 17 Patatian A, Delestre-Delacour C, Percoco G, Elian E, Benech P and Follet-Gueye ML
- designed the research study;
- 20 Peno-Mazzarino L, Driouich A and Lati E contributed to provide essential reagents and tools;
- 22 Patatian A, Delestre-Delacour C, Percoco G, Ramdani Y, Benech P, Follet-Gueye ML
- analyzed the data;
- Patatian A, Delestre-Delacour C, Percoco G, Benech P and Follet-Gueye ML wrote the paper
- 27 Benech P and Follet-Gueye ML are the senior co-authors.

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