Potential role of polycyclic aromatic hydrocarbons as mediators of cardiovascular effects from combustion particles

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Abstract

Air pollution is the most important environmental risk factor for disease and premature death, and exposure to combustion particles from vehicles is a major contributor. Human epidemiological studies combined with experimental studies strongly suggest that exposure to combustion particles may enhance the risk of cardiovascular disease (CVD), including atherosclerosis, hypertension, thrombosis and myocardial infarction. In this review we hypothesize that adhered organic chemicals like polycyclic aromatic hydrocarbons (PAHs), contribute to development or exacerbation of CVD from combustion particles exposure. We summarize present knowledge from existing human epidemiological and clinical studies as well as experimental studies in animals and relevant in vitro studies. The available evidence suggests that organic compounds attached to these particles are significant triggers of CVD. Furthermore, their effects seem to be mediated at least in part by the aryl hydrocarbon receptor (AhR). The mechanisms include AhR-induced changes in gene expression as well as formation of reactive oxygen species (ROS) and/or reactive electrophilic metabolites. This is in accordance with a role of PAHs, as they seem to be the major chemical group on combustion particles, which bind AhR and/or is metabolically activated by CYP-enzymes. In some experimental models however, it seems as PAHs may induce an inflammatory atherosclerotic plaque phenotype irrespective of DNA- and/or AhR-ligand binding properties. Thus, various components and several signalling mechanisms/pathways are likely involved in CVD induced by combustion particles. We still need to expand our knowledge about the role of PAHs in CVD and in particular the relative importance of the different PAH species. This warrants further studies as enhanced knowledge on this issue may amend risk assessment of CVD caused by combustion particles and selection of efficient measures to reduce the health effects of particular matters (PM).

Keywords: Air pollution, Combustion particles, Polycyclic aromatic hydrocarbons, Cardiovascular disease, Atherosclerosis

Background

According to the World Health Organization (WHO) air pollution is the preponderant environmental risk factor, being responsible for about one in every nine deaths globally [1]. Exposure to particulate matter with an aerodynamic diameter of 2.5 μm and less (PM2.5) has been found to have vascular effects leading to ischemia, myocardial infarction, stroke and other cardiovascular diseases (CVD) [2–4].

Cardiovascular health consequences of air pollution are generally equal to or exceed those due to pulmonary diseases [3, 5]. As is the case for lung cancer, it is no apparent threshold for adverse cardiovascular effects due to PM2.5 in the dose range humans are exposed [6]. The aim of this review was to highlight the hazard potential of polycyclic aromatic hydrocarbons (PAHs) as mediators of PM-induced CVD, as this has received limited attention by particle toxicologists.

Particulate matter and polycyclic aromatic hydrocarbons in ambient air

A number of factors affects PM toxicity, including size, shape, structure, surface reactivity, bio-persistence and...
presence of soluble components (Table 1) [7, 8]. There are large regional differences in composition of PM depending on sources [9]. Toxicological studies have identified several transition metals, organic carbon species, semi-quinones, and endotoxins as specific PM-related components with potential to induce oxidative stress and inflammation [3]. Combustion engines, in particular diesel engines are major contributors to PM$_{2.5}$ in urban environments. Thus, combustion particles such as diesel exhaust particles (DEP) are frequently used to explore mechanisms of PM-induced CVD [7, 10–12]. Combustion particles consist of carbon cores in the ultrafine PM size-range (< 100 nm) with complex mixtures of organic chemicals adhered to the surface [13, 14]. Composition and amount of organic chemicals present in DEP vary, dependent on fuel burned, temperature, engine load, drive-cycles and type of combustion technology. Average levels of organic chemicals in DEP generally range from 20 to 40% of total mass, but may reach as much as 90% [15, 16]. PAHs are the most well-known of these chemicals [17]. Other known chemical groups include n-alkanes, hopanes and steranes [18].

PAHs are produced by incomplete combustion of organic materials such as coal and fossil fuels, cigarette smoking and various industrial activities [19]. The major sources for the global total atmospheric emission of PAH16 have been estimated to be residential/commercial biomass burning (60.5%), open-field biomass burning (agricultural waste burning, deforestation, wildfires (13.6%), and petroleum consumption by motor vehicles (12.8%) [20]. The majority of airborne PAHs with low vapor pressure is adsorbed to PM [17, 21]. PAHs containing five or more aromatic rings are mainly found bound to PM, while PAHs containing four or less aromatic rings seem predominately to occur in the gas phase. However, three and four-ring PAHs such as phenanthrene and pyrene are so abundant in outdoor air, that they also tend to be the most abundant PAHs bound to DEP and other combustion particles [17]. As temperature and vapor pressure is connected, a considerably larger proportion of airborne PAHs will be bound to PM during winter, while a relatively larger fraction will be in the gas phase during summer. In line with this, winter PM$_{2.5}$ from Milan (Italy) was found to contain 10-fold higher PAH content (% of PM mass) compared to summer PM$_{2.5}$ [22].

Possible mechanisms linking PM to CVD

WHO has estimated that approximately 75% of deaths attributable to ambient air pollution are due to stroke or ischemic heart disease [1]. PM$_{2.5}$ exposure is associated with endothelial dysfunction in CVD–risk groups [23, 24], but recent findings indicate that environmental exposure to PM$_{2.5}$ may cause endothelial injury even in young healthy adults [25]. Furthermore, it has been suggested that air pollution may cause hypertension, and increase in long-term PM$_{2.5}$ exposure as low as 3 $\mu$g/m$^3$ has been associated with vascular dysfunction [26, 27]. Double-blinded cross-over exposures have also revealed that diesel exhaust increases systolic blood pressure in healthy participants [28].

Combustion particles may contribute to development of CVD via several mechanisms (Fig. 1 and Table 2). Exposure of pulmonary macrophages and epithelial cells may cause oxidative stress, further triggering release of pro-inflammatory mediators into the circulation. These mediators have potential to harm endothelial cells and cause systemic effects [25, 29]. PM$_{2.5}$/DEP may affect platelets and coagulation, increasing the risk of vascular clotting [30–32]. It has also been suggested that inhaled diesel exhaust may trigger receptors in the autonomic nervous system of the respiratory tract and thus affect cardiac control [33, 34]. In addition, constituents of PM$_{2.5}$/DEP may have more direct cardiovascular effects [11, 35, 36]. Recently, inhaled gold nanoparticles were found to accumulate at sites of vascular inflammation in mice and humans [37]. However, only a small amount of gold nano-particles (less than 0.3%) reach the circulation [38]. By contrast, it has been shown that when combustion particles deposit in the alveolar region the majority of their available PAH-load may rapidly detach from the particles, and is transferred across the epithelial barrier and diffuses into the bloodstream in an un-metabolized state [17, 39, 40].

Due to the complex composition of PM$_{2.5}$, there is no single causative chemical, chemical group or component behind the various cardiovascular effects [3, 41, 42]. However, even though particle cores sometimes may be

| PM surface: charge, reactive groups /«stable radicals» / redox-active metals |
|-----------------------------|
| PM inorganic chemicals: redox-active metals |
| PM organic chemicals: ROS and reactive metabolites via metabolism of PAH within cells |

Table 1 Combustion particle properties linked to redox activity
involved, biologic effects of combustion particles seem largely dependent on organic chemicals. Notably, animal studies have shown that DEP denuded of organic chemicals lost their potential to induce atherosclerosis [43]. Furthermore, experimental studies in vitro have illustrated that some effects of PM2.5/DEP relevant for CVD, are linked to extractable chemicals from these particles [44–48]. Thus, as PM2.5/DEP contains substantial amounts of organic chemicals, their vascular effects may presumably be linked to these chemicals [11, 14, 35, 37].

**Inflammation and atherosclerosis**

Atherosclerosis may lead to myocardial infarction, cerebrovascular and peripheral vascular disease, making it the major cause of deaths due to CVD [49, 50]. It is an inflammatory disorder of the arteries, initiated by dysfunction of endothelial cells, which precede the histopathological changes. The process involves oxidative stress and leads to increased levels of local inflammatory mediators including cytokines, chemokines and adhesion molecules that lead to extravasation of monocytes. These monocytes accumulate oxidized low-density lipoproteins (oxLDL) and develop into foam cells and deteriorate, leading to atheroma. Several mediators among others matrix metalloproteinases (MMPs) destabilize atherosclerotic plaques ultimately causing rupture and thus infarction [51].

Inflammation in endothelial cells and/or the lung is considered a central link between ambient PM-exposure and CVD [52]. Inflammatory reactions may be directly caused by PM-induced chemokine/cytokine release as well as indirectly through PM-induced cytotoxicity [53, 54]. Oxidative stress is central in both processes [54–56]. Reactive oxygen species (ROS) can be generated directly by particles and particle components or more indirectly through various metabolic and inflammatory processes (Tables 1 and 2) [57, 58].

After exposing healthy men to DEP, Törnqvist and co-workers observed impairment of endothelium-dependent vasodilatation suggested to be due to early systemic oxidative stress [59]. Animal experiments have shown that DEP exposure increases size and complexity of lesions in atherosclerotic mice [60]. In an Apo E−/− mice model, DEP caused marked effects on buildup of plaques in arterial walls, while DEP denuded of organic chemicals was without effect [43], indeed supporting an important role of these chemicals in atherosclerotic effects of DEP. That DEP may aggravate development and progression of atherosclerosis is further supported by in vitro studies. In a co-culture model, wood smoke particles and DEP increased adhesion of monocytes to endothelial cells [61], which is often linked to enhanced migration of inflammatory cells from the bloodstream. DEP has been
shown to impair endothelial function [62, 63], increase formation of lipid-loaded foam cells from macrophages [64], and trigger inflammatory reactions in endothelial cells [48].

**Aryl hydrocarbon receptor**

The aryl hydrocarbon receptor (AhR), plays a central role in regulating toxicity of PAHs and other environmental pollutants such as dioxins and co-planar polychlorinated biphenyls [65, 66]. In its classical mode of action, ligand-activated AhR dimerizes with the AhR nuclear translocator (ARNT) and binds to so-called xenobiotic response elements (XREs) in promotor regions of target genes such as cytochrome P450 (CYP) enzymes CYP1A1/CYP1B1 (Table 2). Metabolism of PAH from DEP by various CYP-enzymes may form ROS and reactive electrophilic metabolites with potential to trigger inflammation [67, 68]. Furthermore, it has now become clear that a number of pro-inflammatory genes are directly regulated by the AhR [69–71], and at least some of these such as interleukin (IL)-1β and IL-8 (CXCL8) contain xenobiotic response elements (XREs) in their promotor region [72, 73]. AhR may also mediate inflammatory signals via non-classical pathways; this includes cross-talk with the nuclear factor-κB (NF-κB) family of transcription factors as well as other transcription factors and signaling molecules, independent of ARNT activation [74–76]. In addition to its transcriptional role, AhR is also believed to mediate toxic effects via non-genomic signals including increases in intracellular concentration of calcium [Ca$^{2+}$] [77, 78].

AhR is important for cellular functions. Increasing evidence suggests that AhR plays a central role in development and maintenance of the cardiovascular system, and that xenobiotics may affect homeostasis and trigger CVD-pathogenesis by modulating biological responses of critical cell types through activation of AhR [79–84]. Knockdown of AhR results in cardiac hypertrophy and specific AhR-knock-down in vascular endothelial cells cause hypotension [85, 86]. Furthermore, overexpression of AhR has been shown to induce endothelial dysfunction [87]. AhR expression and polymorphisms were also associated with risk of coronary arterial disease in a Chinese population [88]. Compared with controls, blood

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**Table 2** Initial molecular effects of combustion particle/PAH-Parent compound, reactive oxygen species (ROS) and electrophilic metabolites

| Extracellular effects (blood) | HDL to dysfunctional HDL |
|-------------------------------|--------------------------|
| **Cellular effects**          |                          |
| Altered membrane structure and function |                          |
| Membrane receptors, transporters and channels |                          |
| Ca$^{2+}$ signaling |                          |
| ROS signaling                  |                          |
| Phosphorylation signaling (including inhibition of phosphatases) |                          |
| **Intracellular receptors including aryl hydrocarbon receptor (AhR)** |                          |
| Canonical genomic: CYP enzymes |                          |
| Non-canonical genomic: Cross talk with e.g. NF-κB |                          |
| Non-genomic signaling: |                          |
| protein phosphorylation via kinases (ScC) |                          |
| membrane order /caveola |                          |
| Ca$^{2+}$ signaling via SOCE/ROCE |                          |
| **Cellular induced ROS formation/oxidatitv stress** |                          |
| Redox cycling of eg PAHs metabolites |                          |
| Activation of membrane bound oxidases |                          |
| Damage to mitochondria |                          |
| Activation of ROS defense (macrophages) |                          |
| **Genotoxic effects** |                          |
| DNA damage and mutation |                          |
levels of AhR were found to be significantly increased in patients with coronary arterial disease [88]. In line with this, DEP-exposure has been reported to induce cardiac dysfunction and remodeling (left ventricular dilation) through an AhR-dependent mechanism [89]. Furthermore, the prototypical environmental AhR ligand, 3,4,7,8-tetrachlorodibenzop-p-dioxin (TCDD) has been reported to induce cardiomyopathies, cardiac lesions, arteritis, and atherosclerosis in rodents, and increase the risk of CVD in humans [83]. Recently it was also shown that TCDD inhibits cardiomyocyte differentiation from human embryonic stem cells via AhR-regulated mechanisms [90].

**Calcium signaling**
The cytosolic concentration of calcium \([\text{Ca}^{2+}]\) is central to pathophysiological processes including AhR-genomic signaling, oxidative stress and inflammation [91, 92]. In endothelial cells \([\text{Ca}^{2+}]\), regulates blood pressure and flow, especially through control of vascular smooth muscle cells via myo-endothelial micro-domains and eNOS [93–96]. Furthermore, \([\text{Ca}^{2+}]\), is involved in regulation of endothelial permeability, a central step in the pathogenesis of atherosclerosis [97, 98].

Activation of \([\text{Ca}^{2+}]\)-channels in the plasma membrane such as transient receptor potential (TRP) channels, results in \([\text{Ca}^{2+}]\)-influx [99]. Notably, a number of studies suggest that combustion particles including DEP and wood smoke particles, and chemicals attached may trigger health effects by affecting \([\text{Ca}^{2+}]\) flux through TRP-channels [100, 101]. Some of the TRP-channels appear to be activated through direct interaction with particles or attached chemicals, while others seem to be activated by more indirect mechanisms such as transactivation. Importantly, several TRP-channels are central to endothelial homeostasis, and seem to play a role in development of CVD, especially by affecting endothelial function [102–104].

\([\text{Ca}^{2+}]\), is also regulated via \([\text{Ca}^{2+}]\)-release from intracellular stores such as the endoplasmic reticulum or mitochondria. This may result from activating \(\mathrm{G}\) protein-coupled receptors (GPCRs) or receptor tyrosine kinases (RTKs) [105, 106]. \(\beta_1\) and \(\beta_2\)-adrenergic receptors (ADRs) regulate cardiopulmonary function and immune responses, and are among the main drug-targets in CVD treatment [107–109]. Certain PAHs known to be present in DEP may increase \([\text{Ca}^{2+}]\), in human micro-vascular endothelial cells (HMEC-1) and other cell types via \(\beta_2\)ADRs [110–112]. In human bronchial epithelial BEAS-2B cells exposed to 1-nitropyrene (1-NP), \(\beta_2\)ADRs appeared to be involved in \([\text{Ca}^{2+}]\)-increase and induction of the pro-inflammatory cytokine CXCL8 [111].

Transporters, channels and receptors cluster in membrane micro domains [113], and their activity may also be affected by their presence inside or outside such ordered domains [114, 115]. Several xenobiotics including DEP-extracts and PAHs have been found to affect membrane microstructure, thus possibly affecting \([\text{Ca}^{2+}]\), or other signaling mechanisms by altering the membrane physiology [116–119].

**Search strategy and review structure**
As a starting point the following search terms were used in PubMed: (((“Cardiovascular Diseases”[Mesh]) OR “Blood Pressure”[Mesh]) AND (((“Air Pollutants”[-Mesh]) OR “Air Pollution”[Mesh]) OR “Environmental Exposure”[Mesh]) OR “Inhalation Exposure/adverse effects”[Mesh]) AND “Polycyclic Aromatic Hydrocarbons”[Mesh]) (29.5.2018). Using this approach 121 studies were found. Only 12 of these studies were linked to general population when excluding studies on health effects of cancer therapy (eg. with anthracyclines) and occupation. Thus, we additionally included occupational studies of environmental setting to the papers reviewed. Studies of PAH at high non-environmental settings (e.g. coke oven workers) were also commented as they were regarded to present relevant information.

Given the difficulty of identifying relevant animal and in vitro mechanistic studies linking PAH to CVD from other literature, additional strategies were also used. A number of searches were performed in PubMed using combinations PAH or specific PAH and terms linked to CVD including endothelial dysfunction, foam cells and cardiovascular development. Some papers were identified by tracking the citation network (cited and citing papers) of identified papers, while some were from the authors personal databases.

Publications identified were screened at abstract level. A total of 19 epidemiological studies exploring cardiovascular effects of exposure to environmental levels of PAHs and CVD were included. No formal analysis of these studies was however undertaken.

With regard to available animal and mechanistic research, we highlight research suggesting that extractable organic material of combustion particles, PAHs and AhR and intracellular calcium could be linked to cellular processes central in development and exacerbation of CVD. Concentrations or exposure routes used in experimental studies with pure PAH-exposure were not evaluated. Information from these studies were included to explore possible mechanisms involved and added as proof of principle.

**The role of organic chemicals and PAH in mediating CVD**

**Human exposure and epidemiological studies**
Exposure to PM2.5/DEP has been found to cause dysfunction of cells and biological processes of the cardiovascular system linked to CVD, including atherosclerosis, hypertension,
myocardial infarction, stroke, thrombosis and restricted valve motion (Table 3) [3, 4]. Furthermore, accumulating evidence suggests that PM/DEP with the highest portion of organic chemicals have the greatest effects on vascular outcomes [2, 11, 35, 120, 121]. A recent review reported that most epidemiological studies found significant positive association between PAHs exposure and manifest CVD, as well as major risk factors predisposing for CVD including elevated blood pressure [122].

Importantly, we are not only exposed to PAHs through polluted air. As reviewed elsewhere tobacco smoke and foods are among the major sources in addition to occupational exposures [21]. The relative contribution of PAHs from air pollution versus other sources with regard to CVD will depend on the location, activity and dietary habits of the population in study. However, the majority of PAHs absorbed via the gastro-intestinal tract will go through first-path metabolism and elimination in the liver. By contrast, it has been shown by Gerde et al. [40] that inhaled B[a]P taken up through the alveolar region primarily enters the circulation, reaching the heart and vasculature in an un-metabolized state. Thus, the importance of air pollution as a source for circulatory levels of parent PAHs should not be underestimated.

Urinary 1-hydroxypyrene, a metabolite of pyrene, is among the most commonly used biomarkers. Although 1-hydroxypyrene concentrations are correlated to smoking, certain PAH-rich food items and occupational exposure studies have shown that there is a statistically significant correlation between urinary 1-hydroxypyrene concentrations and ambient air levels of pyrene and benzo[a]pyrene (B[a]P), in subjects that smoke less than 20 cigarettes daily [21]. Thus, it has been argued that 1-hydroxypyrene is a valid biomarker also of PAH exposure from ambient air.

Heart disease and mortality rates
PM and PAH exposures may occur in occupational settings at levels 1–3 orders of magnitude higher than those in environmental settings [123]. Notably, heart disease mortality rates in occupational cohorts such as aluminum smelters are typically lower than those in the general population [124, 125], likely due to the “healthy worker effect” bias which has been suggested to be strong for diseases of the cardiovascular system [126]. The relation between exposure to PAH and mortality from ischemic heart disease (IHD; 418 cases) was studied in a cohort of 12,367 male asphalt workers from various nations. Both cumulative and average exposure indices for B[a]P were positively associated with mortality, and demonstrated a consistent exposure–response relation for this association [127]. Recent morbidity studies among aluminum smelters have reported associations of adverse cardiovascular effects with PM and PAH exposure, by using biomarkers of CVD, such as markers of inflammation, blood pressure, and heart rate variability. Ischemic heart disease mortality was associated with B[a]P in the highest exposure category. A monotonic, but non-significant trend was observed between chronic B[a]P exposure and acute myocardial infarction. When follow-up was restricted to active employment, hazard ratio for ischemic heart disease was 2.39 in the highest cumulative B[a]P category. The stronger associations observed during employment suggests that risk may not persist after exposure cessation [128].

In a cohort of autoworkers, modest evidence that occupational exposure to PM3.5 containing PAHs may increase risk of ischemic heart disease mortality was reported [129]. In a population-based case-reference study of myocardial infarction and occupational exposure to motor exhaust and other combustion products, relative risk of myocardial infarction was 2.11 among highly exposed and 1.42 among those intermediately exposed to combustion products from organic material. Furthermore, exposure-response patterns in terms of both maximum exposure intensity and cumulative dose, were found [130].

Exposure to traffic increased the risk of myocardial infarction in susceptible subjects [131]. Increased onset of chest pain was observed immediately and 6 h after traffic

| Blood clotting and thrombosis |
| Blood pressure increased |
| **Endothelial dysfunction** increased; adhesion molecules, disrupted calcium homeostasis and inflammatory reactions |
| Inflammation local and systemic increased |
| Development of foam cells increased |
| Cardiovascular development dysregulated |

Table 3 Effects of combustion particles/PAHs on processes linked to CVD
exposure. Exacerbated heart symptoms among myocardial infarction survivors have been linked to particle-associated organic compounds [132]. The study showed an association between PM, adhered organic chemicals and daily symptoms.

The association between particle-associated PAH concentrations and symptom-severity among myocardial infarction survivors, suggests a major influence on cardiovascular health [133]. In the general population, cross-sectional studies have reported that concentration of urinary mono-hydroxy metabolites of phenanthrene, was significantly associated with self-reported CVD [19]. Subjects with middle and highest tertile of fluorene and phenanthrene metabolites had a similar significantly higher prevalence of peripheral arterial disease as compared to subjects within the lowest tertile [134]. A Chinese multi-provincial cohort study found an association between the 10-year risk of atherosclerotic cardiovascular disease and PAHs exposure measured as urinary OH-PAH metabolites, more specifically: 2-hydroxyfluorene, 9-hydroxyfluorene, 1-hydroxyphenanthrene, and ΣOH-PAHs levels [135]. Increased serum C-reactive protein (CRP) has also been reported from subjects with elevated levels of urinary PAHs, indicating inflammation [136].

Occupational PAH exposure among boilermakers [137, 138] and coke oven workers [137] has been associated with altered heart rate variability. Recently, an association between background PAH exposure and heart rate variability in the general population was also reported. Increased urinary OH-PAH metabolites and Framingham risk scores were dose-dependently related to decreased heart rate variability [139].

**Inflammatory markers**

Urinary 1-hydroxypyrene was increased among taxi-drivers compared to non-occupationally exposed subjects, and positively correlated with pro-inflammatory cytokines (IL-1β, IL-6, IL-10, TNF-α, IFN-γ and hs-CRP) and biomarkers of oxidative damage (serum levels of oxidized LDL, auto-antibodies and homocysteine), but negatively correlated with antioxidants. As higher levels of inflammatory biomarkers and homocysteine, represent important predictors for cardiovascular events, these data suggest a possible link between PAH exposures and CVD in an occupational setting [140]. In support of this, a study comparing taxi drivers with and without co-morbidity found that urinary 1-hydroxypyrene levels were associated with carotid intima-media thickness and with serum homocysteine levels [141]. Occupational exposure to soot, rich in PAH, has also been associated with increased risk of CVD. In a cross-sectional study of chimney sweepers, it was found that chimney sweepers had up to 7-fold higher concentrations of PAH metabolites in urine than controls. Furthermore, early markers of CVD, namely homocysteine, cholesterol, and reduced levels of high density lipoproteins (HDL), were affected in serum from chimney sweepers. PAH metabolites correlated positively with the percentage of time spent soot sweeping. A study of school-age children identified increases in oxidative stress biomarkers in association with urinary PAHs [138].

**Blood pressure**

Coke oven workers are highly exposed to PAHs, and studies have associated coke oven emission to hypertension and abnormal electrocardiography [142–144]. Furthermore, the levels of 2-hydroxyphenanthrene, 3-hydroxybenzo[a]pyrene, and 3-hydroxy-benzo[a]anthracene were positively associated with diastolic blood pressure in chimney sweepers [145]. Contrasting findings were reported in a study of blood pressure changes and exposure to PAHs in outdoor workers. The results showed that in the total group, levels of urinary 1-hydroxypyrene were negatively associated with systolic and diastolic blood pressure [146]. The authors suggest that occupational exposure to PAHs may significantly influence blood pressure, probably acting via the autonomic nervous system [146].

In a population of elderly, the oxy-PAHs, chrysene-5,6-dione and B[α]P-3,6-dione were significantly associated with increased systolic blood pressure and pulse pressure [147]. PAHs were also associated with hypertension in a study among relatively young individuals, mostly women [148]. An association between PAHs in housewives’ hair and hypertension has been investigated [149]. Seven PAHs in hair samples were measured with detection rate > 70%. Only acenaphthylene was found to be associated with an increased risk of hypertension. Among children in Saudi Arabia, proximity to an oil refinery was associated with pre-hypertension and increased PAH and PM exposure. However, urinary PAH metabolites (1-hydroxypyrene and hydroxyphenanthrene) were not associated with cardiovascular outcomes in the study [150]. In a repeated measures study of 106 residents in China, eight urinary PAHs metabolites (OH-PAHs) were analyzed. Association of OH-PAHs with high blood pressure and increased risk for atherosclerotic CVD were found, and was suggested to be partially mediated by obesity [151].

**Embryonic development**

Passive diffusion seems to control trans-placental transfer of PAHs from maternal serum to the fetal circulation [152]. There is evidence from experimental systems that exposure to PAHs results in congenital heart defects [153], as discussed later. However, results from a large population-based study do not support an association between potential maternal occupational exposure to PAHs and various congenital heart defects [154].
**Evaluation of epidemiological studies**

Overall, the epidemiological data suggest a possible association between PAH-exposure and CVD, although some inconsistencies have been reported amongst others for hypertension. Furthermore, the number of available studies is limited, and the role of PAHs can be difficult to disentangle from other combustion-related pollutants, especially in non-occupational cohort studies. It should also be emphasized that no formal analysis of the strengths of these epidemiological studies was performed in this review. Thus, conclusions should not be drawn based on these epidemiological data alone.

**Experimental studies in animals and mechanistic considerations**

In vivo animal studies addressing cardiovascular effects of PM were recently summarized by Møller and co-workers [155]. Some challenges regarding extrapolation from animal studies to humans were discussed. Atherosclerosis is not as common in rodents as in man, e.g. wildtype mice do not develop atherosclerosis; even in the often used atherosclerosis-prone ApoE−/− (knock-down) mouse model, plaques formed do not rupture. In humans plaque rupture is a pivotal event in acute myocardial infarction and stroke [155].

**Effects of DEP on CVD considering content of organic chemicals**

As animal models seem to be less sensitive, quantitative consideration based on these studies are less relevant. However, they do give important knowledge with regard to possible mechanisms involved. Most studies on the widely used automobile-derived DEP sample A-DEP (50% organic chemicals) reported effects on either atheroma development or vasomotor function, as reviewed by Møller et al. [155]. ApoE−/− mice exposed to SRM 1650 (20% organic chemicals), had increased plaques progression [156]. Studies with SRM 2975, which has a low content of organic chemicals (5%) [157, 158], are somewhat contradictory and less convincing. In Wistar rats exposed to SRM 2975 by intra-tracheal instillation, vasomotor function was unaffected although the rats had extensive lung inflammation [159]. However, SRM 2975 disturbed vasomotor function of hypertensive rats [160]. Furthermore, SRM 2975 has been reported to increase both lung inflammation as well as plaque formation in ApoE−/− mice [60].

Several studies on DEP with various content of organic chemicals have suggested that they are important triggers of effects on CVD. Compared to PM2.5, ultrafine particles contained more than twice the amount of organic chemicals, and induced significantly more proatherogenic effects in vivo [35]. Keebaugh and coworkers have possibly conducted the most compelling in vivo evidence that organic chemicals attached to combustion particles is important for atherosclerosis development [43]. ApoE−/− mice exposed to concentrated ambient particles (CAPs) had accelerated atherosclerosis development and reduced cardiac function, while these responses in mice exposed to CAPs denuded of organic chemicals were not statistically different from mice exposed to air.

**Effects on heart**

PAHs such as B[a]P are known to bind to and activate the AhR, leading to increased expression of xenobiotic metabolic enzymes (XME) such as CYP1A1, CYP1A2, CYP1B1, NAD(P)H:quinone oxidoreductase-1 (NQO1), and glutathione S-transferase A1 (GSTA1) [68], enhanced production of ROS and reactive PAH species leading to lipid peroxidation and tissue damage. Notably, studies from our lab and others, suggest that AhR-activation and induction of CYP1-expression may be the most sensitive endpoint in DEP- and PM-exposed cells [161, 162]. AhR and AhR-regulated xenobiotic metabolizing enzymes appear to be highly expressed in the cardiovascular system. The various AhR-regulated genes are differentially expressed in different parts. Particularly high levels are found in the endothelium of aorta, coronary arteries and ventricles [163]. Disruption of synthesis and/or metabolism of endogenous substances, such as arachidonic acid (AA), prostaglandins (PGs), and thyroid hormones has been suggested to contribute to the pathogenesis of CVD [164]. Studies in rats and chicken embryos have demonstrated that CYP1A1 mediates metabolism of arachidonic acid to hydroxyeicosatetraenoic acid (HETE), epoxyeicosatrienoic acid (EET), and Prostaglandin E2 metabolites [165, 166]. These endogenous substances may thus be targets for PAH-mediated cardiotoxicity via AhR-induced metabolism. While cardiac AhR-regulated CYPs are involved in CVD pathogenesis; the AhR-regulated enzymes NQO1 and GST are regarded as more cardio-protective. An imbalance in expression of cardio-toxic and cardio-protective xenobiotic metabolizing enzymes has thus been suggested as a main determinant of PAH-mediated cardiotoxicity [163]. However, as AhR also appear to play a central endogenous role in development and homeostasis of the cardiovascular system, impact of PAHs on CVD is likely not restricted to regulation of xenobiotic metabolism.

Experimental studies support the epidemiological findings indicating that exposure to AhR agonists in the environment may lead to cardiovascular toxicity. AhR agonists such as 3-methylcholanthrene (3-MC) and B[a]P increased the heart to body weight ratio as well as levels of hypertrophic markers, such as atrial natriuretic peptide and brain natriuretic peptide in rats [167]. In contrast, treatment with B[c]P, an isomer of B[a]P that is a
poor AhR-ligand, did not induce cardiac hypertrophy, supporting a role of AhR in the pathogenesis of cardiac hypertrophy. However, phenanthrene, which is also generally considered a poor activator of AhR-mediated CYP-1 expression, was reported to induce cardiac hypertrophy through a mechanism involving reduced miR-133 expression [168]. In the same study, effects on cardiomyocyte size and protein content in vitro, was observed at low nanomolar concentrations of phenanthrene (0.05–0.5 nM). Additionally, other AhR-ligands [163, 169] have been reported to cause cardio-toxic effects in humans and experimental animals. Exposure to B[a]P, induced atherosclerosis to a greater extent in mice with high-affinity AhR (B6) and had a huge impact on gene expression of the aorta when compared to mice with low affinity AhR (B6.D2) [170]. This study also indicated an endogenous role of AhR signaling in regulating heart size.

Genome-wide association studies have identified susceptibility loci and candidate genetic variants that predispose to coronary artery disease in humans [171]. Such studies have identified variation at 6q23.2 to be associated with this disease in Caucasian and Han Chinese populations [172], and identified TCF21 as the causal gene in this locus [173]. Recent findings suggest that TCF21 and AhR cooperate to activate a pro-inflammatory gene expression program in coronary artery smooth muscle cells to activate an inflammatory gene expression program that is exacerbated by TCDD or PAHs, and may contribute to the overall risk for coronary arterial disease [71]. Furthermore, the study showed that AhR gene expression was increased in atherosclerotic lesions in mice, using laser capture microdissection. The AhR protein was localized in human carotid atherosclerotic lesions, where it was associated with protein kinas with a critical role in innate immune responses.

**Oxidative stress, inflammation and atherosclerosis**

There are studies suggesting that PAHs like B[a]P aggravate atherosclerosis via increased oxidative stress [174]. Oxidative stress in the vascular endothelium will decrease availability of NO, a key regulator of vascular tone and blood pressure. Overexpression of antioxidant enzymes in ApoE-deficient mice suppressed B[a]P-accelerated atherosclerosis [6]. Some studies find that B[a]P and other AhR agonists increase expression of pro-inflammatory cytokines such as CXCL8 and TNF-α, which in turn cause infiltration of macrophages and neutrophils into the lungs [73]. Pulmonary oxidative stress and inflammation may result in “systemic spill-over” suggested to be important for development of CVD [3].

Oxidative stress may also lead to alteration in circulating lipids. Elevations of serum cholesterol-associated triglyceride and LDL are regarded as primary causative factors for the development of atherosclerosis. These pro-atherogenic molecules diffuse into sub-endothelial cells and cause further vascular dysfunction [2]. Microarray studies after TCDD exposure in rodents revealed AhR involvement in regulation of fatty acid and cholesterol synthesis [175]. In humans, industrial exposure to dioxin has been associated with lipid metabolism disruption and high levels of circulating cholesterol and triglyceride [176].

Atherosclerosis induced by PAHs has similarly been linked to AhR-dependent effects on cholesterol synthesis [177]. Furthermore, B[a]P-exposed ApoE−/− mice show enhanced monocyte chemoattractant protein-1 (MCP-1) gene expression and marked increase of macrophage content in atherosclerotic plaques [178]. Furthermore, in vitro studies have found that B[a]P-induced MCP-1 gene and protein expression was inhibited by AhR antagonism, but not antioxidant treatment, suggesting AhR-mediated induction [179]. B[a]P also enhanced TNFα-induced expression of MCP-1, thus supporting the concept that induction of inflammation is a crucial process in PAH-enhanced atherogenesis.

**DNA damage and atherosclerosis**

Environmental chemical carcinogens including PAHs have also been suggested as risk factor for atherosclerosis as a result of their mutagenic effects [180, 181]. Atherosclerosis is associated with DNA damage in both circulating and vessel-wall cells. Furthermore, DNA adducts derived from exposure to environmental mutagens are abundant in atherosclerotic vessels, and PAH-DNA damage has been detected in human endothelial and smooth vascular muscle cells [182]. It has been suggested that PAHs could cause aberrant smooth muscle cell proliferation, a focal point in the genesis and progression of atherosclerosis [183–185]. As seen in smooth muscle cells isolated from B[a]P-treated quail, serial sub-culture of smooth muscle cells exposed to B[a]P in vitro yielded a fast-growing population of cells after the fifth passage [183]. The pathogenic relevance of mutation-related molecular damage in atherosclerosis has only partial support in experimental animal models. Both the number and size of arterial lesions in the brachiocephalic arteries in atherosclerosis-susceptible (“White Carneau”) and atherosclerosis-resistant (“Show Racer”) females pigeons were significantly enhanced after long-term dosing with B[a]P [186]. Exposure to B[e]P did not enhance lesion development, and no effects were observed in males. On the other hand, chronic exposure to B[a]P only induced larger and phenotypically different atherosclerotic plaques in ApoE-knockout mice while their location or number seemed unaffected [187]. Furthermore, morphometrical analysis show that B[e]P causes a similar increase in plaque size. Thus, in some models it seems that PAHs may induce
an inflammatory atherosclerotic plaque phenotype irrespective of DNA- and/or AhR-binding properties [178].

**Endothelial dysfunction**

Endothelial dysfunction is a key initiating event in numerous CVD including atherosclerosis. Endothelial cells have the highest induction of CYP1A1 in the vasculature of rat models, and these cells have ability to metabolize B[a]P [188]. The vascular endothelium is susceptible to damage, as it is in constant contact with circulating xenobiotics including PAHs. Accordingly, B[a]P-DNA adducts have been found in the endothelium from human atherosclerotic lesions [182].

In vitro studies indicate that DEP may impair endothelial function [62]. Dysfunction of the endothelium is marked by increased adhesiveness caused by the presentation of cellular adhesion molecules, such as intercellular adhesion molecule 1 (ICAM-1) and vascular cell adhesion molecule-1 (VCAM-1). These are essential initiating events of atherosclerosis, which result in retention of macrophages and monocytes in the sub-endothelial space. DEP have been shown to increase endothelial cell permeability through down-regulation of tight junction proteins, decreased trans-endothelial resistance and redistribution of vascular endothelial cadherin from cell membrane intracellularly [189]. Data from in vitro studies on human endothelial cells demonstrate that B[a]P is able to increase ICAM-1 through a caveolae- and AhR-mediated pathway, thereby increasing monocyte adhesion [190].

Endothelial dysfunction is also characterized by disruption of calcium homeostasis [93, 95] and increased expression of pro-inflammatory markers [191]. Interestingly, in a human exposure study, the calcium blocker verapamil lost its vasodilatory effect after exposure to DEP, indicating effects on [Ca²⁺]i regulation [192]. In a recent in vitro study, DEP applied on the epithelial side of an alveolar 3D tri-culture, rapidly induced pro-inflammatory and AhR-regulated genes in basolateral endothelial cells [48]. Further analysis of endothelial cells showed that lipophilic extracts of DEP containing most of the PAHs, modified calcium homeostasis and increased expression of pro-inflammatory markers via an AhR-dependent pathway [48, 116]. The authors hypothesized that PAHs detached from DEP translocate through alveolar epithelial cells and modify calcium homeostasis and increase inflammatory reactions, thereby triggering endothelial dysfunction.

In human endothelial HMEC-1 cells, B[a]P triggered a relatively rapid and transient increase of [Ca²⁺]i, which was prevented by co-treatment with β2ADR inhibitors, anti-β2ADR antibodies, or siRNA-mediated knockdown of β2ADR expression [112]. B[a]P was shown, to bind β2ADR directly with high affinity (Kd = 10 nM), as assessed by in vitro binding assays and molecular modeling [112]. This interaction also appear to cause a desensitization of β2ADR signaling, leading to reduced responsiveness towards epinephrine [193]. In vitro studies on lung epithelial cells suggest that 1-nitropyrene may induce Ca²⁺-signaling at least partly through activation of β2ADR, and that both β2ADR- and Ca²⁺-signaling may be involved in CXCL8 up-regulation [111]. However, the 1-nitropyrene concentration required to activate β2ADR-induced Ca²⁺-signaling, appeared to be an order of magnitude higher than B[a]P [111]. Other G protein-coupled receptors (GPCRs) have also been implicated in DEP-induced Ca²⁺-signaling and inflammation. Li et al. [100] found that organic chemicals extracted from DEP increased [Ca²⁺]i, via the GPCR, pro tease activated receptor 2 (PAR-2), in primary human bronchial cells. PARs are central for regulating vascular function, and activation may promote conversion of endothelial cells into a pro-inflammatory phenotype in conditions associated with endothelial dysfunction [194]. Recently it was reported that both induction of inflammation-associated genes and [Ca²⁺]i, partly depended on PAR-2 in endothelial cells exposed to lipophilic organic chemicals from DEP [48, 195].

Pyrene and B[e]P, two PAHs normally considered to have low affinity for the AhR, also induced marked [Ca²⁺]i induction in the HMEC-1 endothelial cell line. Especially pyrene appeared to be a potent inducer of Ca²⁺-signaling, triggering approximately 2-fold higher [Ca²⁺]i responses compared to B[a]P or chrysene [112]. Surprisingly, pyrene-induced [Ca²⁺]i seemed to depend on a selective activation of non-genomic AhR-signaling, occurring in absence of effects on CYP1A1 expression [119]. This differs distinctly from B[a]P-induced Ca²⁺-signaling through β2ADR [112], and suggest that PAHs may affect [Ca²⁺]i-regulation in endothelial cell through different mechanisms. Notably, studies on effect of lipophilic organic chemicals extracted from DEP, suggest that their induction of [Ca²⁺]i, could involve both AhR- and β2ADR-dependent responses [119, 195].

Phenanthraquinone, another PAH-derivative found in DEP, has a potent inhibitory effect on eNOS activity [196]. TCDD was also reported to suppress eNOS, but induced iNOS [197]. Moreover, increased eNOS activity and enhanced vessel elasticity was observed in AhR-deficient mice, while AhR overexpression ex vivo suppressed the migratory capacity of endothelial cells, impaired eNOS activation and suppressed NO production [198]. Furthermore, AhR was reported to be involved in cigarette smoke-induced increases in carotid intimal thickening in mice, through activation of iNOS expression in smooth muscle cells [199]. Notably, eNOS levels are reduced and iNOS levels increased during endothelial aging and dysfunction [200]. Thus, it has been speculated that...
increased AhR expression could result in vascular inflammation through induction of iNOS and thereby further contribute to endothelial dysfunction and vessel aging [198].

**Effects on blood pressure**

Exposure to PAHs in experimental animals has produced equivocal effects on hypertension. 3-MC-induced hypertension was found to be associated with eNOS inactivation [201]. NO produced by endothelial cells is a key regulator of vascular tone and blood pressure. Thus, an inactivation of eNOS may cause suppression of NO-mediated vaso-relaxation and elevation of blood pressure. In another study, intranasal B[a]P was found to alter circadian blood pressure patterns and cause lung inflammation [202]. Atherosclerosis and hypertension are risk factors for development of abdominal aortic aneurysms. Thus, it is interesting to note that B[a]P aggravates development of abdominal aortic aneurysms in ApoE-knockout mice [203]. Furthermore, ischemia-induced angiogenesis by B[a]P was found to depend on AhR [204].

**Development of foam cells**

Air pollution particles increased formation of lipid loaded foam cells from macrophages in vitro [64]. The formation of foam cells in sub-endothelial lesions play a key role in atherogenesis, in part by releasing pro-inflammatory cytokines. Most interestingly, in vitro studies with the human macrophage cell line U937 showed that organic extracts from ultrafine particles (UFP) and DEP as well as TCDD induced more expression of CXCL8, TNF-α, and COX-2 mRNA, than stripped particles, whereas total particles (parent and stripped particles) led to a greater production of C-reactive protein and IL-6 mRNA [64]. UFP-, DEP- and organic extract-induced expressions of COX-2 and CYP1A1 were suppressed by AhR antagonist, indicating that these effects are mainly mediated by organic components which can activate the AhR. In contrast, induction of C-reactive protein and IL-6 seem to be particle-related effects that are AhR-independent. The inflammatory responses induced by PM and their respective organic extracts were also associated with a subsequent increase of cholesterol accumulation. B[a]P has been found to cause similar lipid accumulation in primary human macrophages obtained by GM-CSF-mediated differentiation of monocytes [205]. The effect was linked to an AhR-dependent repression of Niemann–Pick type C1 protein (NPC1). Interestingly, NPC1 is a transmembrane protein containing a sterol-sensitive domain that participates in cholesterol trafficking from the late endosome/lysosome to the plasma membrane, and is therefore thought to protect cells from intracellular accumulation of cholesterol [206].

Taken together, these data illustrate that vascular inflammatory responses as well as foam cell-formation caused by PM, are tightly linked to organic chemicals triggering AhR responses. Even though foam cell formation caused by PM is likely triggered by several components, PAHs seem to represent possible candidates.

**Embryonic development**

Several studies indicate that in utero exposure to PAHs dysregulate cardiovascular development. Following in utero B[a]P exposure of Long Evans Hooded rats, systolic blood pressure was significantly elevated in offspring on postnatal day P53 in the middle and high exposure groups as compared to controls [207]. A number of studies have also investigated effects of PM and PAHs on cardiovascular development in zebrafish embryos. PM-induced disruption of cardiovascular development in zebrafish embryos has been suggested in part to be due to AhR-mediated suppression of Wnt-signaling, and has been attributed to PAHs [208–210]. Defects in cardiac function preceded characteristic morphological abnormalities induced in zebrafish embryos exposed to a mixture of PAHs [153]. Most interestingly, the relative toxicity of different mixtures was directly proportional to the amount of phenanthrene, or dibenzothiophene-phenanthrene total in the mixture, apparently due to direct effects on cardiac conduction.

Pyrene induced a different syndrome of anemia, peripheral vascular defects, and neuronal cell death in zebrafish, similar to effects previously described for potent AhR ligands [153]. Embryonic exposure of zebrafish to pyrene, has also been shown to disrupt normal cardiac development resulting in heart abnormalities, such as pericardial edema and cardiac looping defects, and alter expression of defective cardiac differentiation-related genes [211]. Furthermore, in zebrafish embryos exposed to phenanthrene it has also been observed abnormally looped and enlarged hearts along with increased expression of matrix metalloproteinase-9 (MMP-9) and transforming growth factor [212]. Low level co-exposure with silica nanoparticles and B[a]P induced inflammatory response and hyper-coagulable conditions in zebrafish embryos [213]. Thus it appears that different PAH species may have distinct and specific effects on early cardiovascular development in fish. It seems that these effects are due to mechanisms that differ from classical AhR-ligand binding and/or formation of electrophiles.

Although existence of high-affinity physiological activators of AhR are still debated, also endogenous signaling of the AhR is believed to be important in the development and function of cardiovascular system, based on studies with AhR gene-deficient mice [83]. These studies show that AhR-knockout mice develop cardiac hypertrophy, abnormal vascular structure in
multiple organs and altered blood pressure depending on their host environment and point to a role for genetic variations of this gene.

**Conclusion**

Human epidemiological studies combined with experimental studies strongly suggest that exposure to combustion particles enhance risk of CVD, including atherosclerosis, hypertension, thrombosis and myocardial infarction. Overall, epidemiological studies both in occupational settings and the general population suggest possible associations between environmental PAH exposure and CVD including well known CVD-risk factors. However, it should be noted that the literature is limited and some inconsistencies have been reported.

Animal models seem less sensitive than epidemiological studies, but combined with in vitro experimental models they increase our understanding of possible biological mechanisms through which exposure to PM may cause adverse effects linked to CVD. Experiments suggest that organic compounds attached to combustion particles are of importance for triggering CVD, and furthermore that their effects are mediated at least in part by AhR. This is in accordance with a role of PAHs, a well-known group of chemicals present on combustion particles which bind to AhR and/or are metabolically activated by CYP-enzymes.

A number of cardiovascular effects of TCDD and AhR knock-down or overexpression are now well documented, highlighting the central role of AhR in CVD. Specific studies with PAHs show that also B[a]P is cardio-toxic, and increased the heart to body weight ratio as well as levels of hypertrophy markers via AhR-dependent mechanisms. Likewise, certain PAHs may produce equivocal effects on hypertension in experimental animals. Furthermore, PAHs accelerate development of atherosclerosis, induce major changes in gene expression depending on AhR, and B[a]P DNA adducts are found in atherosclerotic lesions.

There are studies that support mechanisms of cardiovascular toxicity involving AhR, ROS and/or reactive electrophilic metabolites. On the other hand, it seems as PAHs in some models may induce an inflammatory atherosclerotic plaque phenotype irrespective of their DNA- and/or AhR-ligand binding properties. Importantly, cardiovascular effects of PAHs may not be restricted to B[a]P, but has also been reported for pyrene, phenanthrene and B[e]P. Additionally, animal knock-out studies clearly link AhR as such to CVD, pointing to the possibility that exposure to PAH may disturb AhR-regulated gene-expression linked to endogenous ligands important for a well-functioning cardiovascular system.

There is still a need to expand our knowledge on the role of PM-composition for development and/or exacerbation of CVD. The key drivers of cardiovascular effects observed in combustion-PM exposed populations still remains to be clearly identified. Nevertheless, mechanistic studies in animals and cell models suggest that PAHs adhered to combustion particles may be among the critical determinants in CVD. This notion seem to be supported by epidemiological studies. Several uncertainties regarding the suggested mechanisms involved and importance of different PAH species remain to be elucidated, and studies assessing the association between PAHs and CVD in the general population remains scarce. This warrants further studies as enhanced knowledge may have implication for risk assessment of combustion particles and their associated PAHs.

**Abbreviations**

[Ca^{2+}]: Cytosolic concentration of calcium; ADRs: Adrenergic receptors; AhR: Aryl hydrocarbon receptor; ARNT: AhR nuclear translocator; B[a]P: Benzo[a]pyrene; CVD: Cardiovascular diseases; CYP: Cytochrome P450; DEP: Diesel exhaust particles; HDL: High density lipoproteins; NF-κB: Nuclear factor-κB; oxLDL: Oxidized low density lipoproteins; PAHs: Polycyclic aromatic hydrocarbons; PM: Particular matter; ROS: Reactive oxygen species; TRP: Transient receptor potential; WHO: World Health Organization

**Acknowledgements**

A summary of this review will be included in a suggested WHO document initiated by the Task Force on Health (TFH) working group on Polycyclic Aromatic Hydrocarbons (PAHs) at the TFH meeting in May 2017. The first PAH-literature search was done by Swiss Tropical and Public Health Institute, Basel, Switzerland. The objective of the working group on PAHs is to “Evaluate the current knowledge on the health risk of polycyclic aromatic hydrocarbons and identify critical gaps” linked to air pollution.

**Authors’ contributions**

JAH and BCB drafted various parts of the first version of the manuscript and wrote the final version in collaboration with MR, ML and JØ. All authors read, commented and approved the final manuscript.

**Funding**

The work was supported by the Research Council of Norway, through the Environmental Exposures and Health Outcomes- and Better Health programs (grants no. 228143 and 260381).

**Availability of data and materials**

“Not applicable”.

**Ethics approval and consent to participate**

“Not applicable”.

**Consent for publication**

“Not applicable”.

**Competing interests**

The authors declare that they have no competing interests.

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