Waterpipe smoke condensate influences epithelial to mesenchymal transition and interferes with the cytotoxic immune response in non-small cell lung cancer cell lines

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Abstract. Waterpipe tobacco smoking (WPS) continues to spread globally and presents serious health hazards. The aim of the present study was to investigate the effects of treatment with WPS condensate (WPSC) on lung cell proliferation and plasticity as well as tumor cell recognition and killing by natural killer (NK) cells using cytotoxicity assays. The results indicated that exposure of normal and cancer lung cell lines to WPSC resulted in a decrease in their in vitro growth in a dose-dependent manner and it induced tumor senescence. In addition, WPSC selectively caused DNA damage as revealed by an increase in γH2AX and 53BP1 in tumor lung cells. To gain further insight into the molecular mechanisms altered by WPSC, we conducted a global comprehensive transcriptome analysis of WPSC-treated tumor cells. Data analysis identified an expression profile of genes that best distinguished treated and non-treated cells involving several pathways. Of these pathways, we focused on those involved in epithelial to mesenchymal transition (EMT) and stemness. Results showed that WPSC induced an increase in SNAI2 expression associated with EMT, ACTA2 and SERPINE2 were involved in invasion and CD44 was associated with stemness. Furthermore, WPSC exposure increased the expression of inflammatory response genes including CASP1, IL1B, IL6 and CCL2. While immune synapse formation between NK and WPSC-treated lung cancer target cells was not affected, the capacity of NK cells to kill these target cells was reduced. The data reported in the present study are, to the best of our knowledge, the first in vitro demonstration of WPSC effects on lung cellular parameters providing evidence of its potential involvement in tumor physiology and development.

Introduction

Smoking is the leading cause of chronic obstructive pulmonary disease, a progressive and eventually debilitating lung disease (1). Furthermore, smokers of tobacco are 20-40 times more at risk of developing lung cancer in comparison to non-smokers (2).

Tobacco can be smoked using different ways, two common ways include the use of cigarettes or waterpipes. Several studies have investigated the chemical composition of cigarette smoke and more than 5,000 bioactive chemical compounds have been isolated, including over 60 carcinogens (3). Waterpipe smoke analysis has revealed that it contains significant concentrations of toxicants including 27 known or suspected carcinogens thought to cause dependence, heart disease, lung disease and cancer (4). Despite the adverse health effects among users, waterpipe usage is growing in popularity and public health interventions remain below what is present for cigarette smoking (5).

Several studies have addressed the in vivo effects of WPS on waterpipe smokers’ health. Smokers are found to have high urinary concentrations of several toxins including carcinogens (6), resulting in profound effects on lung function (7). Waterpipe smokers were also observed to have 6-fold greater risk of developing lung cancer (8). At the molecular level, DNA repair gene expression was reported to be decreased in the blood of waterpipe smokers, while DNA damage-related gene expression was increased (9). It has also been reported...
that WPS induces endothelial cell dysfunction, inflammation, and impaired repair mechanisms with implication in vascular disease (10). In this respect, nicotine, present in WPS, induces bronchial epithelial cell apoptosis and senescence via ROS-mediated autophagy-impairment (11). WPS also induces cell cycle arrest and cellular senescence mediated by the p53-p21 pathway in alveolar type 2 cell disease (10), whereas it induces apoptosis in human aortic endothelial cells (10,12). All these data highlight the damaging effects of WPS. More importantly, WPS may contribute towards EMT, tumor heterogeneity and immune escape. These processes are known to play critical roles in tumor plasticity and are important factors impacting both the diagnosis and treatment of cancer patients (13).

The aim of the present study examined the changes in tumor lung cell gene expression related to DNA damage, inflammation, EMT and stemness. In addition, the consequence of WPSC treatment on immune recognition and killing by NK cells was investigated. Our results emphasized the potential impact of WPSC on tumor lung cell behavior and provide insights into their associated transcriptomic response including DNA damage, inflammation, and cell plasticity.

Materials and methods

Waterpipe smoke sampling and analysis. Waterpipe smoke collection was performed as previously described (14). Briefly, 17.5 g double apple flavor tobacco (mou'assel) was placed in the head piece of the waterpipe which was then tightly wrapped using a perforated aluminum foil. Two pieces of quick lighting charcoal briquettes were used to heat the tobacco. The generated smoke was collected using a robotic machine (IREADY LLC) that simulates the human smoking process. The puff duration was set at 5 sec per puff with 15 sec inter-puff duration, for a total of 80 puffs per session. Collection of the smoke condensate was carried out on pre-conditioned glass wool fibers packed inside a T-shaped tube. It is important to note that under our experimental conditions, the cells were exposed to the waterpipe smoke condensate samples. To identify the chemical composition of the condensate and to eliminate any masking effect of the large glycerin peak during gas chromatography-mass spectrometry (GC-MS), successive extraction steps were performed. The extraction procedure was carried out by mixing 72.6 mg of the extract in 4 ml of toluene. The mixture was stirred for 24 h and allowed to separate. In this step, glycerin is not expected to move into the toluene layer. Then, 0.15 ml of the remaining components of the extract were dissolved in 15 ml of ethanol followed by a dilution of 1:40 in ethanol prior to gas chromatography mass spectrometry (GCMS) analysis to eliminate detector saturation. Specifically, 2 ml of toluene was added to the smoke condensate (20 ml), agitated for 2 h on an orbital shaker (Z206A), and spun at 100 x g for 5 min at room temperature. The toluene was then separated prior to dissolving in 2 ml ethanol which was used for the GCMS analysis. It is worth noting that reproducibility was assured by using internal standards. Tridecane and 1-octadecene were used as internal standards for toluene extract while dibutyl phthalate was used for ethanol solution prior to GCMS analysis. The toluene and ethanol extract were labelled as ‘toluene’ and ‘toluene-ethanol’, respectively. The toluene and ethanol solutions were then analyzed using GCMS-QP2010 Ultra instrument (Shimadzu Corporation). Rtx-5MS capillary column (30 m in length, 0.25 µm in thickness, and a diameter of 0.25 mm) was used. The column oven temperature was held at 40˚C for 3 min then ramped at a rate of 5˚C per min to 300˚C where it was held constant for 15 min. Helium was used as the carrier gas at a column flow of 1.0 ml/min. The safety, hazard and toxicity evaluation of the toluene and ethanol extracts was performed using ToxNet and PubChem resources.

Cell culture and proliferation assay. A549 (gift from Professor Fathia Mami Chouaib, Gustave Roussy, Villejuif Cedex, France; RRID:CVCL_0023) and H460 cells (AddexBio C0016003 RRID:CVCL_0459) were grown in complete RPMI-1640 medium, (cat. no. 61870010; Gibco; Thermo Fisher Scientific, Inc.) supplemented with 10% heat-inactivated fetal bovine serum (cat. no. 10270-106; Gibco; Thermo Fisher Scientific, Inc.), 1% penicillin-streptomycin (cat. no. 15140-122; Gibco; Thermo Fisher Scientific, Inc.) and 1% sodium pyruvate (cat. no. 11360-039; Gibco; Thermo Fisher Scientific, Inc.). BEAS-2B cells (ECACC 95102433 RRID: CVCL_0168) were grown in BEGM media (cat. no. CC-3170; Lonza) on collagen (cat. no. A1048301; Gibco; Thermo Fisher Scientific, Inc.).-coated tissue culture dishes according to the manufacturer’s protocol. NK92 cells (gift from Professor Fathia Mami Chouaib) were cultured in complete RPMI-1640 medium supplemented with 200 IU of IL2 recombinant human protein (cat. no. PHC0021; Gibco; Thermo Fisher Scientific, Inc.). H460 and BEAS-2B were purchased from a commercial source that guarantees cell line authenticity. We tested and confirmed that all the cell lines were mycoplasma-free.

Trypan blue exclusion assay was used to determine the cell number and viability of A549, H460 and BEAS-2B cells exposed to various concentrations and durations of WPSC as described earlier in the text. Subsequently, 50,000 cells (A549 and H460) or 100,000 cells (BEAS-2B) were plated on 3.5-cm dishes for 24 h after which the cells were treated with 1 ml of complete media containing WPSC at the indicated concentrations. Plates were counted, every day for 7 days and the number of live and dead cells was determined.

RNA extraction/cDNA and qPCR. RNA was purified using Easy Blue (cat. no. 17061; Intrion Biotechnology, Inc.) according to the manufacturer’s protocol. RNA quality and quantity were assessed by nanodrop and 1% agarose gel electrophoresis. cDNA synthesis was performed using high capacity cDNA reverse transcription kit (cat. no. 4374966; Thermo Fisher Scientific Inc.). qPCR was performed using SYBR-Green (cat. no. 4309155; Thermo Fisher Scientific Inc.). Ab 7500 FAST Real-Time PCR system. Analysis was performed using the ΔΔCq method (15). Forward (F) and reverse (R) primers used in this study were: GAPDH forward, (5'-GCCACATCGTCGACACAC-3') and GAPDH reverse, (5'-CCAGAGTFAAAGACGAC-3'); CCL2 forward, (5'-CAGCCAGATGCAATGCAAATG-3') and CCL2 reverse, (5'-TGGAAATCTGTAACCACCTTCT-3'); CASP1 forward, (5'-TTTCGCCAAAGTGGTATTTTCA-3') and CASP1 reverse, (5'-GGCATCTGCGCTCTACCATC-3'); IL1B forward, (5'-TTGCACATGGGATAACGAGG-3') and IL1B reverse,
with 300,000 NK92 cells in the presence of IL-2 for 30 min in
immunological synapse 100,000 lung cells were incubated
For visualizing
Immunological synapse formation assay. LSM 800).
Fisher Scientific, Inc.) and visualized on Zeiss LSM 800 with
antibody staining. Cells were then mounted on glass slides
antibodies as described below with 3x5 min washes after each
The cells were then stained with a primary and secondary
15 min at room temperature. Prior to staining, the cells were
was performed at 0.7 V/cm and 300 mA for 35 min. The slides
were left for 30 min for DNA unwinding and electrophoresis
horizontal electrophoresis tank and soaked in cold alkaline
holes in the second layer, and was allowed to solidify. After
gelling, the slides were immersed in chilled lysing solution
containing 2.5 M NaCl, 100 mM EDTA, 0.1% TX-100, 0.5% NaDOC, 0.1% SDS,
gelling, the slides were immersed in chilled lysing solution
containing 2.5 M NaCl, 100 mM EDTA, 10 mM Tris base
pluggin on ImageJ software. The tail length was considered
per slide and the data were analyzed using OpenComet (17)
(0.2 mg/100 ml) and were visualized at x20 magnification on
neutralized using 0.4 M Tris (pH 7.4). To capture images,
were then removed from the electrophoresis buffer and were
were loaded on 10 or 12% SDS-PAGE, and transferred onto a nitrocellulose membrane (cat. no. GE10600004; Sigma-Aldrich; Merck KGaA). Proteins were quantified following brief sonication by Pierce BCA protein assay kit (cat. no. 23225; Thermo Fisher Scientific Inc.). Then, 8-12 mg of proteins were loaded on 10 or 12% SDS-PAGE, and transferred onto a nitrocellulose membrane (cat. no. GE10600004; Sigma-Aldrich; Merck KGaA) at 80 V for 3 h for low molecular weight proteins or 80 V for 8 h for high molecular weight proteins. After incubation with 5% BSA in TBST (10 mM Tris, pH 8.0, 150 mM NaCl, 0.5% Tween-20) for 60 min at room temperature, the membrane was washed once with TBST and incubated with the listed antibodies overnight at 4˚C according to their data sheets.
β-galactosidase senescence assay. Cells were stained for
β-galactosidase using Senescence β-galactosidase Staining Kit (cat. no. 9860S; Cell Signaling Technology) according to manufacturer's protocol.
NK92-mediated cytotoxicity assay. Cytotoxicity assay was performed as previously described (18) with the following modifications: IL-2-activated NK92 effector cells were incubated with target A549 or H460 cells with the E/T ratios, 1:10, and 1:20 in a 5% CO₂ incubator at 37˚C in a 96-well plate in triplicates. After 6 h of co-culture, NK activity was measured using the LDH Cytotoxicity Assay Kit (cat. no. 88954; Pierce™) according to the manufacturer's protocol.
Microarray analysis. Total RNA concentrations were determined using the Qubit® RNA HS Assay Kit (cat. no. Q32852; Invitrogen; Thermo Fisher Scientific, Inc.) using Qubit® 3.0 (cat. no. Q33216; Invitrogen; Thermo Fisher Scientific, Inc.) and RNA quality was assessed by running a 2% agarose gel and checking the 18S and 28S rRNA bands.

(5'-TTTTTGGCTGTGAGTCCCGGAG-3'); CD44 forward, (5'-TGCCGCTTTGGAGTCTATT-3') and CD44 reverse, (5'-CCGATGCTGGTTGGTTTCC-3'); SERPINE2 forward, (5'-TGGTGAAGAGGATACCGGCCGTA-3') and SERPINE2 reverse, (5'-TGGTGAAGAGGATACCGGCCGTA-3'); SNAI2 forward, (5'-CCGCAAGGCACATCTGGTTA-3'); ACTA2 forward, (5'-CTATGAGGCTATGCTTGGC-3') and ACTA2 reverse, (5'-GGTGGAGCAGTGTAAAGGAAGA-3'); IL6 forward, (5'-ACTACCTTCTTCAAGAACATTTG-3') and IL6 reverse, (5'-CCATCTTTTGGAGGTTCCAGTGG-3').

Alkaline comet assay. Cells were harvested and prepared as described in Olive and Banáth (16). Briefly, the treated cells were centrifuged at 100 x g at room temperature for 5 min. Cell pellet was re-suspended in 1X PBS and approximately 5,000 cells per slide were considered for the assay. Cells were mixed with 0.75% low-melting agarose (LMA) at 37˚C, and the mixture was loaded onto glass slides pre-coated with 1.5% normal melting agarose. Coverslips were then placed gently on the slides to allow even spreading of the gel and then placed at 4˚C for gelling. A third layer of the 0.75% LMA was added onto the slide to fill any residual holes in the second layer, and was allowed to solidify. After gelling, the slides were immersed in chilled lysing solution containing 2.5 M NaCl, 100 mM EDTA, 10 mM Tris base (pH 10) with 1% Triton X-100 and 10% DMSO and were kept overnight at 4˚C. After lysis, the slides were placed in a horizontal electrophoresis tank and soaked in cold alkaline electrophoresis buffer (1 mM EDTA, 300 mM NaOH). Slides were left for 30 min for DNA unwinding and electrophoresis was performed at 0.7 V/cm and 300 mA for 35 min. The slides were then removed from the electrophoresis buffer and were neutralized using 0.4 M Tris (pH 7.4). To capture images, the slides were stained with 100 ml of ethidium bromide (0.2 mg/100 ml) and were visualized at x20 magnification on Zeiss LSM 800 with Airyscan. In total, 50 images were taken per slide and the data were analyzed using OpenComet (17) plugin on ImageJ software. The tail length was considered as a suitable parameter for measuring the extent of DNA damage.

Immunofluorescence. Cells were fixed in 4% paraformaldehyde (cat. no. 28906; Thermo Fisher Scientific Inc.) in 1X PBS at 10 min for room temperature. Cells were then washed with 1X PBS and permeabilized with 0.1% TX-100 in PBS for 15 min at room temperature. Prior to staining, the cells were blocked in 2% BSA in 1X PBS for 1 h at room temperature. The cells were then stained with a primary and secondary antibodies as described below with 3x5 min washes after each antibody staining. Cells were then mounted on glass slides using Prolong gold antifade reagent (cat. no. P36930; Thermo Fisher Scientific, Inc.) and visualized on Zeiss LSM 800 with Airyscan. The comet assays were visualized using a 20x plan apochromat objective, with Zeiss Axio cam 305 mono (Zeiss LSM 800).

Immunological synapse formation assay. For visualizing immunological synapse 100,000 lung cells were incubated with 300,000 NK92 cells in the presence of IL-2 for 30 min in a 37˚C incubator. Cells were subsequently fixed and stained for the markers described in the text.

Antibodies used in this study. Mouse monoclonal anti-human Phospho-histone H2AX (cat. no. 05-636; 1:1,000 dilution; Merck Millipore), rabbit anti-human histone H2AX (cat. no. P16104; 1:2,000 dilution; Ray Biotech), rabbit anti-human p21 (cat. no. 2147; 1:1,000 dilution; Cell Signaling Technology), rabbit anti-human 53BP1 (cat. no. 4937; 1:1,000 dilution; Cell Signaling Technology), mouse anti-human p53 (cat. no. 48818; 1:1,000 dilution; Cell Signaling Technology), mouse monoclonal anti-gizzard β-actin (cat. no. sc-47778; 1:5,000 dilution; Santa Cruz Biotechnology), phallolidin (cat. no. A12379; 1:1,000 dilution; Thermo Fisher Scientific Inc.), DAPI (cat. no. D1306; 1:36,000 dilution; Thermo Fisher Scientific Inc.). Secondary antibodies used are goat anti-mouse Alexa Fluor 568 (A11004; 1:1,000 dilution; Thermo Fisher Scientific, Inc.), goat anti-rabbit Alexa Fluor 488 (cat. no. A11034; 1:1,000 dilution; Thermo Fisher Scientific Inc.), goat anti-mouse Alexa Fluor 488 (cat. no. A11001; 1:1,000 dilution; Thermo Fisher Scientific Inc.), goat anti-rabbit Alexa Fluor 568 (cat. no. A11011; 1:1,000 dilution; Thermo Fisher Scientific Inc.).

Immunoblotting. Cells grown in 3.5-cm plates were washed once with 1X ice-cold PBS and lysed in 100 ml of RIPA (150 mM NaCl, 0.1% TX-100, 0.5% NaDOC, 0.1% SDS, 50 mM Tris-CL pH 8.0) with protease inhibitor cocktail (cat. no. P2714; Sigma-Aldrich; Merck KGaA). Proteins were quantified following brief sonication by Pierce BCA protein assay kit (cat. no. 23225; Thermo Fisher Scientific Inc.). Then, 8-12 mg of proteins were loaded on 10 or 12% SDS-PAGE, and transferred onto a nitrocellulose membrane (cat. no. GE10600004; Sigma-Aldrich; Merck KGaA) at 80 V for 3 h for low molecular weight proteins or 80 V for 8 h for high molecular weight proteins. After incubation with 5% BSA in TBST (10 mM Tris, pH 8.0, 150 mM NaCl, 0.5% Tween-20) for 60 min at room temperature, the membrane was washed once with TBST and incubated with the listed antibodies overnight at 4˚C according to their data sheets.

β-galactosidase senescence assay. Cells were stained for β-galactosidase using Senescence β-galactosidase Staining Kit (cat. no. 9860S; Cell Signaling Technology) according to manufacturer's protocol.
Subsequently, 100 ng of good quality RNA was prepared for transcriptome profiling using the Human Clariom™ D Assay (cat. no. 902922; Applied Biosystems). All steps were performed following the manufacturer’s protocol, where in brief, the GeneChip™ WT PLUS Reagent Kit (cat. no. 902281; Applied Biosystems) was first used to generate amplified and biotinylated sense-strand DNA that was then hybridized on the Human Clariom™ D Arrays (Applied Biosystems™). These arrays include over 6,765,500 probes that can detect genes, exons, and alternative splicing events giving rise to coding and long non-coding RNA isoforms. The arrays were washed and stained on the GeneChip Fluidics Station 450 (cat. no. 00-0079; Applied Biosystems) and scanned with the GeneChip™ Scanner 3,000 7G (cat. no. 00-0213; Applied Biosystems). The scanned images and DAT files were inspected for the absence of bubbles and proper scanner alignment, respectively, while the generated raw files (CELL) were imported to the Transcriptome Analysis Console (TAC) 4.0 software (Applied Biosystems) for data analysis. Expression (Gene + Exon) analysis was carried out on TAC by applying the Gene + Exon-Signal Space Transformation-Robust Multi-Array Average (SST-RMA) algorithm. Default settings were used for determining relative gene expression levels of each transcript, which included applying the ebayes (Empirical Bayes Statistics for Differential Expression) ANOVA method for statistical testing, as well as setting the threshold for gene expression fold change (FC) between the treated and untreated samples at ≤ -2 or ≥2. Only coding and multiple complex loci with P-value <0.05 were considered for further analysis. A heatmap was generated by hierarchical clustering of common deregulated genes using complete linkage with Euclidean distance on Heatmapper (19). Venn diagram was plotted using Venny (20).

### Table I. Major chemical compounds identified in the toluene and ethanol extracts.

<table>
<thead>
<tr>
<th>Toluene</th>
<th>Ethanol</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benzaldehyde</td>
<td>Hexadecane, 2,6,10,14-tetramethyl- (CAS)</td>
</tr>
<tr>
<td>Benzaldehyde, 3-ethoxy-4-hydroxy- (CAS)</td>
<td>Hexadecanoic acid, 1-(hydroxymethyl)1,2-ethanediyl ester</td>
</tr>
<tr>
<td>Benzene, 1,3-dimethyl-</td>
<td>4H-Pyrane-4-one, 2,3-dihydro-3,5-dihydroxy-6-methyl-</td>
</tr>
<tr>
<td>Benzyl alcohol</td>
<td>5-Hydroxymethylfurural</td>
</tr>
<tr>
<td>Benzenepropanoic acid, 3,5-bis(1,1-dimethyl)ethyl-4-hydroxy-, octadecyl ester</td>
<td>1H-Indene, 1-hexadecyl-2,3-dihydro-</td>
</tr>
<tr>
<td>Cyclopetanone, 1-ethyl-2-methyl-, cis-</td>
<td>Nonyl tetracetyl ether</td>
</tr>
<tr>
<td>Cyclohexane, ethyl-</td>
<td>Nicotine</td>
</tr>
<tr>
<td>Decane, 3,8-trimethyl-</td>
<td>octadecanoic acid, 3-oxo-, ethyl ester</td>
</tr>
<tr>
<td>DECANE, 3,3,7-TRIMETHYL-</td>
<td>Octadecanoic acid, 2,3-dihydroxypropyl ester</td>
</tr>
<tr>
<td>Dihydro methyl jasmonate</td>
<td>Phenol, 2,4-bis(1,1-dimethyl)ethyl- , phosphite (3:1)</td>
</tr>
<tr>
<td>2,3-Dihydroxypropyl icosanoate, 2TMS derivative</td>
<td>Pyrrolidine-D4</td>
</tr>
<tr>
<td>Docosane</td>
<td>Ethanol, 2-[(triethylsilyl)oxy]-</td>
</tr>
<tr>
<td>Dodecane, 4,6-dimethyl-</td>
<td>1,2,3-Propanetriol, 1-acetate</td>
</tr>
<tr>
<td>Ethylbenzene</td>
<td>Pyridine, 3-(1-methyl-2-pyrrolidinyl)-(S)-(CAS)</td>
</tr>
<tr>
<td>Hexadecane, 2,6,10,14-tetramethyl- (CAS)</td>
<td>Propanal, 3-ethoxy- (CAS)</td>
</tr>
<tr>
<td>1-Hexadecanol</td>
<td>Trans-Anethole</td>
</tr>
<tr>
<td>Hexane, 3,3-dimethyl- (CAS)</td>
<td>Tris(2,4-di-tert-butylphenyl) phosphate</td>
</tr>
<tr>
<td></td>
<td>Trans-4-hydroxymethyl-2-methyl-1,3-dioxolane</td>
</tr>
</tbody>
</table>

Statistical analysis. Statistical analyses were carried out using GraphPad Software version 6 (GraphPad Software, Inc.). All data are expressed as means ± SEM. Significant differences were found using two-way analysis of variance (ANOVA) followed by Bonferroni post hoc test.

**Results**

Toxic compounds identified in WPSC. We first attempted to identify chemical compounds in WPSC by analyzing the toluene extract and the ethanol solution using GCMS. Many peaks were identified in the chromatograms including some that are present at trace levels. Major peaks are shown in Fig. 1. An internal standard was used to evaluate the reproducibility of the analysis and to allow the calculation of relative concentrations. Table I summarizes the major chemical compounds identified in both the toluene and ethanol solutions. Most of these compounds are known to cause serious health problems. The safety, hazard and toxicity evaluation of the toluene and ethanol extracts were performed using ToxNet and PubChem resources. Several compounds were identified as carcinogens or may cause various toxicities and health hazards. Several harmful compounds were identified and may be associated with the burning of coal as reported in a previous study (14). For example, ethyl benzene, a component previously identified in coal smoke was also...
Figure 1. GCMS chromatograms for WPSC. For this step, 2 ml of toluene was added to the smoke condensate (20 ml), agitated for 2 h at room temperature on an orbital shaker (Z206A), and spun at 100 x g for 5 min at room temperature. The toluene was then separated prior to dissolving in 2 ml ethanol which was used for the GCMS analysis. Toluene extraction is shown in blue; ethanol extraction is shown in orange.

Figure 2. WPSC effects on lung cell proliferation. Growth curves for A549, H460 and BEAS-2B cell lines grown in the presence of 0.5% WPSC are shown. The cell number was determined by counting in a hemocytometer after detachment of cells with trypsin every day up to 8 days. The number of live cells is shown in A, C and E. The number of dead cells is shown in B, D and F. Results represent means of three independent experiments, were run in duplicates and data represent mean ± standard error of mean. *P≤0.05, **P≤0.01 and ***P≤0.001.

A549 cell growth

A549 cell death

H460 cell growth

H460 cell death

BEAS-2B cell growth

BEAS-2B cell death
found in WPSC. It is classified by the International Agency for Research on Cancer (IARC) as a possible human carcinogen and may cause kidney failure as a result of prolonged exposure at low concentrations (14). Indene, another compound found in WPSC, is a polycyclic aromatic hydrocarbon which was identified in coal smoke (14). Furthermore, several identified compounds are known to cause irritation to the skin, eyes and respiratory tract, including 3-ethoxy-4-hydroxy-benzaldehyde, 1,3-dimethyl-benzene, benzyl alcohol, docosane, ethylbenzene, 3-ethoxypropionaldehyde, and 2,3-dihydro-3,5-dihydroxy-6-methyl-4H-pyran-4-one. Many of the identified compounds are known to interfere with and alter the functions of the central nervous system including 1,3-dimethyl benzene, benzyl alcohol, ethyl cyclohexane, ethyl benzene and nicotine. In addition, several compounds are known to induce DNA damage including benzaldehyde, 1,3-dimethyl Benzene, and 5-hydroxymethyl furfural (23).

Effect of WPSC on lung cancer cell proliferation. We first tested the cytotoxic effect of WPSC on two non-small cell lung cancer cells, A549, and H460 as well as on a normal bronchial epithelial cell line, BEAS-2B. Both cell number and viability of tested cells were measured in the presence of various concentrations of WPSC condensate (0.5, 1 and 2%) for 8 consecutive days with re-administration of fresh WPSC at day 4. Additionally, WPSC treatment resulted in a dose- and time-dependent decrease in cancer cell proliferation (Fig. 2A-D). Hemocytometer images are shown in Fig. S1. At higher doses of 1 and 2% an increase in cell death was observed. By contrast, the lowest concentration of 0.5% resulted in a significant increase in the cell death of BEAS-2B normal lung cells following 2 days of treatment (Fig. 2E and F); consequently, these cells were not analyzed further. Subsequent experiments on lung cancer cells were performed using 0.5% WPSC, considering the absence of cell death.

WPSC increases DNA damage and cellular senescence in lung cancer cells. DNA damage often arises as a result of normal cellular processes, as a by-product of the cell's own metabolic activity. It can also be induced as a result of environmental exposure to chemical agents. We therefore investigated whether WPSC induced DNA damage and used the alkaline comet assay to detect and quantify the degree of gross DNA damage. As shown in Fig. S2A and B, WPSC induced an increase in tail length in the cancer cell lines A549 and H460 when compared to untreated cells, indicating an increase in the frequency of DNA breaks. Comet assay reflects the physical status of genomic DNA while 53BP1/γH2AX staining represents processes related to the biological DNA damage response (DDR) (24). We investigated the effect of WPSC on the DDR using two markers: Phosphorylated histone H2AX at Ser139 (γH2AX) and 53BP1. Foci formation indicates the aggregation of these proteins induced by DNA damage. Consistent with the comet tail formation, in response to WPSC treatment, an increase
In the nuclear foci for both γH2AX and 53BP1 in A549 and H460 cells was observed (Fig. S2C-E).

Since waterpipe smoke condensate resulted in decreased cell growth, we tested whether the lung cancer cells were undergoing apoptosis or cell cycle arrest and senescence. Consistent with cell cycle arrest and induction of senescence, we found that WPSC induced an increase in p21 protein expression as revealed by western blot analysis (Fig. 3A).

Using confocal microscopy analysis, we demonstrated an accumulation of both p21 and p53 proteins in the nuclei of A549 and H460 cells (Fig. 3B).
both cancer cells (Fig. 3B). This was also supported by an increase in β-galactosidase activity in these cells indicating the induction of cellular senescence (Fig. 3C and D).

Transcriptomic changes associated with WPSC treatment. To determine whether WPSC treatment impacts the molecular pathways associated with EMT, stemness and immune evasion and to identify novel gene expression patterns in response to WPSC treatment, a microarray analysis was performed. RNA derived from biological duplicates of A549 and H460 cell lines 8 days post-treatment with WPSC, as well as their untreated counterparts was analyzed. Considering only coding and multiple complex loci with fold change of ≤ -2 and ≥2 and P-value of <0.05, volcano plots were generated. A greater number of differentially expressed gene transcripts in A549 (1452 upregulated and 1272 downregulated) in comparison with H460 (452 upregulated and 604 downregulated) (Fig. 4A and B) was observed. Among these differentially expressed transcripts, 184 were common to both A549 and H460 (Fig. 4C and D). Of these 70 transcripts showed opposite patterns of deregulation: upregulated in one cell line and downregulated in the other or vice versa. Additionally, 50 transcripts were upregulated, and 64 transcripts were downregulated in the two cell lines, respectively (Fig 4D). Pathway enrichment analysis revealed that the commonly deregulated genes clustered in pathways involved EMT, the cell cycle, apoptosis, DNA repair and the inflammatory response, among others (Fig. 4E).

WPSC activates the inflammatory response in lung cancer cells. Waterpipe smoking has been shown to trigger the inflammatory response in humans as measured in the blood of waterpipe smokers (25). Using microarray analysis, gene expression in the inflammatory pathway (Fig. 5A and B)
was detected. IL-1β is a key mediator of inflammation and promotes invasiveness (26), whereas the chemokine CCL2 regulates inflammatory responses and also plays a role in tumor progression and metastasis (27). In addition, IL-6 is a pro-inflammatory cytokine with key roles in EMT (28).

Fig. 5A and B shows a significant increase in expression levels of these genes in response to WPSC. We also investigated the expression levels of caspase-1, the cysteine protease that converts the inactive proform of IL-1β to the active inflammatory cytokine (29). The results showed that the expression level was slightly increased in the two cells (Fig. 5A and B), consistent with the increase in IL-1β.

WPSC induces EMT and stemness properties. EMT and stem cell-like traits are intertwined processes that dictate tumor aggressiveness. It is well established that EMT generates cells with stem cell properties (30). Using microarray analysis, several genes involved in EMT were found to be enriched including IL-6, IL-15, GLIPR1, CD44 and COL5A. Using qPCR analysis, we demonstrated that both EMT and stemness genes are regulated (Fig. 5C and D), an increase in the cancer stem cell marker CD44 was observed. CD44 is also known to promote EMT (31). Furthermore, an increase in SNAI2, an EMT key transcription factor (32), SERPINE2, a protease that promotes ECM deposition and invasion (33) and ACTA2 coding for alpha smooth muscle actin that promotes motility and invasion (34) were also increased in A549 cells. Cell morphological changes in response to WPSC were observed. WPSC treatment converted the cells from a ‘cuboidal’ epithelial structure into an elongated mesenchymal shape (Fig. 5E).

WPSC effect on synapse formation and NK cell-mediated killing. We examined whether WPSC treatment of cancer cells interferes with tumor cell recognition and killing. As is evident in Fig. 6A, the immunological synapse between NK cells and WPSC-treated A549 and H460 cancer cell was not affected. Nevertheless, as shown in Fig. 6B, WPSC induced a significant decrease in the killing of A549 cells but had no significant effect on the H460 cells.

Discussion

In the present study, we investigated the effects of WPSC, using a new method based on a smoking topography stimulating machine on non-small-cell lung cancer cells. We demonstrated that WPSC treatment resulted in a dose-dependent decrease in cell proliferation with cell death at higher concentrations. Both cellular senescence and apoptosis were induced in cancer cells at the lower WPSC concentration of 0.5%. However, such treatment resulted in cell death of normal lung cells. As such, we are in the process of addressing the effects of short-term duration treatment with various concentrations of WPSC on normal lung cells. Furthermore, we did not explore doses lower than 0.5% as this did not inhibit growth in cancer cells. However, the molecular changes described in this study may be induced at lower doses. Adaptive cellular responses are activated upon microenvironmental stress, such as WPS
exposure, and are critical in the prevention of tissue damage and transformation. The net effect of these responses depends on the duration, nature, and intensity of the stress. Indeed, at shorter duration treatment of 72 h we did not observe any cell death (data not shown), which is consistent with previous findings showing that only senescence was generated at this time point in A549 cells (10).

Prolonged exposure to environmental stresses may become deleterious in promoting cell death, excessive inflammation, and tissue remodeling. Indeed, we observed that WPSC treatment resulted in increased DNA damage and a subsequent increase in γH2AX and 53BP1, which is consistent with the recent finding of Yoshida et al., indicating that tobacco smoking resulted in the increase in mutational burden of lung cells (35), and with a known increase in the expression of both γH2Ax and 53BP1 in several cancers including breast, bladder, lung, head and neck (36). By contrast, cigarette smoke has been shown to induce a decrease in nucleotide excision repair mechanisms, and does induce DNA damage as measured by COMET assay (37,38), which raises the interesting possibility that other factors present in WPS may induce a toxic response independent of DNA damage in these cells.

We then analyzed the effects of 8-day treatment of WPSC on the transcriptomic profile of A549 and H460 cells. The results showed an increase in the expression of genes involved in inflammation, EMT and in the generation of cancer stem cells. It is well known that inflammation plays an important role in the tumor-specific microenvironment, which is thought to facilitate all phases of tumorigenesis, from initiation to metastasis (39). Inflammatory mediators also exhibit the potential to induce the acquisition of CSCs properties (40). In the course of this study, we demonstrated an increase in IL-1β, IL-6, CCL2 and caspase-1 levels, known to be associated with the induction of EMT (41-43). IL-6 and IL-1β are also found to be senescence-associated secreted factors (44). WPS in humans has a significant association with systemic inflammation (25,45,46). Our results are consistent with this role for WPSC and further highlight the importance of inhibiting inflammatory processes in WP smokers as a means for more effective cancer treatment.

Of the EMT-related genes, we observed an increase in the expression of SNAI2, SERPINE2, ACTA2 as well as CD44, also is a regulator of cancer stemness. WPS was shown to induce EMT in breast cancer cell lines through FAK and Erk1/Erk2 expression (47) and resulted in an increase in invasion and migration properties of breast cancer cells (48). Our results therefore suggest that WPSC exposure to lung cancer cells may change their physiology to promote EMT processes and acquire CSCs properties.

NK cell-mediated immune surveillance strengthens host defense against certain microbial agents and cells undergoing malignant transformation. We investigated the effects of this WPS treatment on the ability of NK cells to recognize and kill the lung cancer cell targets. To date, no data are available as to the effect of WPS on NK cell function. Nevertheless, in mice, chronic exposure to WPS has been reported to promote immune suppression (49). In addition, in cigarette smokers lower populations of NK cells are found (50), and patients with lung cancer had a marked decrease in NK activity compared to non-smokers (51). However, other reports indicate that cigarette smoking was associated with an increase in NK cell tumoricidal activity without any alteration in the absolute number of NK cells in blood (52). Furthermore, NK cells from cigarette-exposed mice produced more IFN-γ following stimulation with IL-12, IL-18, or both (53). Under our experimental conditions, the results showed that A549 cell killing was reduced as compared to non-treated cells, whereas H460 cell survival was not affected. However, the capacity of the NK cells to form synapses with the WPSC-treated cancer cells was not affected in the two cell lines. This is most likely due to genetic variations in the two cell lines, engaging communication networks that could also include the release of inflammatory mediators that signal to the immune system. This is interesting as one possibility is that WPSC could induce several pathways that affect susceptibility to lysis. Nevertheless, we did not observe a correlation between the EMT/CSC induction in A549 cells and a decrease in their NK-mediated cell lysis.

As a limitation of the study, we did not explore the molecular mechanisms of cell apoptosis, as we found WPSC affecting other hallmark properties of cancer including EMT-, stemness- and genomic instability (DNA damage)-related changes. These characteristics are not unique to waterpipe smoke as cigarette smoke has also been shown to induce EMT cell properties (54) in addition to apoptosis (55).

WPSC can induce apoptosis (12). At the same time, WPSC also drives hallmark cancer properties (senescence and EMT, stemness and genomic instability) that promote survival of these cells. In addition, these cells also acquire properties to escape immune surveillance, which complicates the situation further. These characteristics are not unique to WPSC as cigarette smoke has also been shown to impair NK-mediated immune surveillance (54) and to induce EMT cell properties (54) as well as apoptosis (55). We are currently culturing the cells in WPSC media for longer durations (up to 4-5 months) and we see that these tumor cells keep dividing (unpublished data). As DNA damage contributes to increased genomic instability and mutational burden, WPSC can also contribute to genetic heterogeneity of tumors.

Our results indicate that WPSC is a contributing factor in the pathogenesis of lung cancer, through impairing cell growth, inducing inflammation and DNA damage. If we are to assume that a cancer patient continues to smoke, WPSC may serve as fuel to the cancer cells and may contribute to metastases. In fact, continued smoking is considered a strong adverse predictor of survival and increases the risk of a second lung cancer compared to those patients who stopped smoking (56). Thus, therapy modalities can be more effective by eliminating smoke exposure to cancer patients and by targeting the inflammatory mechanisms in order to control the emergence of aggressive cancer clones with EMT/CSCs features.

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Availability of data and materials

The datasets used and/or analyzed during the current study are available with the corresponding author on reasonable request.

Authors' contributions

SC conceived, significantly guided in the design, analysis, and interpretation of the findings of this study. RFZ, PP, NZ, AR and ST conceived and designed experiments, acquired and interpreted data. RAK, FA performed the microarray-related experiments and analysis with contributions from GHV. YES performed the WPSC purification and analysis. HN conducted the confocal imaging. All authors discussed the results and contributed to the final manuscript.

Ethical approval and consent to participate

Not applicable.

Patient consent for publication

Not applicable.

Competing interests

All authors declare that they have no competing interests.

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