Ambient Fine Particulate Matter Induces Apoptosis of Endothelial Progenitor Cells Through Reactive Oxygen Species Formation

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Key Words
PM • Endothelial progenitor cells • ROS • Apoptosis

Abstract
Background/Aims: Bone marrow (BM)-derived endothelial progenitor cells (EPCs) play a critical role in angiogenesis and vascular repair. Some environmental insults, like fine particulate matter (PM) exposure, significantly impair cardiovascular functions. However, the mechanisms for PM-induced adverse effects on cardiovascular system remain largely unknown. The present research was to study the detrimental effects of PM on EPCs and explore the potential mechanisms. Methods: PM was intranasal-distilled into male C57BL/6 mice for one month. Flow cytometry was used to measure the number of EPCs, apoptosis level of circulating EPCs measured using ELISA. To determine the role of PM-induced ROS in EPC apoptosis, PM was co-administrated with the antioxidant N-acetylcysteine (NAC) in wild type mice or used in a triple transgenic mouse line (TG) with overexpression of antioxidant enzyme network (AON) composed of superoxide dismutase (SOD)1, SOD3, and glutathione peroxidase (Gpx-1) with decreased in vivo ROS production. Results: PM treatment significantly decreased circulating EPC population, promoted apoptosis of EPCs in association with increased ROS production and serum TNF-α and IL-1β levels, which could be effectively reversed by either NAC treatment or overexpression of AON. Conclusion: PM exposure significantly decreased circulating EPCs population due to increased apoptosis via ROS formation in mice.
Introduction

Endothelial dysfunction or injury is considered one of the major factors that contribute to the development of cardiovascular diseases like atherosclerosis and coronary heart disease [1, 2]. Endothelial progenitor cells (EPCs) play a critical role in vascular re-endothelialization, angiogenesis, and prevention of neointima formation after vascular injury [3-5]. Therefore, it is important to maintain the number and function of EPCs to normal level in pathological conditions, such as hyperlipidemia and diabetes.

PM is responsible for 3.2 million deaths per year and 76 million years of healthy life lost [6]. The majority of mortality following PM exposure has been shown to be related to cardiovascular diseases [6]. Different sources of PM contain different components. The composition of PM is a mixture of various particles including metals, crustal material and bioaerosols [7, 8]. It has been reported that PM exposure is able to produce many deleterious effects on cardiovascular system such as vascular dysfunction, reduced heart rate variability and enhanced coagulation-thrombosis risk [7, 9]. Long-term exposure of PM accelerated the mice with high fat diet [10]. Recent studies have found that PM significantly suppressed the number and function of EPCs in both mouse and human [11, 12]. However, the mechanisms for the detrimental effects of PM on EPCs remain largely unknown.

It has been shown that exposure to PM leads to increased production of reactive oxygen species (ROS) and oxidative stress [13]. The present study was designed to test the hypothesis that increased ROS formation could mediate the effect of PM on the population and increased apoptosis of EPCs. To further test the hypothesis that ROS induced by PM was the main cause for EPCs apoptosis, ROS production was blocked by using either antioxidant N-acetylcysteine (NAC) or a transgenic mouse model (TG) with concomitant overexpression of an antioxidant enzyme network (AON) of human copper/zinc superoxide dismutase (SOD)1, extracellular SOD3, and glutathione peroxidase (Gpx-1) with decreased ROS formation. We observed that NAC treatment or AON overexpression could effectively rescue PM-induced EPCs apoptosis. Taken together, our data demonstrated that PM-mediated ROS production was indeed the major cause for increased apoptosis of EPCs, thus, preventing excessive ROS formation might be a potential effective approach to the prevention and treatment of PM-induced cardiovascular disorders.

Materials and Methods

PM exposure and animal model

All the animal experiments were performed in accordance with the Guidelines of the Animal Care Committee of the Ohio State University Medical Center, Columbus, Ohio, USA. The experimental protocols for the present study were reviewed and approved by the Animal Care Committee of the Ohio State University Medical Center. PM-<4μm (Standard Reference Materials 2786) was purchased from The National Institute of Standards and Technology (NIST), which has a mean particle diameter of 2.8 μm (particle-size characteristics in atmospheric particulate material and similar matrices) with the major components including polycyclic aromatic hydrocarbons (PAHs), nitro-substituted PAHs (nitro-PAHs), polybrominated diphenyl ether (PBDE) congeners, hexabromocyclododecane (HBCD) isomers, sugars, polychlorinated dibenzo-p-dioxin (PCDD) and dibenzofuran (PCDF) congeners, and inorganic constituents [14]. PM was dispersed in solution by ultrasonic in endotoxin-free PBS for 30 min at a concentration of 0.5 μg/μl [15]. Each mouse was treated with 10 ug PM three times per week for 1 month via intranasal instillation that has been widely used and validated as described [16-19]. Endotoxin-free PBS was used as control. Wild-type (WT) male C57 BL/6 mice (6-8 weeks old) were purchased from Jackson Lab (Maine, USA). To evaluate the role of ROS formation induced by PM, the mice were pre-treated with NAC (1mg/ml in the drink...
water) for 24 hours prior to PM exposure. To further evaluate the role of ROS production in mediating the effects of PM, a TG mouse model with concomitant overexpression of AON with decreased ROS production (6-8 weeks old, male) were used to repeat the experiment. The generation of TG mouse that have been backcrossed at least 10 generations onto the C57 BL/6 background was detailed previously [20]. The AON enzyme overexpression level and their activities were also determined recently [21], and confirmed in the present study. The littermate WT male C57BL6 mice were used as the control.

**Histological examination of lung sections**

To assess the effects of PM on airway inflammation, lungs were excised and fixed in 10% formalin, washed in methanol, dehydrated, embedded in paraffin, and cut into 5-µm sections. The preparations were then mounted on slides, stained with hematoxylin and eosin, and examined microscopically in a blinded fashion.

**Proinflammatory cytokines measurement**

Mouse blood was collected after 1 month of exposure to PM. The serum was prepared from the blood sample with centrifugation for 20 minutes at 300g. The levels of serum proinflammatory cytokines TNF-α and IL-1β were determined using ELISA kit from Biolegend (San Diego, CA, USA) according to manufacturer’s recommendation.

**Flow cytometry analysis for cell apoptosis, intracellular ROS formation and EPCs**

After exposure of mice to PM or PBS for 1 month, mouse blood was collected and the red blood cells (RBC) were eliminated with RBC-lysis buffer as described [22]. The blood cell apoptotic rate was determined with FACS using the apoptosis kit from BD Pharmingen (CA, USA). The early apoptotic cells were defined as Annexin V FITC positive cells, while the late apoptotic cells was defined as Annexin V FITC and propidium iodide (PI) double positive cells as described [23]. The level of intracellular ROS formation in blood cells was determined using the ROS Detection Reagents-FITC (Invitrogen) as described [22]. The cells were incubated with the reagent for 10 min at 37°C. The labeled cells were washed twice with PBS and then suspended in warm PBS for analysis using flow cytometry. The fluorescence-positive cells were quantitatively evaluated using an LSRII (BD Bioscience, CA, USA) at the wavelength of 525 nm as described [22]. For the circulating EPC analysis, the blood CD34+/CD133+ cell population was determined using flow cytometry as described [24] with a total of 50,000 cells used for each test. The antibody CD34+ AF700 was obtained from Becton Dickinson Biosciences (N), USA, and CD133+ PE was from Biolegend (San Diego, CA, USA).

**Statistical Analysis**

All the data were presented as means ± standard deviation (SD), and statistically analyzed using unpaired Student t-test (two-sided) for two groups of data or one way ANOVA (analysis of variance) (PRISM Version 4.0; GraphPad Software, Inc., San Diego, CA) followed by post hoc conservative Tukey's test for three or more groups of data to minimize type I error as appropriate. The differences were considered statistically significant when a two-tailed p < 0.05.

**Results**

**PM treatment decreased circulating EPC number in association with increased apoptosis**

Blood cells were collected for EPC analysis after PM exposure. Flow cytometry analysis showed that PM exposure significantly decreased the population of CD34+/CD133+ cell by 50% as compared to the control group (Fig. 1A). In search for the reason for EPC suppression by PM, the blood cell apoptotic rate was determined. As shown in Fig. 1B, the early apoptosis rate of circulating EPCs was significantly increased up to 17% compared to the rate of 9% in control group. In addition, the late apoptosis rate was also substantially increased up to 3 folds over the control group (Fig. 1C).

**PM treatment increased the serum levels of TNF-α and IL-1β and induced inflammation infiltration in the lung**

TNF-α and IL-1β are closely related to cell apoptosis [25, 26] and upregulated by PM exposure [27]. To explore the mechanisms for increased EPC apoptosis induced by PM,
Fig. 1. PM treatment decreased murine circulating EPC level with increased EPC apoptosis. A. After exposure of C57BL/6 mice with PM or PBS through intranasal distillation for 1 month, the blood cells were collected after red blood cell lysis and stained with CD34-AF700 and CD133-PE antibody for EPC (CD34+/CD133+) measurement. The EPC level was significantly decreased in C57BL/6 mice with PM exposure compared to the PBS control. The annexin V (B) and PI (C) were used to incubate the blood cells for apoptosis analysis. Both early (B) and late (C) apoptotic rate of blood cells in the mice with PM exposure was significantly increased as compared to the PBS control group. WT+PBS: C57BL/6 mice with PBS treatment; WT+ PM: C57BL/6 mice with PM exposure. * WT+ PM vs WT+PBS, P<0.01, n=8.

Fig. 2. PM increased the levels of serum proinflammatory cytokines and induced lung inflammation infiltration. The murine serum was prepared for proinflammatory cytokine TNF-α and IL-1β measurements, and lung was collected for inflammation infiltration analysis after exposure in C57BL/6 mice with PM or PBS for 1 month. There was a significant increase in both TNF-α (A) and IL-1β (B) in the serum in the mice with PM exposure compared to the PBS control. The inflammation level was significantly increased in murine lung with PM exposure. WT+PBS: C57BL/6 mice with PBS treatment; WT+ PM: C57BL/6 mice with PM exposure. * WT+ PM vs WT+PBS, P<0.01, n=8.

serum levels of TNF-α and IL-1β were measured. The serum levels of the inflammatory factors TNF-α and IL-1β were significantly elevated to 9 pg/ml ± 1.8 pg/ml and 156.9 pm/ml ± 37.8 pg/ml in the mice with PM exposure compared to 4.3 pm/ml ± 1.1 pg/ml and 69.8 pg/ml ± 25.1 pg/ml in the mice with PBS treatment, respectively (Fig. 2A and B). To determine if the increased serum levels of TNF-α and IL-1β were associated with tissue inflammation, we examined the inflammation infiltration in the lung. Indeed, after exposure with PM for 1 month, the inflammation infiltration of mouse lung was induced compared to the PBS control (Fig. 2C).
PM increased intracellular ROS production in circulating EPCs

ROS production could induce proinflammatory cytokine secretion [28]. Thus, we hypothesized that PM-induced proinflammatory cytokine secretion might be through ROS production. We observed that intracellular ROS level was indeed significantly increased in the circulating EPCs in the mice with PM exposure (Fig. 3).

NAC treatment or AON overexpression attenuated the detrimental effects of PM on EPCs

To determine whether ROS was the cause for EPC apoptosis induced by PM, both pharmacological and transgenic approaches were employed to block ROS generation. When the WT-mice were co-treated with PM and NAC, ROS formation was effectively inhibited. We also used a transgenic (TG) mouse model that over-expressed AON with reduced ROS production. We observed that blood intracellular ROS production induced by PM exposure was effectively blocked in NAC-treated mice and in the TG mice overexpressing the AON (Fig. 4A).

To determine whether the increased serum proinflammatory cytokine level and inflammation infiltration in the lung in the mice exposed to PM were mediated by ROS, the lung tissue inflammation infiltration and the serum TNF-α and IL-1β levels were evaluated in NAC-treated mice and in TG mice. Consistent with our hypothesis, the TNF-α and IL-1β levels were reversed to the normal level in the mice exposed to PM and treated with NAC or over-expressing AON compared to their controls (Fig. 4B and 4C). In addition, murine lung inflammation infiltration level was significantly decreased in TG mice or NAC-treated WT mice (Fig. 4D). These data suggested that increased serum proinflammatory cytokine level and lung inflammation infiltration in the mice exposed to PM were indeed mediated by increased ROS formation.

Next, we determined whether NAC treatment or over-expressing AON could prevent PM-induced EPC apoptosis. As expected, both early and late apoptotic rate (17% for early apoptotic and 14% for late apoptotic rate after PM treatment) of EPCs were significantly reversed in either NAC treated mice or TG mice following PM exposure (Fig. 4E and F).

Finally, we examined whether inhibition of ROS production could maintain EPC population during PM exposure. As shown in Fig. 4G, decreased EPC population by PM was completely reversed by NAC treatment or AON overexpression. Thus, our data suggested that ROS induced by PM exposure was indeed a major cause for decreased population of circulating EPCs due to increased apoptosis.
Fig. 4. NAC treatment or AON overexpression decreased ROS production and reversed the detrimental effects of PM. (A) Increased blood intracellular ROS production was effectively blocked by NAC treatment or AON overexpression in the mice exposed with PM. The murine lung inflammation was much lower in TG mouse and WT mouse with NAC treatment than the WT mouse with PM exposure. Increased level of serum proinflammatory cytokine TNF-α (B) and IL-1β (C) as well as lung inflammation infiltration (D) was significantly decreased by NAC treatment or AON overexpression in the mice exposed with PM. Increased early apoptotic (E) and late apoptotic (F) rate were effectively reversed by NAC treatment or AON overexpression in the mice exposed with PM. (G) Decreased circulating EPC level was completely restored by NAC treatment or AON overexpression in the mice exposed with PM. WT+PBS: C57BL/6 mice with PBS treatment; WT+PM: C57BL/6 mice with PM exposure; WT+PM+NAC: C57BL/6 mice with PM exposure and NAC treatment; TG+PM: TG mouse with PM exposure. * WT+PM vs WT+PBS or WT+PM+NAC or TG+PM, P<0.01, n=8.
Discussion

In the present study, we demonstrated that PM exposure significantly decreased the circulating EPC population through proinflammatory cytokine TNF-α and IL-1β mediated apoptosis. We further demonstrated that treating the mice with antioxidant NAC or overexpression of AON significantly decreased the intracellular ROS level in circulating EPCs, reduced serum TNF-α and IL-1β level, diminished EPC apoptotic rate, and restored EPC population in the mice with PM exposure. To our knowledge, this was the first time to report that PM-induced ROS production was responsible for the impaired number of EPCs through ROS-TNF-α/IL-1β-apoptotic pathway as shown in Fig. 5.

Air pollution or second hand smoke (SHS) has significant impact on the number and function of EPCs. Some studies have shown that SHS, PM and other pollutants such as nickel and acrolein are able to decrease the number and function of circulating EPCs [11, 29]. The potential mechanisms are believed to be related to impaired VEGF-mediated signaling that compromised the mobilization of bone marrow-derived EPCs to the circulation [30] or blocking nitric oxide production [31]. On the other hand, SHS has been shown to increase EPC level through increased VEGF and MCP-1 level [32]. Some clinical studies demonstrated, that brief inhalation of coarse PM [12] or SHS [31] could increase the number, but decrease the function of EPCs. However, O’Toole et al. reported that exposure of higher concentration of PM (for 24 hours) could lead to a reduction in circulating EPCs in young adults [33]. Further studies are required to dissect the detailed molecular mechanisms to clarify the conflicting observations.

ROS and oxidative stress are involved in EPC dysfunction in many disease states including hyperlipidemia, diabetes and coronary artery disease [3-5]. Particles especially PM widely exist in the environment and may carry ROS within gas phase [13] or water phase (aerosol) [34] into the lower respiratory tract to create an increased risk on health. There is growing evidence for oxidative stress in response to air pollution in different organs [35]. ROS could function as signaling molecules in PM-triggered autophagy in human epithelia A549 cells [36]. Oxidative stress could be triggered by PM and result in alterations in mitochondrial gene expression in brown adipose tissue [37]. Clinical studies indicated that ROS-initiated oxidative stress and inflammation by PM was closely related to the pediatric asthma [38]. Tonne et al. described a relationship between ambient PM_{2.5} oxidative burden and carotid intima-media thickness (a measure of subclinical atherosclerosis) [39]. PM extracts could deplete antioxidants from a simulated respiratory tract lining fluid model which contained three major water soluble antioxidants (glutathione, urate, and ascorbate) at physiological
concentrations that served as the first line defense in the airway against the oxidative activity of PM [40]. In addition, Strak et al. delineated that PM- and PM_{10}-related oxidative burden did not have an important impact on acute changes in exhaled NO or lung function, while a close relationship was present for ultrafine particles and NO2/NOx [41]. These observations were consistent with our hypotheses that ROS production might be a major cause of decreased number of EPCs following PM exposure.

The role of inflammatory cytokine-induced ROS production in apoptosis has been established [42]. ROS could also directly induce inflammatory cytokine expression/production via regulation of immune signaling through TLR4-mediated NF-κB activation [28]. It was reported that the Mn^{2+}-mediated induction of inflammatory cytokine production was associated with increased production of H$_2$O$_2$, and effectively attenuated by the H$_2$O$_2$-scavenger dithiothreitol, and partially prevented by inhibitors of NF-κB and p38MAP kinase [43]. ROS has been shown to induce gene expression of inflammatory mediators, such as IL-1 and TNF-α [44]. Inflammatory cytokines have significant effect on nearly all cell types including EPCs through their receptors (which are ubiquitously present on nearly all cell types), and p75-related TNF receptor that is predominantly expressed in lymphoid cells as well as other hematopoietic and endothelial lineage cells like EPCs [45]. In the present study, we demonstrated that PM exposure induced ROS production that increased TNF-α and IL-1β level, thus leading to increased apoptosis of EPCs.

Antioxidant enzyme and antioxidant supplementation have been examined for its impact on cardio-respiratory effects of PM exposure. Animal studies have shown an increase in the levels of antioxidant gene expression in epithelial cells after exposure to diesel exhaust particles [46]. Romieu et al. [47] reported that omega-3 polyunsaturated fatty acid blunted the adverse effect of PM on heart rate variability. Additionally, Tashakkor et al. demonstrated that antioxidant supplementation such as Vitamin C and E had multiple beneficial and protective effects against the damage of different air pollution on human lung [48]. Antioxidant probucol and vitamins were able to rescue cigarette smoke-dependent impairment of ischemia-induced neovascularization via improvement of EPC function [49]. However, to date, no study has linked the beneficial effects of antioxidant treatments to EPC survival. The present study illustrated that either antioxidant NAC or overexpression of AON was able to reverse the adverse effect of PM on EPCs. These therapeutic strategy to treat cardiovascular diseases associated with air pollution.

Of course, there are lots of questions that need to be addressed on PM-induced structural and functional impairment on EPCs. For example, does PM also affect bone marrow stem cells which are the main source of EPCs? Does PM affect the proliferation of EPCs in addition to induce apoptosis? Does PM also trigger autophagy of EPCs? Does PM induce other inflammatory factors to alter the EPC population? How does ROS regulate the production and secretion of TNF-α and IL-1β? What are the source of TNF-α and IL-1β following PM treatment? All these questions require further studies.

In conclusion, we reported that PM exposure significantly decreased circulating EPC population through enhanced ROS-TNF-α and IL-1β-apoptosis pathway, and the antioxidant NAC or overexpression of AON was able to reverse the adverse effect of PM on EPCs. These findings suggested that preventing excessive ROS production could constitute a novel therapeutic strategy to treat cardiovascular diseases associated with air pollution.

**Disclosure Statement**

The authors declare no conflict of interest.

**Acknowledgements**

This work was supported by a US NIH grants NIH R01 HL094650 (to ZL), RO1ES018900 (QS), and an American Heart Association grant to HZ (AHA 12SDG12070174)
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