The role of synthetic coolants, WS-3 and WS-23, in modulating E-cigarette-induced reactive oxygen species (ROS) in lung epithelial cells

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1. Introduction

During the past few years, adolescent use of e-cigs or various electronic nicotine delivery systems (ENDS) has significantly increased, thus leading to an increase in the prevalence of E-cigarette (E-cig) or Vaping Associated Lung injury (EVALI) across the United States [1,2]. As of February 18, 2020, a total of 2807 EVALI-related hospitalizations or deaths were reported to the Centers for Disease Control (CDC) from all 50 states [1,2]. Consequently, the Food & Drug administration (FDA) implemented an enforcement policy to remove all flavored cartridge or/pod-based e-cigs except tobacco and menthol-flavored pods from the market [3].

Following the FDA’s 2020 flavor-enforcement policy, menthol-flavored e-cig sales had significantly increased in the US; specifically, there was a 54.5 % increase in the market share of menthol-flavored e-cigs over four weeks and an 82.8 % increase over eight weeks following the FDA’s ruling [4]. Menthol induces a perception of a cool sensation by activating the cold receptor found in the oral cavity [5,6]. Menthol reduces the bitterness associated with inhaled nicotine and increases its smoothness upon inhalation, thus increasing e-cigarette appeal [7].

Other than menthol, several synthetic cooling agents have been added to e-cigarette/e-liquid formulations as a replacement for menthol due to...
these synthetic agents giving similar cooling sensations upon aerosol inhalation as menthol [8-11]. Methyl diisopropyl propionamide (WS-23) and N-Ethyl-2-isopropyl-5-methylcyclohexanecarboxamide (WS-3) are examples of these added cooling agents [5,9].

A recent report has showed that e-cigarette flavors with "ice," "chilled," "Cooled," and "Polar" in their name, and other flavors consisting of flavor combinations with fruity and drink flavors like "melon-ice," "blueberry-ice," and "iced-pink punch" contained WS-23 and WS-3 in their formulations [12]. The significant increase in the marketing of “iced/cooled” flavored e-cigs in the U.S. had occurred right around the time when sales of disposable e-cigs surged following the FDA’s implementation of its March 2020 e-cig flavor enforcement policy [12]. One study had found, via GC-MS and LC-HRMS/MS, that both WS-3 and WS-23 were major components found in nicotine-containing vaping fluids provide by patients apart of the 197 reported-cases of EVALI in New York State to the Wadsworth Center at the New York State Department of Heath from August 2019 through June 2021 [13]. Additionally, one study [9] found that WS-23 was present in e-cigs marketed in the US at levels that may potentially result in exceeding the Margin of Exposure (MOE), a risk assessment parameter for toxic compounds used by World Health Organization (WHO) [9], Jabba et al. [14] results suggest that those who use e-liquids comprised of WS-3 or WS-23 are potentially at risk for long-term pulmonary health issues [9].

Aerosols generated by e-cigs or other ENDS modalities have been found to contain dangerous chemicals, including formaldehyde and acetaldehyde, which are known to cause lung cancer and cardiovascular disease [15]. Also, consistently, it has been found that dysregulated inflammatory cytokine output is an effect of chronic e-cig exposure in both in vivo and in vitro models [16-18]. Moreover, previous studies have shown that aerosols generated by flavored e-cigs produce significant levels of acellular reactive oxygen species (ROS) and induce cellular ROS in small airway epithelial cells (SAEC) [19-21]. ROS, either exogenous or when produced in excess endogenously, can lead to a redox imbalance in the lungs [22]. One study found tobacco smoke to contain a significant amount of free radicals, ~1 x 10^15 radicals per puff [14,23, 24]. ROS in smoke generated from conventional cigarettes, when inhaled, will react with antioxidants in the epithelial lining fluid (ELF) covering airway epithelial cells [23]. Moreover, ROS in tobacco smoke, after reaching the ELF of airways, can lead to the destruction of endogenous antioxidants, thus significantly reducing cellular antioxidant capacity [24]. Oxidative stress induced by this redox imbalance has been implicated in the pathology of many types of lung diseases, such as acute respiratory distress syndrome (ARDS), asthma, and chronic obstructive pulmonary disease (COPD) [22]. Similarly, studies have shown that exposure of e-liquids, e-liquid solvents, and e-cig aerosols have led to significant increases in the levels of cellular ROS produced by cultured pulmonary airway cells [25,26].

Studies so far have shown that exposure to e-cig aerosols induces oxidative and carbonyl stress in the lungs [17,21,27]. Regarding ROS-related e-cig studies, studies have shown that total acellular ROS levels in e-cig aerosols are dependent on brand, flavor, operational voltage, and puffing protocol, but no studies so far have sought to investigate the role of synthetic coolants have in modifying total acellular ROS in aerosols generated from e-liquids and cellular ROS levels from BEAS-2B cells exposed to those e-liquids. In this study, we seek to understand the role of WS-23 and WS-3 in modifying acellular and cellular ROS levels due to exposure to e-liquids.

2. Materials and methods

2.1. Procurement of e-liquid constituents and composition of e-liquid solutions

Propylene Glycol (PG), Vegetable Glycerin (VG), WS-23 solution (30% suspended in PG), and Koolada (10% WS-3 in PG) were purchased online from Flavor Jungle. 100 mg/mL nicotine salt solution (50:50 PG-to-VG ratio) was purchased online from PERFECTVAPE. E-liquid solutions comprising of PG, VG, salt nicotine, Koolada, and WS-23 were made. For our acellular ROS assays, the following e-liquids were made (Table 1). Six different e-liquid formulations were used in this study; all of them containing a mixture of PG and VG at a 50:50 vol percentage ratio, and some of them differing in their volume concentrations of nicotine (0% or 5%) and Flavor Jungle synthetic coolant-containing solution (0% or 3%).

2.2. Generation of aerosols, fluorescence spectroscopy, and acellular ROS quantification

Each e-liquid solution was added to a new, empty refillable pod (OVNStech, Shenzhen, GD, China) (Mo: W01 Pods) and aerosolized using a pod-based device. Specifically, each pod-based device was attached to a Buixo Individual Cigarette Puff Generator (Data Sciences International, St. Paul, MN, USA) (Cat#601–2055–001), and subsequently, its component e-liquid was aerosolized and “bubbled” through 10 mL of freshly made fluorogenic dye within a 50 mL conical tube (Fig. 1). Cell permeant 2′,7′-dichlorodihydrofluorescein diacetate (H2DCFDA) (EMD Biosciences, San Diego, CA, USA) (Cat# 287810) dissolved in 0.01 N NaOH, phosphate buffer, PO4, and horseradish peroxidase (Thermo Fisher Scientific, Waltham, MA, USA (Cat# 31491)) were used to make the fluorogenic dye. The aerosols generated from each e-liquid solution were individually bubbled through 10 mL of H2DCFDA solution at 1.5 L/min. A schematic of the e-cig aerosolization procedure is shown in Fig. 1. Each pod containing a respective e-liquid solution had undergone three separate puffing regimens to create three separate samples of bubbled dye solution. The same puffing regimen was used for “bubbling” filtered air through fluorogenic dye for a negative control. For our positive control, the smoke generated from a research cigarette (Kentucky Tobacco Research & Development Center in the University of Kentucky, Lexington, KY, USA) (Mo: 3R4F) was bubbled through the fluorogenic dye. After “bubbling,” each resulting fluorogenic dye sample was placed in a 37 °C degree water bath (VWR 1228 Digital Water Bath) for fifteen minutes; subsequently, the solution was analyzed via fluorescence spectroscopy using a spectrofluorometer (Turner Quantech fluorometer, Mo. FM109535) in fluorescence intensity units (FIU). Readings on the spectrofluorometer were measured as H2O2 equivalents using a standard curve generated using the 0–50 μM H2O2 standards made.

2.3. Cells culture conditions and treatments

BEAS-2B cell lines (ATCC, Manassas, VA) were used in this study for subsequent oxidative stress detection assays. Cells were maintained in

Table 1 Composition of E-liquids Analyzed.

| Composition of E-liquid solution | PG:VG Ratio by volume | Nicotine concentration (% by volume) | Cooling solution added | Cooling solution concentration (% by volume) |
|----------------------------------|-----------------------|------------------------------------|------------------------|---------------------------------------------|
| PG:VG                            | 0.0                   | 5.0                                | None                   | 3.0                                         |
| PG:VG (Nicotine)                 | 0.0                   | 5.0                                | Koolada                | FlavorJungle                               |
| PG:VG + Koolada                  | 0.0                   | 5.0                                | Koolada (10% WS-3 in PG) | FlavorJungle Koolada (10% Koolada in PG)   |
| PG:VG + WS-23                    | 0.0                   | 5.0                                | FlavorJungle           | FlavorJungle WS-23 (30% PG)                |
| PG:VG (Nicotine) + Koolada       | 0.0                   | 5.0                                | FlavorJungle           | FlavorJungle WS-23 (30% PG)                |
| PG:VG (Nicotine) + WS-23         | 0.0                   | 5.0                                | FlavorJungle           | FlavorJungle WS-23 (30% PG)                |
The levels of acellular ROS generated by the PG:VG solution (2.02–2.60 µM H₂O₂) were significantly higher (p < 0.05), than those generated by the filtered air control (0.96–1.66 µM H₂O₂) (Fig. 2a). When the levels of acellular ROS generated by the PG:VG solution containing nicotine (5 %) (1.13–1.84 µM H₂O₂) and the filtered air control (0.96–1.66 µM H₂O₂) were compared, the generated ROS levels did not significantly differ (p > 0.05) (Fig. 2b). The levels of ROS generated by the PG:VG with WS-23 solution (1.21–4.16 µM H₂O₂) did not significantly differ from those generated by the aerosolized PG:VG solution nor from the levels of acellular ROS generated by the filtered air control (p > 0.05) (Fig. 3a). However, the levels of acellular ROS generated by the aerosolized e-liquid solution containing PG:VG with nicotine (5 %) and WS-23 (3 %) (1.94–2.95 µM H₂O₂) were significantly higher than those generated by the filtered air control (p < 0.05) (0.96–1.66 µM H₂O₂) (Fig. 3b). In contrast, the levels of acellular ROS generated by the PG:VG solution containing nicotine and WS-23 (1.94–2.95 µM H₂O₂) did not differ significantly from those generated by the PG:VG solution containing nicotine (p > 0.05) (Fig. 3b). When the levels of acellular ROS generated by the PG:VG solution containing nicotine and WS-3 (2.27–2.57 µM H₂O₂) and the filtered air control were compared, the generated ROS levels were significantly different (p < 0.01) (Fig. 4a). However, the difference in acellular ROS levels between aerosolized PG:VG with WS-3 solution and aerosolized PG:VG solution was not significant (p > 0.05) (Fig. 4a). Additionally, the levels of ROS generated by the PG:VG solution with nicotine and WS-3 (1.79–3.35 µM H₂O₂) did not significantly differ from those generated by the aerosolized PG:VG with nicotine solution nor the filtered air control (p > 0.05) (Fig. 4b).

The levels of acellular ROS generated by the PG:VG (50:50) with WS-3 (3 %) solution did not significantly differ from those generated by the PG:VG (50:50) with WS-23 (3 %) solution (p > 0.05) (Fig. 5a). Additionally, neither the difference in acellular ROS levels between the aerosolized PG:VG + WS-3 solution and the filtered air control nor that between the aerosolized PG:VG + WS-23 solution and the filtered air control were significant (p > 0.05) (Fig. 5a). When comparing the levels of ROS generated by the PG:VG with WS-3 and nicotine solution to those generated by the PG:VG with WS-23 and nicotine solution, it did not significantly differ (p > 0.05) (Fig. 5b). Moreover, neither the difference in acellular ROS levels between aerosolized PG:VG + WS-3 solution and the filtered air control nor that between the aerosolized PG:VG + WS-23 solution and the filtered air control were significant (p > 0.05) (Fig. 5b).

Our data show that regardless of nicotine content (0% or 5%), minimal differences in acellular ROS levels exist when comparing the addition of WS-3 and WS-23 to e-liquid base (PG:VG) (Fig. 5a-b). For the acellular ROS assays conducted, smoke generated from a 3R4F research cigarette was used as a positive control with 45.87–49.42 µM H₂O₂ equivalents (n = 2).

3.2. WS-3 and WS-23 modify ROS levels in BEAS-2B cells

Cellular oxidative stress in BEAS-2B cells was assessed through using analyses of fluorescent intensities within cells exposed to CellROX Green reagent; these fluorescent intensities being used as a measure of ROS levels. Fluorescent imaging showed that cells exposed to e-liquid containing PG:VG and cells exposed to PG:VG (nicotine) contained significantly higher levels of cellular ROS than untreated cells (p < 0.05) and (p < 0.01), respectively (Fig. 6a-b). However, the levels of cellular ROS generated by BEAS-2B exposed to PG:VG with WS-23 solution did not significantly differ from those generated by cells exposed to the PG:VG (p > 0.05) nor those generated by untreated cells (p > 0.05) (Fig. 7a). In contrast, the levels of the cellular ROS generated...
by BEAS-2B cells exposed to PG:VG (nicotine) with WS-23 were significantly higher than those generated by untreated cells (p < 0.05), but they did not significantly differ from those generated by cells exposed to PG:VG (nicotine) (p > 0.05) (Fig. 7b). Regarding the levels of ROS generated from cells treated with PG:VG containing WS-3; while they were significantly higher than those generated by untreated cells (p < 0.05), they did not differ significantly from those generated from cells exposed to PG:VG solution (p > 0.05) (Fig. 8a). Similarly, while our data showed that the difference in generated cellular ROS levels between the PG:VG (nicotine) + WS-3 treatment and the PG:VG treatment was not significant (p > 0.05), cells exposed to PG:VG (nicotine) + WS-3 did produce significantly higher levels of ROS than untreated cells (p < 0.01) (Fig. 8b). Additionally, the levels of ROS generated by cells exposed to PG:VG with WS-3 solution did not significantly differ from those generated by the PG:VG with WS-3 solution nor from those generated by the untreated cells (p > 0.05) (Fig. 9a). Similar results were observed when comparing ROS levels generated by cells exposed to both treatments of e-liquid formulations containing nicotine and synthetic coolants. Overall, there was no significant difference between the levels of ROS generated by the PG:VG (nicotine) with WS-3 treatment and the PG:VG (nicotine) with WS-3 treatment (p > 0.05) (Fig. 9b). Cellular ROS levels from cells exposed to e-liquid formulations...
containing either WS-3 or WS-23 were significantly higher than the untreated cells. It should be noted that cell viability was slightly reduced in the PG:VG (Nicotine) + WS-23 treatment groups (personal observation).

4. Discussion

With the surge in e-cig use amongst youth in the United States in 2021 and the recent influx of "iced" e-cig flavors in US marketplaces, there is a greater need to fill the knowledge gap on the safety of inhaling synthetic-coolant additives [28]. Our study sought to determine whether adding a widely used synthetic coolants, WS-3 and WS-23, to nicotine and non-nicotine-containing e-liquids modifies the level of acellular ROS generated in e-cig aerosols and cellular ROS levels generated by the BEAS-2B airway epithelial cell lines. When comparing acellular ROS levels generated by the PG:VG with nicotine and WS-23 (3.0%) solution and the PG:VG solution, the levels of acellular ROS did not differ significantly. Similarly, there was no significant difference between acellular ROS levels from PG:VG (Nic) with WS-3 groups and PG:VG (Nic) with WS-23 groups as those in Figs. 3 and 4. Fig. 5A-5B include the air, PG:VG, PG:VG (Nic), PG:VG with WS-3, and PG:VG (Nic) with WS-23 groups in order to compare the acellular ROS levels between specific e-liquid formulations.
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and the PG:VG with nicotine solution, the levels of generated cellular ROS did not significantly differ. However, we did note that the difference in BEAS-2B cell generated ROS levels between the PG:VG (50:50) treatment and the untreated cells (p < 0.01) was lower than the difference in BEAS-2B generated ROS levels between the PG:VG (nicotine) treatment and the untreated cells (p < 0.01). Also, we observed significantly higher levels of ROS production by BEAS-2B cell treated with PG:VG (nicotine) with WS-23 (3 %) when compared to untreated cells (p < 0.01). Also, we observed a significant difference in BEAS-2B ROS production between the PG:VG with WS-3 (3 %) treatment and untreated cells (p < 0.05). Our data suggests that the addition of WS-23 and WS-3 by themselves to e-liquid base solution, PG:VG, and e-liquid based solution containing nicotine, PG:VG (nicotine), does not lead to significantly modifying cellular ROS levels generated by BEAS-2B cells.

However, these differences in generated cellular ROS levels between e-liquid base solution treatments and synthetic coolant-containing e-liquid base solution treatments are noteworthy and were observed regardless of nicotine content (5 %). Cellular ROS levels were increased by WS-3 and WS-23 treated in e-cig nicotine groups.

Regarding our acellular ROS data, we see that the difference in acellular ROS levels between the aerosolized WS-23 (3 %) solution and the filtered air control (p > 0.05) is higher than that between the aerosolized PG:VG with nicotine solution and the filtered air control (p > 0.05). Likewise, our acellular ROS data suggest that adding WS-23 to nicotine-containing e-liquid base leads to noteworthy changes in generated acellular ROS levels. Our findings are similar to that of previous studies showing that treatments with e-liquids induce significant levels of ROS production in BEAS-2B cells when compared to untreated

Fig. 6. Comparative analysis of cellular ROS levels generated from treatments with PG:VG, PG:VG with nicotine, and an untreated control. Comparisons between the cellular ROS levels was conducted using CellROX green reagent generated by BEAS-2B which were left untreated and, (a) cells treated with PG:VG (50:50) and PG:VG containing nicotine for four hours. After an additional four hours, cell medium was aspirated, and the live cells were stained with 5 μM CellROX Green Reagent (in DMEM 0% FBS). After PFA fixation, nuclei were counterstained with Hoechst stain. Data are represented as mean ± SEM, and significance was determined using an unpaired t-test. * p < 0.5 and **p < 0.01 (n=3). (b) ROS labelled nuclei were assessed by Cytation 5 imaging (BioTek) reader and CellROX fluorescent signals were analyzed using Image J software (n = 3). Scale bar = 200 μm. The fluorescent images showing stained untreated cells were common to both the treatment groups, and were used for comparative analyses between the groups. Histograms as Arbitrary Unit (AU) representing CellRox Fluorescence in untreated cells (a) were used to prepare the comparative graph shown (b).
Fig. 7. Comparative analysis of cellular ROS levels generated by PG:VG, PG:VG + WS-23, PG:VG with nicotine, PG:VG with nicotine + WS-23, and an untreated control. Comparisons between the cellular ROS levels was conducted using CellROX green reagent generated by BEAS-2B which were left untreated and, (a) Cells were treated with PG:VG and PG:VG + WS-23, (b) Cells were treated with both PG:VG containing nicotine and PG:VG containing nicotine and WS-23 for four hours. After an additional four hours, cell medium was aspirated and the live cells were stained with 5 μM CellROX Green Reagent (in DMEM 0% FBS). After PFA fixation, nuclei were counterstained with Hoechst stain. Data are represented as mean ± SEM, and significance was determined using One-way ANOVA. * p < 0.05 and **p < 0.01 (n = 3). ‘NS’ is abbreviated for “Non-Significant” versus air control (p > 0.05). Data were compared with images on Fig 6. ROS labelled nuclei were analyzed by Cytation 5 imaging (BioTek) reader and CellROX fluorescent signals were measured using Image J software. Scale bar = 200μm.

Fig. 8. Comparative analysis of cellular ROS levels generated by PG:VG, PG:VG + WS-3, PG:VG with nicotine, PG:VG with nicotine + WS-3, and an untreated control. Comparisons between the cellular ROS levels was conducted using CellRox green reagent generated by BEAS-2B which were left untreated and those which were treated; treatments are as follow: (a) Cells were treated with PG:VG and PG:VG + WS-3, (b) Cells were treated with both PG:VG containing nicotine and PG:VG containing nicotine + WS-3 for four hours. After an additional four hours cell medium was aspirated and the live cells were stained with 5 μM CellRox Green Reagent (in DMEM 0% FBS). After PFA fixation, nuclei were counterstained with Hoechst stain. Data are represented as mean ± SEM, and significance was determined using One-Way ANOVA. * p < 0.05 and **p < 0.01 (n = 3). ‘NS’ is abbreviated for “Non-Significant” versus air control (p > 0.05). Data were compared with images on Fig 6. ROS labelled nuclei were assessed by Cytation 5 imaging (BioTek) reader and CellROX fluorescent signals were analyzed using Image J software. Scale bar = 200μm.
cells [29]. Regarding our understanding of the potentially harmful effects of WS-23, our data seems to suggest that WS-23 itself has a limited impact in altering e-cig-generated acellular ROS levels as well as a limited effect on modifying the levels of ROS generated by BEAS-2B cells. Similarly, our data also seems to suggest that the addition of WS-3 to e-liquids does not significantly modify e-cig generated acellular ROS levels nor cellular ROS levels generated by treated BEAS-2B cells. Concerning the findings of other studies investigating the physiological effects of using coolant containing e-liquids, using human bronchial epithelial cell cultures, previously, another study found that treatment with menthol significantly increased mitochondrial ROS via the TRPM8 receptor [30]. BEAS-2B cell exposures reported by these investigators consisted of aerosol treatments, in which cells were exposed for two separate 1.5-minute durations, separated by an incubation period; however, our cell treatment protocol consisted of a direct e-liquid exposure and those apart of the respective control group [34]. Neutrophils are a major sources of endogenous ROS production. Likewise, future studies aimed at understanding the role of WS-23 in modulating e-cig induced oxidative stress should involve measurements of intracellular and extracellular ROS using isolated Polymorphonuclear Neutrophils (PMNs) [35]. More specifically, PMNs isolated from blood collected from mice exposed to aerosolized e-liquids of varying WS-23 concentrations can be analyzed via luminol enhanced chemiluminescence exposure [35]. The proposed experiment can provide insight into the differences between intra-and extra-cellular ROS of PMNs isolated from mice exposed to various concentration of WS-23 as reported recently [35].

Recent studies investigating the potential of exposures to synthetic coolant-containing e-liquids may have an effect on pulmonary pathophysiology including cytotoxicity evaluations on BEAS-2B cells exposed to aerosols generated by various flavored e-cigs containing either WS-3, WS-23, or both [33]. Omaiye et al. [33] used Lactase Dehydrogenase (LDH), Neutral Red Uptake (NRU), and MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) tetrazolium reduction assays to assess the role of exposure to aerosols generated by e-liquids containing WS-3 and WS-23 have an effect in inducing cytotoxicity in human bronchial epithelial cells [33]. Similar to this study [29], our study involved analyzing the cellular responses of BEAS-2B cells to e-liquids containing either WS-3 or WS-23, and showed some cytotoxic responses when WS-3 and WS-23 combined with e-cig nicotine. However, in contrast to Omaiye et al. [33], which assessed the cytotoxicity induced by different treatments, our study assessed differences in ROS production. Additionally Omaiye et al. [33] used aerosol exposures, whereas our study conducted cell-culture exposures via direct stimulation [32]. We further determined the cytotoxicity of the cooling agents, when WS-23 was treated to BEAS2B cells for hazard characterization. Various toxicological parameters were calculated in a dose-response using a linear response phase (0.05–3 mg/mL), and found the dose > 2.0 mg/mL was more cytotoxic (based on significant LDH release). Further work is in progress to determine the LC50/IC50 of these synthetic coolants.

In rodent studies, rats exposed to aerosolized e-liquid containing WS-23 at tested doses (via acute and subacute exposures) found no substantial changes in histopathologic analyses of vital organs nor relative organ weights [34]. This same study, via a bronchoalveolar lavage fluid (BALF) analysis, found no significant difference in neutrophil concentration between rats which had undergone repeated 28-day WS-23 exposure and those apart of the respective control group [34]. Neutrophils are a major sources of endogenous ROS production. Likewise, future studies aimed at understanding the role of WS-23 in modulating e-cig induced oxidative stress should involve measurements of intracellular and extracellular ROS using isolated Polymorphonuclear Neutrophils (PMNs) [35]. More specifically, PMNs isolated from blood collected from mice exposed to aerosolized e-liquids of varying WS-23 concentrations can be analyzed via luminol enhanced chemiluminescence exposure [35]. The proposed experiment can provide insight into the differences between intra-and extra-cellular ROS of PMNs isolated from mice exposed to various concentration of WS-23 as reported recently [35]. Regarding our understanding of the effects of other e-liquid coolant additives, using human bronchial epithelial cell cultures, one study found that treatment with menthol significantly increased mitochondrial ROS via the TRPM8 receptor [30]. However, in contrast to our study, the studies of Nair et al. [30] BEAS-2B cell exposures consisted of aerosol treatments, in which cells were exposed for two separate 1.5-minute durations, separated by an incubation period; however, our cell treatment protocol consisted of a direct e-liquid treatment for 4 h. Hence, understanding the role of WS-3 and WS-23 in modulating e-cig-induced oxidative stress should involve measurements of intracellular and extracellular ROS using isolated Polymorphonuclear Neutrophils (PMNs) [35] and macrophages, airway immune cells and epithelial cells.

Our findings concur with previous studies showing that aerosolized e-liquids contain significant levels of acellular ROS and induce significant levels of cellular ROS in pulmonary epithelial cells [19,21]. Regarding previous studies that analyzed acellular ROS levels within “cool/iced” flavored e-cigs, one study found differences in generated-acellular ROS levels between Tobacco-Derived Nicotine (TDN) and Tobacco-Free Nicotine (TFN) among cool/iced flavored
e-cigs were minimal compared to tobacco and fruit flavors [20].

Regarding limitations in our study, our study did not include the treatment of airway epithelial cells with aerosolized e-liquids. Previous studies have shown that treatments with aerosolized e-liquids induce significant levels of ROS production in Human Bronchial Epithelial cells (BEAS-2B) [29]. Epithelial cells lining the airways are the first structural cell targets of any inhaled substances [36] and, the inhalation of e-cigs results in pulmonary epithelial cells being exposed to aerosols generated from e-liquids. In comparison to aerosol exposures, e-cig cell exposures using direct stimulation allows for a more precise control of dosage and a more expeditious analysis of different types of e-liquids consisting of various flavors, nicotine concentrations, and coolant concentrations [32]. However, cell cultures conducted with e-liquids via direct stimulation do not emulate the process associated with the actual usage of e-cigs which is the inhalation of aerosolized e-liquids into the lungs ("vaping.")

Future studies analyzing the role of WS-3 and WS-23 in potentially modifying ROS generated from BEAS-2B cells should utilize cell culture exposures via e-cig aerosols for analyzing the cellular oxidative-stress levels [29,37]. Through this proposed assay, an understanding of how exposure to aerosolized synthetic coolants affects mitochondrial ROS production can be obtained. The reasoning for our reason to conduct e-cig cell exposures via direct stimulation rather than using aerosols lied in our understanding of that more studies investigating how the direct addition of e-liquids to pulmonary cells impacts cellular ROS responses are needed [31]. Consequently, after having observed significant levels of ROS generated by e-liquids containing WS-3 and WS-23 in cell-free conditions, we also determined how the addition of these synthetic coolants to e-liquids directly treated to pulmonary cells impact the cellular ROS levels. However, our study has shown that a wide range of ROS generated by e-liquids containing WS-3 and WS-23 to e-liquids, either 0 % or 5 % nicotine, has a minimal effect on modifying the acellular ROS levels from aerosolized e-liquid base solution or the cellular ROS levels generated by BEAS-2B cells exposed to e-liquid base solution. However, ROS levels were increased by WS-3 and WS-23 in e-cig-e-liquid nicotine groups. Thus, these preliminary findings do strongly suggest the need for further evaluation on the potential health risks associated with inhaling the newly marketed e-cigs containing synthetic coolants. Specifically, our findings do underscore the need for further investigation into the role of WS-3 and WS-23 e-cigarettes in disrupting the endogenous oxidant and antioxidant balance in airways upon inhalation.

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CReditT authorization contribution statement

Conceptualization, I.R.; methodology, I.R.; assay performance: S.Y, SBS, software, S.Y., SBS; validation, S.Y, SBS, I.R.; formal analysis, S.Y., SBS; investigation, S.Y, SBS; re-sources, I.R.; data curation, S.Y., SBS; software, S.Y., SBS; validation, S.Y, SBS, I.R.; formal analysis, S.Y., SBS, I.R.; project administration, I.R.; funding acquisition, I.R. All authors have read and agreed to the published version of the manuscript.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

We declare that we have provided all the data in figures.

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Informed consent statement

Not applicable; no human subjects were involved.

Conflicts of interest

The authors declare no conflicts of interest.

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