Comparative Study on the Chronic Vascular Responses Induced by Regular Versus Occasional Waterpipe Smoke Inhalation in Mice

Naserddine Hamadia  Sumaya Beegamb  Nur Elena Zaaba b  Ozaz Elzaki b  Badreldin H. Ali c  Abderrahim Nemmarb,d

aDepartment of Life and Environmental Sciences, College of Natural and Health Sciences, Zayed University, Abu Dhabi, United Arab Emirates, bDepartment of Physiology, College of Medicine and Health Sciences, United Arab Emirates University, Al Ain, United Arab Emirates, cDepartment of Pharmacology and Clinical Pharmacy, Sultan Qaboos University, Muscat, Al-Khod, Oman, dZayed Center for Health Sciences, United Arab Emirates University, Al Ain, United Arab Emirates

Key Words
Waterpipe smoke regimen • Oxidative stress • Inflammation • Thrombogenicity • Vascular alterations

Abstract
Background/Aims: Waterpipe smoke (WPS) is the second most prevalent form of smoking in the world. There are ample evidences about the vascular alterations caused by regular WPS (Reg-WPS). Nonetheless, comparison of the chronic vascular response induced by regular versus occasional WPS (Occ-WPS) exposure is very scarce. Methods: We investigated, in BALB/c mice, the effects of Occ-WPS (30 minutes/day, 1 day/week) versus Reg-WPS (30 minutes/day, 5 days/week) for 6 months on thrombogenicity and platelet aggregation in vivo and in vitro. Moreover, various markers of endothelial integrity, inflammation and oxidative stress were assessed by enzyme-linked immunosorbent assay and colorimetric assay. Control mice were exposed to air. Results: Our results showed that either Occ-WPS or Reg-WPS exposure shortened the thrombotic time in pial microvessels in vivo. Moreover, in pial venules, this effect was more marked in Reg-WPS group (-47%) compared with Occ-WPS (-34%). Similarly, exposure to either Occ-WPS or Reg-WPS reduced the prothrombin time and activated partial thromboplastin time. Platelet count was increased only in Reg-WPS exposure. Exposure to either Occ-WPS or Reg-WPS induced platelet aggregation in vitro. In addition, there was a statistically significant difference between Occ-WPS and Reg-WPS groups in platelet count and aggregation. Plasma concentration of tissue factor (+98%), P-selectin (+14%) and E-selectin (+16%) were significantly increased in Occ-WPS group compared with air exposed group. Likewise, compared with air group Reg-WPS caused an increase in

Professor A. Nemmar
United Arab Emirates University, College of Medicine and Health Sciences, Department of Physiology
P.O. Box 17666, Al Ain (United Arab Emirates)
Tel. +971-37137533, Fax +971 3 7671966, E-Mail anemmar@uaeu.ac.ae; anemmar@hotmail.com
concentration of tissue factor (+193%), P-selectin (+21%) and E-selectin (+42%). Nevertheless, only Reg-WPS induced a decrease (-38%) in the plasma concentration of tissue plasminogen activator. Notably, our results showed a statistically significant difference between Occ-WPS and Reg-WPS groups in the concentration of tissue factor. Erythrocyte numbers, hemoglobin concentration, hematocrit and lactate dehydrogenase activity were augmented only in Reg-WPS group compared with either control or Occ-WPS groups. Likewise, only Reg-WPS induced an increase in proinflammatory cytokines, tumor necrosis factor-α and interleukin-1β compared with either control or Occ-WPS groups. However, markers of oxidative stress including 8-isoprostane and total antioxidants were enhanced in both Occ-WPS and Reg-WPS compared with control group.

**Conclusion:** Our data confirm the vascular toxicity of the chronic Reg-WPS exposure and shows that even occasional chronic exposure to WPS caused thrombosis, platelet aggregation, endothelial alterations and oxidative stress. The latter findings are an additional cause of concern about the long-term toxicity of occasional waterpipe smoking.

## Introduction

Hookah also known as waterpipe, narghile, argileh, shisha, hubble-bubble, goza, borri, qaylan, chica, and mada’a is a form of traditional smoking made of tobacco pipe with a long stretchy tube that draws the smoke through water contained in a bowel [1]. This practice is ceased to be a middle-eastern culture where we can find it around bars in college campuses in the western hemisphere such as the United States and Europe [2]. The popularity of WPS is rising among trendy youth, university students, and even high-school-aged children [3, 4]. There is an erroneous belief among WPS smokers that WPS is less harmful compared with cigarette smoking (CS) [5], and this is based on the misconception that the filtration occurs when the smoke passes through the water will diminish the levels of tar, nicotine, and other toxins [6]. A meta-analysis conducted by Neergaard et al. suggested that daily WPS generates nicotine equivalent to daily smoking of 10 cigarettes. Furthermore, occasional WPS is comparable to smoking two cigarettes during 24-hr [7]. There are increasing epidemiological evidences suggesting the potential of WPS of becoming a major public health problem in most Arab countries [8].

According to the 2015 WHO recommendation and several other reports, WPS probably is addictive as in other forms of tobacco and its consumption may lead to the same detrimental effects caused by cigarette [1, 9-11]. In support of the latter, it has been reported that WPS contains ample quantities of toxicants including nicotine, tar, carbon monoxide, polycyclic aromatic hydrocarbons, nitrosamines, volatile aldehydes, phenols and heavy metals, catechol and hydroquinone [9, 10]. Clinical and experimental studies including our own, demonstrated that WPS causes various diseases and is detrimental to various systems comparable to cigarette smoke, including cancer [10, 12], reproductive system injuries [13], adverse cardiopulmonary effects [14] and metabolic syndrome development which is considered a major risk factor for developing thrombosis [15, 16]. Furthermore, it leads to low birth weight [17, 18] and increased neonatal death rate [19].

It has been reported that occasional or regular WPS users are susceptible to be regular cigarette smokers, signifying the fact that WPS may be a potential leading way for CS smoking [20]. Most of the research on smoking focused on regular smoking. However, recent population data indicate that 22%–33% of US adult smokers are intermittent smokers who do not smoke every day [21, 22]. Moreover, the number of non-daily smokers is anticipated to rise due to the high cost of tobacco and restriction rules in public areas [23, 24]. The classification of occasional smokers is based on the amount and frequency of smoking. The latter includes four categories, (1) smoking two days in the past week, (2) smoking at least 100 cigarettes in their lifetime, but not smoking daily and having smoked in the last 12 months, (3) smoking between 1 and 10 cigarettes per day in the last 30 days, or (4) smoking at least once a week but not every day [25, 26].
Accumulating evidences suggest that light and non-daily cigarette smoking are associated with increased morbidity including a similar risk of cardiovascular diseases as heavier smokers, different types of cancer, respiratory diseases and reproductive health problems [27-31]. Hence, as far as we are aware, there is a scarcity of data on the vascular and systemic effects of occasional WPS and in order to address that, we aimed to investigate the mechanisms underlying the effects of six months exposure to either Occ-WPS (30 minutes/day, 1 day/week) or Reg-WPS (30 minutes/day, 5 days/week) on thrombogenicity, platelet aggregation, endothelial integrity, inflammation, and oxidative stress.

**Materials and Methods**

**Animals and WPS exposure**

BALB/c mice of both genders aged 6–8 weeks, weighing 20–25 g (Taconic Farms Inc., Germantown, NY, USA) were housed in the local central animal facility of the College of Medicine and Health Sciences and maintained in controlled light cycle (12-h light:12-h dark cycle), humidity of 60% and, controlled-temperature (22 ± 1°C). Animals had free access to water and food ad libitum.

After one week of acclimatization to the experimental conditions, the mice were indiscriminately separated into 3 groups, air (control), Occ-WPS and Reg-WPS. The WPS exposure protocol has been performed according to previously described methods [32, 33]. Mice were placed in soft restraints and connected to the exposure tower. Using a nose-only exposure system connected to a waterpipe device, the animals were exposed to either air or WPS by their noses (inExpose System, SCIREQ, Canada). Animals were exposed to mainstream WPS generated by commercially available apple-flavored tobacco. For each daily session, 10 grams of tobacco were placed into the WPS head. At the end of WPS exposure session, the remaining tobacco was discarded.

Control mice were exposed to air only. The duration of the session was 30 min/day. Regarding Reg-WPS, mice were exposed to WPS, 5 days/wk for 6 months [32, 33] and for Occ-WPS group, mice were exposed to WPS, 1 day/wk for 6 months. The WPS exposure procedure was monitored by a computerized system. A computer-monitored puff was produced every 1 min (consisting of a 2 s puff time of WPS after that a 58 s of fresh air). Twenty-four hours following the last exposure session, various vascular endpoints were assessed. The total number of animals used in the present study was 83. They were distributed as described below.

**Induction of thrombosis in pial arterioles and venules of mouse photochemically**

In vivo pial arteriolar and venular thrombogenesis was assessed in separate set of mice at the end of the 6 months exposure period to either Occ-WPS (n=6) or Reg-WPS (n=7) or air (n=7), according to a previously described technique [32, 33]. Briefly, the trachea was intubated after the induction of anesthesia with urethane (1 mg/g body wt ip), and a 2-Fr venous catheter (Portex, Hythe, UK) was inserted in the right jugular vein for the administration of fluorescein (Sigma- Aldrich). Thereafter, a craniotomy was first performed on the left side, using a microdrill, and the dura was stripped open. Only untraumatized preparations were used, and those showing trauma to either the microvessels or underlying brain tissue were discarded. Animals were then placed on the stage of a fluorescence microscope (Olympus, Melville, NY) attached to a camera and DVD recorder. A heating mat was placed under the mice, and body temperature was raised to 37°C, as monitored by a rectal thermoprobe connected to a temperature reader (Physitemp Instruments). The cranial preparation was moistened continuously with artificial cerebrospinal fluid of the following composition (in mM): 23 NaHCO₃, 5 KCl, 3 NaH₂PO₄, 4 MgSO₄, 124 NaCl, 2.5 CaCl, and 10 glucose (pH 7.3–7.4). A field containing arterioles and venules of 15–20 µm in diameter was chosen. Such a field was taped before and during the photochemical insult. The photochemical insult was carried out by injecting fluorescein (0.1 ml/mouse of 5% solution) via the jugular vein, which was allowed to circulate for 30–40 s. The cranial preparation was then exposed to stabilized mercury light. This combination produces endothelium injury of the arterioles and venules. This, in turn, causes platelets to adhere at the site of endothelial damage and then aggregate. The platelet aggregates and thrombus formation grow in size until complete arteriolar or venular occlusion. The time from the photochemical injury until full vascular occlusion (time to flow stop) in arterioles and venules was measured in seconds. At the end of the experiments, animals were euthanized by an overdose of urethane.
Prothrombin time (PT) and activated partial thromboplastin time (PTT) measurement in plasma in vitro

In a separate set of mice, at the end of the 6 months exposure period to either Occ-WPS (n=6) or Reg-WPS (n=6) or air (n=7), animals were anesthetized, and the blood was withdrawn from the inferior vena cava and placed in citrate solution (3.2%) (ratio of the blood to anticoagulant: 9:1) for PT and aPTT measurement according to a previously described technique [32, 33]. The PT was measured on freshly collected platelet-poor plasma with human relipidated recombinant thromboplastin (Recombiplastin, Instrumentation Laboratory, Orangeburg, NY) in combination with a Merlin coagulometer (MC 1 VET, Merlin). The aPTT was measured with automated aPTT reagent from bioMerieux (Durham, NC) using a Merlin coagulometer (MC 1 VET, Merlin). Normal plasma used as the reference for both PT and aPTT was prepared by pooling equal portions of platelet-poor plasmas from the blood of seven untreated mice.

Platelet aggregation in mouse whole blood

The platelet aggregation assay in whole blood obtained from a separate set of mice exposed to Occ-WPS (n=7) or Reg-WPS (n=7) or air (n=6) for 6 months was performed as described before [32, 33]. After anesthesia, the blood was withdrawn from the vena cava and placed in citrate (3.2%). Aliquots of 100 µl were added to the wells of a Merlin coagulometer (MC 1 VET, Merlin, Lemgo, Germany). After incubation with ADP (1 µM) for 3 min at 37.2°C, Blood samples were stirred for 3min. At the end of this period, 25µl samples were removed and fixed on ice in 225 ml cellFix (Becton Dickinson, Franklin Lakes, NJ). ADP induction of platelet aggregation is reflected by a decrease in counted single platelets in the blood obtained from Occ-WPS or Reg-WPS or air-exposed mice.

Measurement of tissue factor, tissue plasminogen activator (tPA), P-selectin and E-selectin concentrations in the plasma

The concentrations of tissue factor, P-selectin and E-selectin in the plasma obtained from mice exposed to Occ-WPS (n=8) or Reg-WPS (n=8) or air (n=8) were measured by enzyme-linked immunosorbent (ELISA) assays using commercially available kits obtained from R&D systems (Duo Set, Minneapolis, MN, USA). Regarding tPA, the ELISA kit was obtained from Molecular Innovations (Novi, Michigan, USA).

Blood count

Erythrocyte and platelet count, hemoglobin concentration and hematocrit were assessed in the same mice reported above. The animals were anesthetized intraperitoneally with pentobarbital sodium (45 mg/kg), and blood was then drawn from the inferior vena cava in EDTA (4%). A sample was used for platelet and red blood cell counts and hematocrit determination using an ABX VET ABC Hematology Analyzer with a mouse card (ABX Diagnostics, Montpellier, France). The remaining blood was centrifuged for 15 min at 4 °C at 900g, and the plasma samples obtained were stored at -80°C until further analysis.

Measurement of lactate dehydrogenase (LDH) activity in the plasma

The LDH activity was measured using commercial kits (Sigma Chemical, St. Louis, MO, USA) which determine the conversion of lactate to pyruvate in the presence of LDH with an equivalent lessening of NAD in the plasma of the mice reported above. The formation of NADH from the reaction can show a difference when measured in absorbance at 340 nm.

Measurement of TNFα, IL-1β, 8-isoprostane and total antioxidant capacity levels in the plasma

The proinflammatory and oxidative stress markers were assessed in the plasma of the mice reported above. The concentration of the pro-inflammatory cytokines, tumor necrosis factor-α (TNFα) and interleukin-1β (IL-1β) were measured by ELISA assays using commercially available kits obtained from R&D systems (Duo Set, Minneapolis, MN, USA). Protein content in each sample was measured by Bradford’s method, as described earlier [32, 33].

The levels of 8-isoprostane and total antioxidants were quantified according to the manufacturer’s instructions provided in the commercially available assay kits obtained from Cayman Chemicals (Michigan, USA).
Statistical analysis

All graphs were produced using GraphPad Prism Version 7 for Windows software (GraphPad Software Inc, San Diego, CA, United States). Data were expressed as means ± SEM. To assess whether the measured parameters were normally distributed, the Shapiro-Wilk normality test was used. Normally distributed data were tested by using one-way analysis of variance ANOVA followed by Holm-Sidak’s multiple comparisons test. P<0.05 was considered significant.

Results

Photochemically induced thrombosis in pial arterioles and venules of mouse in vivo

Fig. 1 shows the effect of Occ-WPS or Reg-WPS exposure on photochemically induced thrombosis in both pial arterioles and venules of mice. The exposure to either Occ-WPS or Reg-WPS showed a prothrombotic tendency in both pial arterioles and venules. Compared with air-exposed mice, the occlusion time in pial arterioles was significantly shortened following exposure to either Occ-WPS (P<0.00001) or Reg-WPS (P<0.00001) (Fig. 1A). Similarly, in pial venules, Occ-WPS (P<0.00001) or Reg-WPS (P<0.00001) caused a marked reduction of the thrombotic occlusion time. Moreover, in the pial venules there was a statistical significance between Occ-WPS and Reg-WPS groups (P<0.00001) (Fig. 1B).

PT and aPTT

Fig. 2 illustrates the impact of 6 months exposure to either Occ-WPS or Reg-WPS on PT and aPTT in mouse plasma. The shortening of PT and aPTT is indicative of hypercoagulability tendency in the blood. The exposure to Occ-WPS or Reg-WPS induced a statistically significant shortening of PT (P<0.00001, Fig. 2A) and aPTT (P<0.00001, Fig. 2B) compared with plasma obtained from air-exposed mice.

Platelet numbers and platelet aggregation in vitro

As shown in Fig. 3A, Occ-WPS exposure did not affect platelet number in the blood compared with air exposed group. However, Reg-WPS exposure induced significant increase in the platelet numbers (P=0.03). In addition, there was a statistically significant increase in the number of platelets in Reg-WPS group compared with Occ-WPS one (P=0.04). We found that whole blood obtained from mice exposed to either Occ-WPS or Reg-WPS and incubated in vitro with ADP displayed a significant platelet aggregation (P=0.005-P=0.00001) compared with blood collected from mice exposed to air. Moreover, there was a statistical significance between Occ-WPS and Reg-WPS groups (P=0.005) (Fig. 3B).

Fig. 1. Thrombotic occlusion time in pial arterioles (A) and venules (B) at the end of 6 months exposure to air, occasional waterpipe smoking (Occ-WPS) or regular WPS (Reg-WPS). Data are mean ± SEM (n=6-7).
Concentrations of tissue factor, tPA, P-selectin and E-selectin in the plasma

Fig. 4 exemplifies the effect of Occ-WPS or Reg-WPS on the plasma concentrations of tissue factor (Fig. 4A), tPA (Fig. 4B), P-selectin (Fig. 4C) and E-selectin (Fig. 4D). Compared with air exposed group, exposure to Occ-WPS or Reg-WPS caused a significant increase ($P<0.00001$) in tissue factor concentration in the plasma and there was statistical significance between Occ-WPS and Reg-WPS groups ($P<0.00001$). Regarding tPA, a significant reduction was observed only in the plasma of mice exposed to Reg-WPS ($P=0.006$). In addition, there was a statistical significance between Occ-WPS and Reg-WPS groups ($P=0.02$). The plasma concentrations of the adhesion molecule P-selectin were significantly increased following Occ-WPS ($P=0.004$) or Reg-WPS ($P=0.0001$) exposure compared to air exposed mice. In addition, the concentration of E-selectin was statistically increased in both Occ-WPS ($P=0.01$) and Reg-WPS ($P<0.00001$) groups compared with air exposed mice. Moreover, there was a statistical significance in the concentration of E-selectin between Occ-WPS and Reg-WPS groups ($P=0.0006$).
Erythrocyte numbers, hemoglobin concentration, and hematocrit

The erythrocyte numbers and hematocrit were not affected following Occ-WPS exposure for a period of 6 months compared to air exposed group. However, we observed a significant increase in the erythrocyte numbers (P=0.0004) and hematocrit (P=0.0002) following exposure to Reg-WPS (Fig. 5A and B). Similarly, as shown in Fig. 5C, hemoglobin concentration was only significantly increased (P=0.003) in Reg-WPS (Fig. 5B) group compared with air-exposed mice. In addition, there was a statistical significance between Occ-WPS and Reg-WPS groups for erythrocyte numbers (P=0.02), hematocrit (P=0.01) and hemoglobin concentration (P=0.01).

Lactate Dehydrogenase (LDH) activity in the Plasma

Our results showed no statistical changes in LDH activity in the plasma of mice occasionally exposed to WPS compared with air exposed group. In contrast, we observed a significant increase (P=0.0001) in the activity of LDH in the plasma of mice regularly exposed to WPS for 6 months (Fig. 6). Moreover, there was a statistical significance between Occ-WPS and Reg-WPS groups (P=0.0001).

Concentrations of pro-inflammatory cytokines in the plasma

The concentrations of pro-inflammatory cytokines, TNFα and IL-1β are shown in Fig. 7. We observed that Occ-WPS exposure for a period of 6 months did not have a significant effect on the plasma concentrations of TNFα and IL-1β compared with air exposed group. On the contrary, compared with air exposed group, the concentration of TNFα (P=0.003) and IL-1β (P=0.0001) in the plasma were significantly increased following Reg-WPS exposure. Moreover, both TNFα (P=0.007) and IL-1β (P=0.0006) were significantly increased in Reg-WPS group compared with Occ-WPS group.
**Fig. 5.** Erythrocyte numbers (A), hematocrit (B) and hemoglobin concentration (C) at the end of 6 months exposure to air, occasional waterpipe smoking (Occ-WPS) or regular WPS (Reg-WPS). Data are mean ± SEM (n=8).

**Fig. 6.** Lactate Dehydrogenase (LDH) activity in the plasma at the end of 6 months exposure to air, occasional waterpipe smoking (Occ-WPS) or regular WPS (Reg-WPS). Data are mean ± SEM (n=8).
Levels of Oxidative Stress Markers in the Plasma

The quantification of the levels of 8-isoprostane and total antioxidants following Occ-WPS or Reg-WPS exposure is illustrated in Fig. 8. Our results showed that the concentration of 8-isoprostane, a marker of lipid peroxidation in the plasma were significantly increased following either Occ-WPS or Reg-WPS compared with air exposed group (P=0.006, Fig. 8A). Likewise, compared with air exposed group, exposure to either Occ-WPS (P=0.006) or Reg-WPS (P=0.009) induced a significant increase in total antioxidants capacity (Fig. 8B).

Discussion

The current study demonstrated that chronic Reg-WPS exposure triggers vascular toxicity and showed that even occasional chronic exposure to WPS caused thrombosis, platelet aggregation, endothelial alterations and oxidative stress.

Occasional smoking poses a real public health because it is always overlooked by the users due to numerous reasons that range from the erroneous perception that Occ-WPS is harmless to the fact that nicotine dependence can be escaped [34]. It is well known that intermittent or light smoking are associated with cardiopulmonary adverse effects and cancer [30, 35]. While the chronic vascular effects of regular WPS exposure have been investigated, little is known about the possible effects of Occ-WPS on thrombogenicity, blood vessels integrity, inflammation and oxidative stress.
It has been reported that smoking enhances systemic coagulability, through increasing circulating thrombin activity, fibrinogen levels and platelet activation [36]. As a novelty in the current work, our results show that either Occ-WPS or Reg-WPS exposure for a period of 6 months significantly shortened the thrombotic occlusion time in pial arterioles and venules in mice suggesting a prothrombotic impact of WPS regardless of the exposure regimen. Moreover, in pial venules, the thrombotic occlusion time was more markedly shortened in Reg-WPS group compared with Occ-WPS group. These findings provide evidence that WPS is not safe as the general misperception suggests. In line with our findings, it has been shown recently that acute exposure to WPS for 7 days increased the tendency to carotid artery thrombosis [37]. A similar effect was demonstrated following short-term exposure to e-cigarette [38, 39]. We have reported previously that short and long-term exposure to WPS leads to thrombogenicity [33, 40]. Clinically, it has been shown that smokers of waterpipe and cigarette had analogous degrees of endothelial dysfunction compared with nonsmokers [41].

Along with prothrombotic effects, we observed the shortening of PT and aPTT in mice following Occ-WPS or Reg-WPS exposure illustrating that Occ-WPS is as harmful as regular smoking in terms of blood hypercoagulability. Several studies including ours, evidenced the procoagulant effect of WPS [42]. Clinical studies have reported that smoking induces hypercoagulability state characterized by platelet count increase and shortening of PT and aPTT [43, 44].

It has been shown that cigarette smoke induces hemostatic disturbance through alteration in platelet aggregation [45, 46]. Previously, we showed platelet aggregation after short-and long-term WPS exposure [33, 40]. The current data show the numbers of circulating platelets were significantly increased in Reg-WPS and that both exposure regimens induced platelet aggregation in vivo. In addition, there was a statistical significance between Occ-WPS and Reg-WPS groups with a higher impact on platelets aggregation observed following Reg-WPS versus Occ-WPS exposure.

P-selectin and E-selectin are cell adhesion molecules that play a major role in the interaction of platelets and endothelial cells with neutrophils and monocytes [47]. Our data show that exposure to Occ-WPS is as effective as Reg-WPS in the enhancement of the plasma concentrations of P-selectin and E-selectin as platelet and endothelial activation markers respectively, suggesting an abnormal vascular reactivity [48]. Moreover, our results showed a higher concentration of E-selectin in Reg-WPS group compared with Occ-WPS group. This observation is well-matched with those of clinical studies that found a significant increase in systemic P-selectin concentration following exposure to conventional cigarettes or e-cigarette vapor with nicotine [49, 50]. In line with the later, in vitro study showed cigarette smoke extract exposure caused an increase in the expression of the adhesion molecule E-selectin in human umbilical vein endothelial [49] and aortic endothelial cells [51, 52]. Moreover, an increase in the expression of E-selectin in the endothelium of human atherosclerotic lesions [51]. Taking these finding together, Occ-WPS or Reg-WPS exposure triggers the secretion of such adhesion molecules in the bloodstream indicating vascular damage and increased thrombosis. In line with our findings, a clinical study has reported that adolescents smoking waterpipe had significantly lower vascular endothelium growth factor levels which might adversely affect their vascular growth and function [53].

Our results show that both exposure regimens induced a significant increase in the concentration of tissue factor. Also, we observed a statistically significant increase in the concentration of tissue factor in Reg-WPS group compared with Occ-WPS group. This finding corroborates the finding of the experimental study that showed exposure to cigarette smoke increases immunoreactivity for tissue factor in animal model of the atherosclerotic plaques [36]. Moreover, an in vitro study showed the induction of tissue factor expression following the incubation of endothelial cells and smooth muscle cells with nicotine and cotinine [54].

Fibrinolysis is considered the main anticoagulation system that depends on the activity of tPA and plasminogen activator inhibitor-1 which are secreted from endothelial cells [55].
Diminished fibrinolytic activity manifested in low levels of tPA in smokers has also been reported in conditions associated with atherosclerosis [56]. Our data show that only Reg-WPS induced significant decrease in the concentration of tPA compared with either the control or Occ-WPS groups. This result corroborates previous findings which showed dysfunctional fibrinolytic system that was manifested in alterations in tPA release associated with cigarette smoking [57] and reduction in tPA concentration following the incubation of endothelial cells with serum collected from cigarette smokers [45]. Moreover, compared with nonsmokers and cigarette smokers, it has been demonstrated that waterpipe smokers had a significantly higher levels of plasma fibrinogen, a risk factor for cardiovascular disease [58].

The present data show a significant increase in erythrocyte numbers, hematocrit and hemoglobin concentrations in Reg-WPS group. In addition, we found a significant difference in the three aforementioned hematological parameters between Occ-WPS and Reg-WPS groups. These findings are indicative of a bone marrow response, causing an increase in red blood cells. We also found that occasional smoking did not provoke such changes. The latter is consistent with a previous study conducted on rats following WPS exposure which showed a significant increase in red blood cells count, hemoglobin concentrations and hematocrit in rats exposed to WPS [59]. In accordance with our findings, a clinical study showed chronic smoking induces increase in red blood cells count, hemoglobin, hematocrit [60].

We also found a significant increase in plasma LDH activity in Reg-WPS exposed mice compared with either the control or Occ-WPS groups. A clinical study conducted on smokers showed high levels of LDH in the plasma [61]. Moreover, we have demonstrated earlier that LDH is increased following short- and long-term exposure to WPS [42, 62, 63]. High levels of LDH in the plasma are usually associated with cell damage caused by smoking [31].

Our data show that Occ-WPS smoking for a period of 6 month did not cause systemic inflammation as no significant changes in the concentrations of IL-1β and TNFα have been observed. By contrast, chronic exposure to Reg-WPS triggered a significant augmentation in the concentration of both pro-inflammatory cytokines. In addition, there was a statistical significance in the concentration of these cytokines between Occ-WPS and Reg-WPS groups. Our findings are consistent with a previous clinical study that showed a significant increase in the levels IL-6, IL-8, IL-1β and TNFα in human plasma following exposure to CS, WPS and dual WPS and CS [64]. The inflammation in the blood suggests that the use of WPS may be a potential risk factor for developing both chronic respiratory and cardiovascular diseases. Moreover, it has been shown that cigarette smoke media stimulates TNFα release by macrophages in vitro [65]. Likewise, we have shown an increase in the concentrations of pro-inflammatory cytokines in the lung and the heart tissue homogenates following WPS exposure [42, 62].

It is well recognized that cigarette smoke and WPS contain many oxidants [66]. Hence, it is well established that the adverse effects of smoking may result at least partly from oxidative damage to cellular components [66]. Such damage could result from both oxidants present in cigarette smoke and reactive oxygen species generated from activated phagocytic cells [67]. Unlike proinflammatory markers, both Occ-WPS and Reg-WPS induced a significant increase in the levels of 8-isoprostane and total antioxidant in the plasma. These findings substantiate other studies results that demonstrated the occurrence of oxidative stress following repeated passive [22] and conventional cigarette smoking [68]. In addition, it is well-established that WPS triggers systemic oxidative stress [14]. Moreover, clinically the increase in oxidative stress markers such as 8-epi-prostaglandin F2α and malondialdehyde has been reported following regular waterpipe smoking [69].

One of the limitations of the current work is the fact that we studied WPS as a whole and it would be relevant to assess the cardiovascular impact of the different components of WPS including the gas and particulate phases. Moreover, the present work has been carried out on healthy mice and it would be important to conduct comparable work on animal models of human diseases including diabetes, hypertension and asthma.
Conclusion

Taken together, our data confirmed the alteration of vascular homeostasis, systemic inflammation and oxidative stress induced by regular WPS inhalation and it showed, for the first time, that even chronic occasional WPS inhalation caused hypercoagulability, vascular alteration and systemic oxidative stress. The latter findings clearly indicate that more measures of awareness are needed to limit the widely spread misperception that occasional smoking has low risk of smoking-related diseases.

Acknowledgements

Author Contributions

NH contributed to the interpretation of data, drafting and revising the manuscript. SB, NEZ and OE performed the experiments. BHA contributed to the design and the editing of the manuscript. AN designed, planned, supervised all the experiments, analyzed and interpreted the data and edited the manuscript. All authors have approved the final version of the manuscript.

Funding

This work was supported by funds of the Zayed Center for Health Sciences (grants # 12R008 and 12R072) and the College of Medicine and Health Sciences (grant # 12M022) of the United Arab Emirates University and Al-Jalila Foundation (grant # AJF201701).

Statement of Ethics

The project was reviewed and approved by the Institutional Animal Care and Use Committee of the United Arab Emirates University (Approval # ERA_2017_5625) and experiments were performed in accordance with protocols approved by the Institutional Animal Care and Research Advisory Committee.

Disclosure Statement

The authors declare they have no conflict of interests.

References

12 El-Zaatari ZM, Chami HA, Zaatari GS: Health effects associated with waterpipe smoking. Tob Control 2015;24:i31-i43.


