Impact of polycyclic aromatic hydrocarbon exposure on cognitive function and neurodegeneration in humans: A systematic review and meta-analysis

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Introduction: This article documents an emerging body of evidence concerning the neurological effect of polycyclic aromatic hydrocarbon (PAH) exposure with regard to cognitive function and increased risk of neurodegeneration.

Methods: Two electronic databases, PubMed and Web of Science, were systematically searched.

Results: The 37/428 studies selected included outcomes measuring cognitive function, neurobehavioral symptoms of impaired cognition, and pathologies associated with neurodegeneration from pre-natal (21/37 studies), childhood (14/37 studies), and adult (8/37 studies) PAH exposure. Sufficient evidence was found surrounding pre-natal exposure negatively impacting child intelligence, mental development, average overall development, verbal IQ, and memory; externalizing, internalizing, anxious, and depressed behaviors; and behavioral development and child attentiveness. Evidence concerning exposure during childhood and as an adult was scarce and highly heterogeneous; however, the presence of neurodegenerative biomarkers and increased concentrations of cryptic “self” antigens in serum and cerebrospinal fluid samples suggest a higher risk of neurodegenerative disease. Associations with lowered cognitive ability and impaired attentiveness were found in children and memory disturbances, specifically auditory memory, verbal learning, and general memory in adults.

Discussion: Although evidence is not yet conclusive and further research is needed, the studies included supported the hypothesis that PAH exposure negatively impacts cognitive function and increases the risk of neurodegeneration in humans, and recommends considering the introduction of a variable “rural vs. urban” as covariate for adjusting analyses, where the neurological functions affected (as result of our review) are outcome variables.

Keywords: polycyclic aromatic hydrocarbon (PAH), cognition, neurological, neurodegeneration, neurobehavioral, meta-analysis, systematic literature search
Introduction

Exposure to air pollution in the environment is now recognized globally by governments, leading research scientists, and civil society as one of the greatest public health hazards of the 21st century (1). Legislation such as "The UK National Air Quality Strategy" (2), and the European Commission’s "Fourth Daughter Directive" (3) have introduced standards to monitor and limit levels of air pollutants posing the greatest risk to human health. Polycyclic aromatic hydrocarbons (PAHs) are a group of pollutants included in such legislation. PAHs are atmospheric organic compounds composed of two or more benzene rings arranged in a variety of different configurations. PAH compounds also include functional derivatives of the PAHs only containing carbon and hydrogen atoms (e.g., nitro-PAHs) and the heterocyclic analogs (e.g., aza-arenes) (4). Over 100 different PAHs were already identified by the beginning of the 21st century (4), and now the list exceeds 300, with an exact number still to be determined, as those studied are mainly selected based on the instrumentation available to each research group and reference standards (5). They are discharged from anthropogenic sources (Supplementary Figure 1), involving the incomplete combustion and pyrolysis of hydrocarbons, predominantly found in coal, oil, wood, and petrol. PAHs exist in the atmosphere in a gaseous state or are adsorbed to particulate matter. Over 80% of particulate-bound PAHs are associated with particulate matter of an aerodynamic diameter ≤ 2.5 μm (PM$_{2.5}$) (6). However, a large number have been also identified in tobacco smoke (5). The study of PAHs and their impact on health has been compounded by their ubiquitousness and the numerous and widespread sources in which they can be found, some of which are also affected by other air pollutants. As PAHs are rather present as part of complex mixtures in air, water, soil, and food, their identification and characterization, for studying their effect on human health, is challenging (5).

Research surrounding PAH exposure and acute short-term health effects in humans has, thus, far focused on vulnerable individuals with pre-existing health conditions: thrombotic effects in individuals with pre-existing coronary heart disease and impaired lung function in asthma sufferers (7). Chronic long-term exposure has implicated PAH’s reactive metabolites as having the ability to bind to proteins and DNA and exert carcinogenic effects (8). Such biochemical disruption and cellular damage have been most extensively researched in occupational studies, whereby high exposure has been associated with increased incidence of lung, bladder, skin, and gastrointestinal cancer (8–11). Additionally, decreased immune function, developing cataracts, and having kidney or liver damage, including jaundice, have also been associated with high exposure (5, 12). Whilst extensive research exists surrounding PAH's genotoxic and carcinogenic properties, an emerging body of evidence concerns PAH's neurotoxic effect through the induction of oxidative stress, inflammation (13), and vascular injury within the brain (14). Recently, research has emerged associating PAH exposure with impaired cognitive function and increased risk of neurodegeneration.

To the best of our knowledge, from the large body of literature on the influence of air pollution on human health, the implications of PAH exposure specifically, on cognitive function and neurodegeneration in humans, have not been systematically reviewed. Prior reviews have addressed the implications of PAH exposure on general health (15, 16) and its carcinogenic outcomes (17, 18). The reviews which have made cognitive function and neurodegeneration the outcome of interest include exposures to a vast mixture of air pollutants (19–22). Therefore, we aim to disentangle the unique neurotoxic effect of PAHs in specific age groups and cognitive-related functions to provide evidence for cognitive research and more vigilant monitoring and tighter restrictions on the main sources of emission, tailored to each age group, given the differential factors affecting the various stages of brain development. The Department for Environment, Food and Rural Affairs currently considers annual monitoring of concentrations of one PAH, benzo(a)pyrene (BaP), to be a sufficient representation of all atmospheric PAHs, and classifies the potential effect on human health of PAHs collectively, as six compounds, categorized as probably or possibly carcinogenic. No mention is made of the adverse neurological impact (2). A possible explanation is the consideration of concentration levels that constitute a risk for cancer, below which the effect

Abbreviations: AD, Alzheimer’s disease; ADHD, Attention deficit hyperactivity disorder; APOE4, Apolipoprotein E4; Aβ1–42, Amyloid beta protein fragment 1–42; α-synuclein, Alpha-synuclein; B[a]P, Benzo[a]pyrene; BDNF, Brain-derived neurotrophic factor; CEBB, Cerebellar antigen; CO, Carbon monoxide; CSF, Cerebrospinal fluid; ETS, Environmental tobacco smoke; IFN γ, Interferon gamma; IgA, Immunoglobulin A; IgG, Immunoglobulin G; IgM, Immunoglobulin M; IL β, Interleukin beta; IL 2, Interleukin 2; IL 6, Interleukin 6; IL 10, Interleukin 10; IQ, Intelligence quotient; MBP, Myelin basic protein; MBP IgA, Myelin basic protein immunoglobulin A; MBP IgG, Myelin basic protein immunoglobulin G; MCP-1, Monocyte chemoattractant protein-1; MOG IgG, Myelin oligodendrocyte glycoprotein immunoglobulin G; MOG IgM, Myelin oligodendrocyte glycoprotein immunoglobulin M; NHANES, National Health and Nutrition Examination Survey; Non-p-tau, Non-phosphorylated tau; NOx, Nitrogen oxide species; NO2, Nitrogen dioxide; OZ IgA, Oculudin/zonulin immunoglobulin A; OZ IgG, Oculudin/zonulin immunoglobulin G; PAH, Polycyclic aromatic hydrocarbons; PAH-DNA adducts, Polycyclic aromatic hydrocarbon- DNA adducts; PD, Parkinson’s disease; PM, Particulate matter; PM$_{2.5}$, Particulate matter with an aerodynamic diameter > 2.5 micrometers; PM$_{10}$, Particulate matter with an aerodynamic diameter > 10 micrometers; p-tau, Phosphorylated tau; S-100 IgG, Astrocytic protein immunoglobulin G, S-100 IgM, Astrocytic protein immunoglobulin M; TDP-43, Transactive response DNA binding protein 43; TRAP, Traffic-related air pollution.
of these pollutants can pass inadvertently. The UK national air quality objective for B(a)P is 0.25 ng m$^{-3}$. However, emissions of B(a)P have been increasing since 2008 and have exceeded this limit in multiple locations at multiple time points (23). Atmospheric PAH concentrations are subject to seasonal variation and climate (24), as seen in pollution level charts that are used by studies to stratify exposure. While such stratification may add granularity to the data, it is often unrealistic given urban movement and the effect of different local government policies e.g., transportation. A more robust stratification would be to contrast urban and rural areas, where the pollution levels known to widely differ. Therefore, a further aim is to explore the difference in PAH concentration in rural vs. metropolitan areas and the influence this could have on cognitive function and neurodegeneration to inform further studies.

Methods
Eligibility criteria

This review was conducted in line with the PRISMA guidelines (25). Studies included were observational cohort studies of both male and female humans. Time of exposure was inclusive of the gestational period and stretched throughout life until death. Exposure quantification was limited to studies that measured the level of exposure to ambient PAHs or PM$_{2.5}$ through environmental air sampling or spatiotemporal modeling. Measures of exposure also included concentration of PAH metabolites in urine and dosimetry of PAH-DNA adducts from DNA extracted from white blood cells. Outcomes included involved a formal assessment of cognitive function, neurobehavioral symptoms of impaired cognition, and pathologies associated with neurodegeneration. Reports were limited to published scientific articles written in the English language. No publication dates were imposed. Studies were excluded if they did not fulfill the inclusion criteria, were not in humans, or where PAH exposure was measured as a component of the diet, environmental tobacco smoke (ETS), or traffic related air pollution (TRAP). Exposure through diet and ETS is not an appropriate representation of a major source of atmospheric PAH that can be geographically differential (i.e., in terms of urban vs. rural areas) or influential in both short- and long-term exposure. Moreover, prior research has elucidated contaminating pollutants within TRAP composition detrimentally affecting cognitive function, and the effect of diet-related benzo[a]pyrene, dibenz[a,h]anthracene, and benzo[h]fluorantracene in human health and cognition (e.g., learning and memory functions). The inclusion of such studies would confound results and prevent us from elucidating the specific impact of ambient PAH on cognition.

Information sources

Studies were identified by searching electronic databases PubMed (1984–2021) and Web of Science (1979–2021). Given the environmental changes seen as the consequence of lockdown policies and movement restrictions mainly in the period April 2020 to December 2021, publications that reflected or analyzed the environmental effect of this “abnormal” period were excluded. “Polycyclic aromatic hydrocarbons” in addition to the following search terms: “brain,” “neurological,” “cognitive,” “cognition,” “neurodegenerative,” “neurodegeneration,” “neurodevelopment,” and “neurodevelopmental” were used to identify articles in both databases. Limitations applied to the search included only the fields “Title” and “Abstract” being searched. In Web of Science, the document type “Articles” was applied. In PubMed, an additional limitation of species, “Humans,” was applied. Eligibility assessment was performed independently in an unbiased standardized manner by one reviewer. Ambiguity concerning the inclusion or exclusion of a study was resolved by a second reviewer being consulted and a consensus taken. Initial screening was performed by reviewing the title and abstract, after which, the full manuscript was reviewed.

Data collection process

A data extraction sheet was developed and pilot tested on five randomly selected included studies, before being refined accordingly. One review author extracted data from the included studies, a second was consulted where ambiguity arose surrounding the appropriate data to extract. One author was contacted through email to provide numerical data that had only been presented graphically. Information extracted from studies comprised sample size, sample characteristics, ratio between sexes, mean age, age range, comorbidities, air pollution component, time of exposure, air pollution data acquisition method, and outcome measure.

Risk of bias in individual studies

The risk of bias was assessed in line with the QUADAS$^2$ guidelines (University of Bristol, 2003). To ascertain the risk of bias within each study included, one reviewer working independently extracted the following information: participant inclusion/exclusion criteria explained, participant withdrawals from the study explained, use of/comparison with a control/low

1 QUADAS-2. URL http://www.bristol.ac.uk/population-health-sciences/projects/quadas/quadas-2/.
exposure population, confounding variables identified, appropriate method/analysis to adjust for confounding variables, outcome assessors aware of exposure status of study participant, intermediate or unexpected results explained/reported, and whether or not the methods of the study were reproducible.

Methods of analysis

Studies included were divided into four subgroups depending on the time at which the exposure was measured: prenatal, childhood, adult, and finally childhood + adult. The category childhood + adult included studies of young individuals from a wide age range, some exposed during childhood, and others where exposure extended through to adulthood, not analyzed separately. Subsequently, studies were further divided into categories depending on the outcome measured: cognitive abilities, neurobehavioral development, or neurodegeneration. This subgroup division was conducted to adjust for heterogeneity between studies. The meta-analysis was performed by extracting odds ratios and 95% confidence intervals (CI) for the effect sizes of reported outcomes or calculating them from the parameters and data given in the original publications using the Practical Meta-Analysis Effect Size Calculator by David B. Wilson, from (https://www.campbellcollaboration.org/escalc/html/EffectSizeCalculator-OR-main.php). Results were double-checked using the following online resources: (https://www.gigacalculator.com/calculators/odds-ratio-calculator.php) and effect size converter (https://www.escal.site/). Forest plots were used to visualize differences in effect sizes between studies within the same subgroup.

Results

Study selection

The search of Web of Science and PubMed provided a total of 490 citations. After adjusting for duplicates, 428 remained. Subsequent screening of the title and abstract resulted in a further 289 being discarded. Of the remaining 139, a further 102 were excluded upon further examination of the full manuscript.

FIGURE 1
Flow chart of the search, study inclusion, and subgroup division.
and application of inclusion criteria. One study (26) reported data from a previous study (27). Sixteen were reviews and did not include any primary data, 47 reported outcomes not relevant to cognitive function or neurodegeneration, and in 38 the recorded exposure to PAH was not in keeping with the specified criteria, resulting in a total of 37 studies included in this review. Subgroup division resulted in 21 pre-natal exposure studies, 15 concerning cognitive development outcomes, and 13 on neurobehavioral development, seven included measures of both. From 11 childhood exposure studies, three were on general cognition, five on neurobehavioral development, and three on neurodegeneration. There were also five adult exposure studies all with outcomes of general cognition, and three childhood + adult studies all with measures of neurodegeneration. One study included measured outcomes for both pre-natal and childhood exposure (28) and two studies involved two different study cohorts: one exposed only during childhood and the other included a mix of childhood and adult-exposed subjects (29, 30). Figure 1 depicts the flow chart for study inclusion and subgroup division. The full dataset can be found at the following link: https://doi.org/10.7488/ds/3031.

Study characteristics

The 37 studies involved populations from nine countries (Figure 2). Study population characteristics including sample size and mean age (±SD) are displayed in Figure 3.

Eleven studies included cohorts from the USA. Six of them selected participants from the Columbia Center for Children’s Environmental Health cohort; however, each study selected different subgroups of the population and measured different outcomes (31–36). Two studies selected a subgroup of participants from the National Health and Nutrition Examination Survey 2001–2002 (NHANES) (37), one of which included additional participants from the NHANES 2003–2004 cohort (38). The remaining three studies involved cohorts from the Childhood Autism Risks from Genetics and the Environment Study (28), the Adolescent Brain Cognitive Development Study (39), and the Asthma Coalition on Community, Environment and Social Stress project (40). Eight studies reported results from populations in China. One study involved a Taiyuan population (41) in addition to two selecting different subgroups from the Taiyuan Mother and Child Cohort Study (42, 43). Three involved populations were from Tongliang (27, 44, 45), and the remaining two were from Shanxi province (46) and Qingdao City (47).

Five studies involved populations from Spain. Two involved a subgroup from the Infancia y Medio Ambiente Project (48, 49) and three studies from the Brain Development and Air Pollution Ultrafine Particles in School Children project (50–52).

Four studies reported on populations in Poland, three including participants from the Krakow Study (53–55) and one from the Polish Mother and Child Cohort Study (56). Four studies reported on populations in Mexico. All involved Mexico City residents (57), where two refer to
Characteristics of the study population involved in each study. Circle size is representative of the sample size. Red bars indicate mean age and SD. Four studies were omitted from this analysis due to insufficient data (28, 34, 43, 60).

Exposure assessment

Of the 37 studies included, seven measured exposure through environmental PAH sampling, five by environmental PM$_{2.5}$ sampling, seven by PM$_{2.5}$ spatiotemporal modeling, 10 by concentrations of PAH metabolites in urine, and eight using dosimetry to measure PAH-DNA adducts (Figure 4).

Outcome assessments

Outcomes included 21 different tests measuring cognitive function, nine different tests measuring neurobehavioral symptoms of impaired cognition, and three different measures of pathologies associated with neurodegeneration (Figure 5).

Pre-natal exposure

Association between pre-natal PAH exposure and cognitive abilities in childhood

Children with a high pre-natal PAH exposure were found to have a delay in overall child intelligence [OR = 1.75, 95% CI, 1.11–2.71] (54), mental development [OR = 0.65 (32)], and average overall development (27, 44) (OR = 0.84, 95% CI, 0.52–1.36; OR = 1.85, 95% CI, 1.13–3.01, respectively). Specifically, the greatest negative effects reported were in verbal IQ (OR = 3.45, 95% CI, 0.95–12.49) (53) and language (OR = 5.99, 95% CI, 1.88–19.02) (47). However, the latter could not be confirmed in five out of six studies (27, 42, 44, 45, 56). Two studies analyzed the effect of PAH on general cognitive abilities with contradictory results: one (31) reported a negative effect (OR = 2.89, 95% CI, 1.33–6.25) while another (56) reported no effect. PAH effect on impaired motor development was inconclusive, as confirmed by four studies (27, 42, 44, 45) (OR = 0.95, 95% CI, 0.58–1.53; OR = 1.91, 95% CI, 1.12–2.97; OR = 1.63, 95% CI, 1.00–2.65; OR = 1.82, 95% CI, 3.21–1.03, respectively), whereas three others could not confirm it. No association was found between PAH exposure and developmental motor ability (56).
FIGURE 4
Pie chart representing the proportion of included studies measuring exposure to PAH as a measure of environmental PAH sampling, environmental PM$_{2.5}$ sampling, PM$_{2.5}$ spatiotemporal modeling, concentration of urinary PAH metabolites, and dosimetry of PAH-DNA adducts.

FIGURE 5
Pie chart representing the number of studies using different tests to measure outcomes. Measures include cognitive function, neurobehavioral symptoms of impaired cognition, and pathologies associated with neurodegeneration.

Fine and gross motor abilities (47), and psychomotor abilities (31). Only one study reported the effect of PAH and reduced adaptive development (27) (OR = 1.77, 95% CI, 1.09–2.88) while four out of five studies reported no association with adaptive domains (42, 44, 45, 47). Size effects reported by the studies mentioned are graphically represented in Figure 6 and listed in Table 1.

Association between pre-natal PAH exposure and neurobehavioral development

Children with a high pre-natal PAH exposure were found to exhibit externalizing and internalizing behavioral problems (OR = 2.49, 95% CI, 1.57–3.95; OR = 2.39, 95% CI, 1.51–3.79, respectively) (55), and infants exhibited a decrease in behavioral development (OR = 2, 95% CI, 1.27–3.15) (43).
Associations with anxious/depressed behavior were found in three out of four studies (33, 34, 55) (OR = 8.89, 95% CI, 1.7–46.51; OR = 8.14, 95% CI, 1.21–54.94; OR = 1.7, 95% CI, 1.08–2.68, respectively), with no association found by one study (36). Three out of five studies reported a negative effect on children’s attentiveness (33, 35, 36) (OR = 1.34, 95% CI, 0.85–1.83; OR = 2.02, 95%, 1.35–3.03; OR = 3.79, 95% CI, 1.14–12.66) whilst two (34, 55) reported no effect. The report from one study (55) about the effect of both withdrawn/depressed and aggressive behavior (OR = 2, 95% CI, 1.27–3.16; OR = 2.29, 95% CI, 1.45–3.62, respectively) was contradicted by another study (36) that reported no effect for either. The latter (36) did, however, report the effect of impaired thought problems (OR = 1.95, 95% CI, 1.3–2.91) which was contradicted by the former (55). Only one out of seven studies reported an association between PAH and social problems (55) (OR = 1.57, 95% CI, 1.00–2.48), and the remaining six reported no effect (27, 36, 42, 44, 45, 47). Two studies (36, 55) found no effect on rule breaking behavior or somatic complaints. One study (35) reported no associations with attention deficit hyperactivity disorder (ADHD) index scores or hyperactive compulsive behavior, nor did another from the same research group (31) regarding total behavioral problems. Studies reporting neurobehavioral effects are reported in Table 2, and effect sizes are depicted in Figure 7.

Association between pre-natal PM$_{2.5}$ exposure and cognitive abilities and neurobehavioral development in childhood

A study (40) examined high PM$_{2.5}$ exposure during early, mid, and late pregnancy with measures of full-scale IQ score, inattentiveness, and adverse memory performance. Boys highly exposed during late pregnancy exhibit lower IQ and inattention when exposure was from mid to late pregnancy. Girls highly exposed during early to mid pregnancy exhibited adverse memory performance. No effect was reported for the remaining domains analyzed by this study (40).

The finding of impaired motor development (48) was not supported by a subsequent study conducted by the same group (49), which reported, however, impaired memory in boys (49). From studies analyzing the impact of pre-natal PM$_{2.5}$ exposure on cognition and neurobehavioral development (Table 3), no effect was found on visual-motor functioning (61), general cognitive ability (28, 49), mental status (48), non-verbal intelligence (61), adaptive function or autism spectrum disorder (28), nor on verbal, perceptive manipulative, and numeric development (49).
TABLE 1 Studies with measured pre-natal PAH exposure on cognitive abilities in childhood.

| References                        | Sample size | Sample characteristics                                                                 | Male: female | Mean age (SD) | Age range          | Comorbidities                                    | Air pollution data acquisition method |
|-----------------------------------|-------------|----------------------------------------------------------------------------------------|--------------|---------------|--------------------|-------------------------------------------------|--------------------------------------|
| Perera et al. (31)                | 183         | Children 3 years of age, mothers 18–35 years, non-smoking, free of diabetes, hypertension, or known HIV, African American and Dominican women residing for a minimum of a year in Washington Heights, Harlem, or the South Bronx in New York City | 84:99        | 3.5 (0.5)     | 3 years to 3 years 12 months | N/A                                           | Environmental samples analyzed for 8 PAHs |
| Jedrychowski et al. (53)          | 170         | Children 7 years of age, mothers ≥18 years of age, non-smoking, singleton pregnancies, no history of illicit drug use, pregnancy related diabetes, or hypertension, no current occupational exposure to PAH or any other known developmental toxicants, and have been resident in Krakow, Poland for a minimum of a year | 80:90        | 7.5 (0.5)     | 7 years to 7 years 12 months | N/A                                           | Cord blood PAH-DNA adduct            |
| Zhang et al. (47)                 | 211         | Infants 12 months of age, free from delivery injuries, neonatal problems, acquired disabilities, developmental dysplasia or other developmental defects, mothers resident in Qingdao city, China for at least 3 years, free from diabetes, known HIV and known neuropsychiatric disease. | 192:156      | 1.0 (0.083)   | 1 year to 1 year 1 month   | N/A                                           | Cord blood benzo(a)pyrene-DNA adduct (ng/mL) |
| Perera et al. (32)                | 380         | Children 2 years of age, mothers 18–35 years, non-smoking, free of diabetes, hypertension, or known HIV, African American and Dominican women residing for a minimum of a year in Washington Heights, Harlem, or the South Bronx in New York City | N/A          | 2.5 (0.5)     | 2 years to 2 years 12 months | N/A                                           | PAH/aromatic DNA adducts in umbilical cord blood samples |
| Polanska et al. (56)              | 406         | Children 1–2 years of age, mothers had single pregnancy up to 12 weeks of gestation, no assisted conception, no pregnancy complications, no chronic disease, resident in Poland | 192:214      | 1.5 (0.5)     | 1 year to 2 years 12 months | N/A                                           | 1-hydroxypyrene metabolites in mothers’ urine |
| Edwards et al. (54)               | 214         | Children 5 years of age, mothers ≥18 years of age, non-smoking, singleton pregnancies, no history of illicit drug use, pregnancy related diabetes, or hypertension, no current occupational exposure to PAH or any other known developmental toxicants, and have been resident in Krakow, Poland for a minimum of a year | 103:111      | 5.5 (0.5)     | 5 years to 5 years 12 months | N/A                                           | Environmental samples analyzed for 8 PAHs |
| Perera et al. (44)                | 217         | Children 2 years of age, born between either March to June 2002 or March to May 2002, mothers ≥20 years, non-smoking, resident within 2 km of Tongliang power plant | 113:104      | 2.5 (0.5)     | 2 years to 2 years 12 months | N/A                                           | Cord blood benzo(a)pyrene-DNA adducts (ng/mL) |
| Tang et al. (45)                  | 110         | Children 2 years of age, born between March and June 2002; mothers ≥20 years, non-smoking, resident within 2 km of Tongliang power plant | 54:56        | 2.5 (0.5)     | 2 years to 2 years 12 months | N/A                                           | Cord blood benzo(a)pyrene-DNA adducts (ng/mL) |
| Tang et al. (26)                  | 215         | Children 2 years of age, born between either March to June 2002 or March to May 2002, mothers ≥20 years, non-smoking, resident within 2 km of Tongliang power plant | 106:109      | 2.5 (0.5)     | 2 years to 2 years 12 months | N/A                                           | Cord blood benzo(a)pyrene-DNA adducts (ng/mL) |

(Continued)
Childhood

Association between childhood PAH exposure and cognitive abilities and neurobehavioral development

Children exposed to high levels of PAH post-natally exhibited lower general cognitive ability and delayed impaired memory (62). Increased inattentiveness was reported by two studies (50, 62), but this finding was contradicted by one study (51). The negative effect of post-natal PAH exposure was not observed in all memory domains. A study (50) found an association between impaired working numeric memory but not on working verbal memory. Short-term memory was not found affected either (62).

The association with ADHD diagnosis reported by one study (38) was not supported by two other studies (50, 51). Neither study found an effect on learning performance (38, 62) or an association with visual spatial skills, non-verbal test performance, executive function, motor performance (62), or behavioral problems (50). Studies reporting childhood PAH exposure can be found in Table 4.

Association between childhood PM$_{2.5}$ exposure and cognitive abilities and neurobehavioral development

From the four studies reporting post-natal PM$_{2.5}$ exposure on general cognition and neurobehavior (Table 5), one reported that children exposed to high levels of PM$_{2.5}$ post-natally displayed impaired selective and sustained attention (63), but this finding was contradicted by two studies (39, 52) reporting no effect on inattentiveness or attention and executive function, respectively. A study’s report of impaired visual information processing speed (63) was again contradicted by another (39) reporting no association with processing speed. No association was found with working memory (39, 52), episodic memory, language (39), cognitive ability, adaptive function, or autism spectrum disorder (28).

Association between childhood PM$_{2.5}$ exposure and neurodegeneration

Only three studies by the same research group analyzed post-natal PM$_{2.5}$ exposure to neurodegeneration (Table 6). Two
TABLE 2. Studies with measured pre-natal PAH exposure on neurobehavioral development.

| References         | Sample size | Sample characteristics                                                                 | Male:female | Mean age (SD) | Age range          | Comorbidities   | Air pollution data acquisition method |
|--------------------|-------------|----------------------------------------------------------------------------------------|-------------|---------------|--------------------|----------------|---------------------------------------|
| Perera et al. (33) | 351         | Children 9 years of age, mothers 18–35 years, non-smoking, free of diabetes, hypertension, or known HIV, African American and Dominican women residing for a minimum of a year in Washington Heights, Harlem, or the South Bronx in New York City. | 163:188     | 9.01 (0.19)   | 9 years to 9 years 12 months | N/A            | Cord blood benzo(a)pyrene-DNA adducts (ng/mL) |
| Perera et al. (33) | 253         | Children 6–7 years of age, mothers 18–35 years, non-smoking, free of diabetes, hypertension, or known HIV, African American and Dominican women residing for a minimum of a year in Washington Heights, Harlem, or the South Bronx in New York City. | 131:122     | 6.5 (0.5)     | 6 years to 7 years 12 months | N/A            | Environmental samples analyzed for 8 PAHs |
| Perera et al. (34) | 215         | Children 3 years 9 months to 5 years 11 months of age, mothers 18–35 years, non-smoking, free of diabetes, hypertension, or known HIV, African American and Dominican women residing for a minimum of a year in Washington Heights, Harlem, or the South Bronx in New York City. | 87:128      | 4.8 (not reported) | 3 years 9 months to 5 years 11 months | N/A            | Cord blood benzo(a)pyrene-DNA adducts (ng/mL) |
| Pagliaccio et al. (36) | 319 | Children 11 years old, mothers 18–35 years, non-smoking, free of diabetes, hypertension, or known HIV, African American and Dominican women residing for a minimum of a year in Washington Heights, Harlem, or the South Bronx in New York City. | 177:142     | 11.5 (0.5)    | 11 years to 11 years 12 months | Early life stress | Environmental samples analyzed for 8 PAHs |
| Nie et al. (43)     | 247         | Infants 3 days of age, mothers ≥18 years, non-smoking, no chronic disease or family history of neurological disease, single gestational viable fetus, who delivered in the Sixth Hospital of Shanxi Medical University and the Eighth People's Hospital of Taiyuan, resident in Taiyuan for at least a year | 132:115     | 3 days (not reported) | 3 days | N/A | Urinary metabolite concentrations of 2-hydroxyfluorene |
| Perera et al. (55)  | 248         | Children from Krakow, Poland, mothers ≥18 years, non-smoking | 122:126     | 7.28 (0.98)   | 6 years to 9 years 12 months | Maternal psychological distress | Personal air monitoring analyzing concentrations of 8 PAHs |
| Tang et al. (45)    | 110         | Children 2 years of age, born between March to June 2002, mothers ≥20 years, non-smoking, resident within 2 km of Tongliang power plant | 54:56       | 2.5 (0.5)     | 2 years to 2 years 12 months | N/A            | Cord blood benzo(a)pyrene-DNA adducts (ng/mL) |
| Zhang et al. (47)   | 211         | Infants 12 months of age, free from delivery injuries, neonatal problems, acquired disabilities, developmental dysplasia or other developmental defects, mothers resident in Qingdao city, China for at least 3 years, free from diabetes, known HIV, and known neuropsychiatric disease. | 192:156     | 1.0 (0.083)   | 1 year to 1 year 1 month | N/A            | Cord blood benzo(a)pyrene-DNA adducts (ng/mL) |
| Perera et al. (44)  | 217         | Children 2 years of age, born between either March to June 2002 or March to May 2002, mothers ≥20 years, non-smoking, resident within 2 km of Tongliang power plant | 113:104     | 2.5 (0.5)     | 2 years to 2 years 12 months | N/A            | Cord blood benzo(a)pyrene-DNA adducts (ng/mL) |

(Continued)
of them found that children highly exposed to PM$_{2.5}$ post-natally exhibited lower amyloid beta protein fragment 1–42 (Aβ$_{1-42}$) and brain-derived neurotrophic factor (BDNF) (29, 30), with one finding, in addition, higher interferon (IFN) γ concentrations in cerebrospinal fluid (CSF) (30). No effect was found with regard to concentrations of biomarkers: non-phosphorylated tau (non-p-tau), vitamin D, tau phosphorylated at threonine 181 (30), cellular prion protein, total tau, interleukin (IL) β, leptin (29, 30), total alpha- synuclein (α-synuclein), oligodendrocyte α-synuclein, hyperphosphorylated tau, tumor necrosis factor alpha, IL 2, IL 6, IL 10, or monocyte chemotactant protein-1 (MCP-1) (29). Of 33 antibodies to neural and tight junction proteins, actin immunoglobulin G (IgG), occludin/zonulin (OZ) immunoglobulin A (IgA), OZ IgG, myelin oligodendrocyte glycoprotein (MOG) IgG, MOG immunoglobulin M (IgM), myelin basic protein (MBP) IgA, MBP IgG, astrocytic protein (S-100) IgG, S-100 IgM and cerebellar antigen (CERE) IgG in serum, and MBP antibodies in CSF were higher in children exposed to high levels of PAH compared to controls (58).

There was one account of impaired cognitive disturbances (60), which was contradicted by two reports of no association with cognitive dysfunction (37, 59) and by additional individual accounts of no effect in approximate number system functioning (41) (i.e., digit span, digit symbol, number of dots tests), confrontational word retrieval, verbal fluency, delayed reaction time between congruent and incongruent stimuli, visual attention, and task switching (59). Mood state, attention/response speed, manual dexterity, or perceptual motor speed were not found associated with PAH exposure in adulthood (41, 46). However, it must be noted that two investigations were population studies (37, 59), while the other three (41, 46, 60) were occupational health studies on brain effects of PAHs in coke ovens (41, 46) or oil spill (60) workers, who are exposed to very high levels of PAHs, especially high molecular weight compounds including benzo(a)pyrene and other compounds with five to six or more hydrocarbon rings. A summary of the population characteristics of the five studies exploring adult PAH exposure and general cognition can be found in Table 7.

### Adult

#### Association between adult PAH exposure and cognitive abilities

Adults highly exposed to PAH exhibited impaired auditory memory (41, 46), individual accounts of memory disturbances (60), and impaired verbal learning and memory (59). However, there was no effect on working memory and executive function, visuospatial memory/attention and planning (59), or visual perception memory (41, 46).

### Childhood and adult

#### Association between childhood and adult PM$_{2.5}$ exposure and neurodegeneration

Details from the three studies reporting on cohorts inclusive of participants exposed to PM$_{2.5}$ only during childhood and some participants through to adulthood can be found in Table 8. In a cohort of mixed exposure to PM$_{2.5}$, the presence of neurodegenerative biomarkers phosphorylated tau (p-tau), α-synuclein, and transactive response DNA-binding protein 43
TABLE 3: Studies with measured pre-natal PM$_{2.5}$ exposure on cognitive abilities and neurobehavioral development in childhood.

| References            | Sample size | Sample characteristics | Male: female | Mean age (SD) | Age range | Comorbidities | Air pollution data acquisition method |
|-----------------------|-------------|------------------------|--------------|---------------|-----------|---------------|--------------------------------------|
| Blazkova et al. (61)  | 169         | Children 5 years of age, born in the summer 2013 to winter 2014, non-smoking mothers, resident in Karvina and Ceske Budejovice, Czech Republic | 78:90        | 5.5 (0.5)     | 5 years to 12 months | Viral diseases, otitis, bronchitis, GIS, HCD | Analysis of 11 OH-PAHs in urine |
| Kerin et al. (20)     | 325         | Children 2–5 years, resident in catchment area of 20 counties in northern California, the central valley and parts of Los Angeles metropolitan area, US, complete history of environmental air exposure, lived with at least 1 biological parent who speaks English or Spanish | 281:44      | (not reported) | 2 years to 12 months | N/A | Residential addresses inputted into Tele Atlas database and software |
| Lertxundi et al. (49) | 560         | Children 4 years male, mothers ≥16 years, resident in Valencia, Sabadell, and Gipuzkoa in Spain | 560:00      | 4.8 (4.9)     | 4 years to 12 months | N/A | Land use regression models |
| Lertxundi et al. (48) | 438         | Children aged ~15 months age, mothers ≥16 years, singleton pregnancies | 198:240     | 1.25 (0.25)   | 1 year 1 month to 1 year 6 months | N/A | Environmental samples from digital DHA-80 high-volume aerosol samplers |
| Chiu et al. (40)      | 119         | Mothers ≥18 years, at 28.4 ± 7.9 weeks gestation between August 2002 and January 2000 in Boston | 00:119      | 6.5 (9.8)     | 6 years to 7 years 3 months | N/A | Use of a hybrid satellite based spatio-temporal prediction model and residential address during pregnancy |

(TDP-43) was confirmed in brainstems (57). The faster increase in concentrations with regard to the age of non-p-tau in CSF was also associated with increased exposure (30). However, no association was found with regard to the concentration of total and oligomer α-synuclein in CSF (29).

Risk of bias within studies

All studies included were of high quality with reproducible accounts of the method employed to assess relevant outcomes, and the inclusion/exclusion criteria used to select the study population were explained in sufficient detail (refer to detailed QUADAS tool responses in the publicly available data at [https://datashare.ed.ac.uk/handle/10283/3892](https://datashare.ed.ac.uk/handle/10283/3892)). Where applicable, all studies provided explanations for participant withdrawal, which were unrelated to both the exposure and the outcome being measured and reported intermediate or unexpected results. Approximately 54% of studies involved the use of a comparison with a low exposure or control population either by dichotomizing exposure data or using a demographically matched control population. The remaining 46% of studies assessed PAH exposure as a continuous variable. All studies correctly identified confounding variables, and the method and analysis were adjusted accordingly. There was, however, a considerable risk of information bias amongst studies, with only 16% of studies reporting the outcome assessor to be blinded and unaware of the exposure status of the study participant. Seventy-three percent of the studies provided no indication as to whether they were or not blinded, and in 11% the outcome assessors were confirmed not blinded.

Discussion

This review found sufficient evidence that pre-natal PAH exposure negatively impacts cognitive function with specific regard to child intelligence, mental development, verbal IQ, memory impairment, average overall development, child attentiveness, behavioral development, and externalizing, internalizing, anxious, and depressed behavioral problems.

Evidence concerning exposure during childhood and as an adult with cognitive function was insufficient to conduct a meta-analysis, due to a reduced number of studies, low consistency, and high heterogeneity in results. However, associations can be observed such as exposure during childhood with lowered cognitive ability, impaired child attentiveness, and exposure as an adult manifesting in memory disturbances.
TABLE 4 Studies with measured childhood PAH exposure on cognitive abilities and neurobehavioral development.

| References | Sample size | Sample characteristics                                      | Male: female | Mean age (SD) | Age range | Comorbidities | Air pollution data acquisition method |
|------------|-------------|------------------------------------------------------------|--------------|---------------|-----------|---------------|--------------------------------------|
| Suter et al. (62) | 31 | Children aged 5–12 resident in Nairobi, Kenya. Infected with HIV and previously enrolled in the Optimizing HIV-1 Therapy Study | N/A          | 6.6 (0.8)     | 5 years to 12 years 12 months | HIV         | Concentration of urinary PAH metabolite 1-hydroxypyrene (1-OHP) |
| Mortamaïs et al. (51) | 242 | Children 7–10 years, resident and enrolled in one of 40 schools in Barcelona, Spain, no dental braces | 123:119      | 8.4 (0.8)     | 7 years to 10 years 12 months | N/A         | Environmental air sampling |
| Abid et al. (38) | 85 | Children 6–15 years of age, part of a civilian population resident in the US | 58:25        | 11.2 (0.5)    | 6 years to 15 years 12 months | N/A         | Urinary metabolite concentrations of 2-naphthol |
| Alemany et al. (50) | 1589 | Children aged 7–11, attending one of 38 schools in Barcelona, Spain, and one school in the adjacent municipality, Sant Cugat del Vallès | 831: 758     | 8.52 (0.87)   | 7 years to 11 years 12 months | APOE e4 allele | Environmental samples analyzed for 7 PAHs |

with specific regard to auditory memory and verbal learning and memory.

Studies concerning PAH exposure during childhood and as an adult were scarce, but an increased risk of neurodegeneration was found through the presence of neurodegenerative biomarkers and increased concentrations of cryptic “self” antigens in serum and CSF, indicative of the neuroinflammatory pathology which precedes Alzheimer’s disease (AD) and Parkinson’s disease (PD).

It is known that some pathways of aryl-hydrocarbon neurotoxicity are common for PAHs, TCDD, dioxin-like agents, polyphenols, and similar xenobiotics. A review of the neuropathological mechanisms of PAHs highlights that these, together with their metabolites, may cross the blood–brain barrier causing neurological abnormalities that may include neuronal damage, impaired neurotransmitter regulation, parasympathetic dysregulation, and neurodegeneration (65). Preclinical studies hint at a common neuropathological mechanism of PAH action being the binding of these compounds to the aryl-hydrocarbon receptor (AhR), a cytosolic transcription factor that initiates a complex pathway leading to alteration of gene regulation. AhR is also present in neural cells and can be involved in the mechanisms leading to PAH-induced neurological disorders (65).

This review differentially addressed the neurological impact of PAHs in three different domains, namely, cognitive abilities, neurobehavioral development, and neurodegeneration, and can be used as evidence for policy surrounding the monitoring of PAHs specifically. In addition, it raises awareness of the potentially confounding effect that different ambient PAH concentrations, in metropolitan and rural settings, can have on research assessing outcomes concerned with cognitive function and neurodegeneration in studies. It was not possible, however, to conclude on the differential impact of PAHs acquired mainly from outdoor sources from those acquired from indoor sources.

A previous review on the impact of PM$_{2.5}$ in disease incidence did not stratify patients by age nor considered differences between urban and rural areas, rather stratifying studies by the pollution level in which the country was considered (i.e., “lightly polluted” vs. “heavily polluted”) (64). Other reviews have highlighted general adverse health conditions such as chronic asthma, increased incidence of premature death and hospital admissions (15), and kidney and liver damage (16). Some focused specifically on the carcinogenic properties and resulting incidence of the lung (17), urinary tract (18), and skin and gastrointestinal tract cancers (16). Those that focused on the neurological impact of air pollution concerned a diverse mixture of compounds. One specifically focused on non-communicable diseases and the roles of nitrogen dioxide (NO$_2$), nitrogen oxide species (NO$_x$), carbon monoxide (CO), and PM$_{2.5}$ (19). Another (20) raises awareness concerning ambient pollution’s adverse effect on cognitive decline and impairment, concurring with findings from (22), where the emphasis was on ozone, PM$_{2.5}$, and PM$_{10}$. A study (21) reported NO$_2$, NO$_x$, black carbon, and PMs as potential risk factors for AD, PD, and multiple sclerosis. Despite the outcomes assessed being oriented toward neurological health, the exposures measured either include multiple pollutants or compounds NO$_2$, NO$_x$, CO, PM, or black carbon, around which extensive research already exists and has culminated in tight air quality restrictions and monitoring, which is closely adhered to by governing bodies. This review raises awareness of the neurological impact PAHs.
TABLE 5  Studies with measured childhood PM₂.₅ exposure on cognitive abilities and neurobehavioral development.

| References          | Sample size | Sample characteristics                                                                 | Male: female | Mean age (SD) | Age range          | Comorbidities | Air pollution data acquisition method |
|---------------------|-------------|----------------------------------------------------------------------------------------|--------------|---------------|--------------------|---------------|--------------------------------------|
| Cserbik et al.      | 10,343      | Children aged 9–10 years, resident in one of 21 study sites in the US                   | 5,410: 4,933 | 9.93 (0.64)   | 9 years to 10 years and 12 months | N/A           | Ensemble-based model approach combining aerosol optical depth models, land-use regression, and chemical transport models |
| Kerin et al.        | 325         | Children 2–5 years, resident in catchment area of 20 counties in northern California, the central valley and parts of Los Angeles metropolitan area, US, complete history of environmental air exposure, lived with at least one biological parent who speaks English or Spanish | 281:44      | (Not reported) | 2 years to 5 years 12 months | N/A           | Residential addresses inputted into Tele Atlas database and software |
| Rivas et al.        | 2,221       | Children 7–10 years old, attending one of 39 schools in Barcelona, Catalonia, Spain, without special needs | 1,133: 1,088 | 8.5 (0.9)     | 7 years to 10 years 12 months | N/A           | Land use regression models           |
| Saenen et al.       | 310         | Children in grades 3–6 in three primary schools, Flanders, Belgium.                     | 158: 152     | 10.2 (1.3)    | N/A                | N/A           | Chronic exposure: spatial temporal interpolation method to model the daily residential exposure. Recent exposure (at schools): portable devices |

has, independent of other pollutants, the importance of which is paramount with the current health impacts of PAHs in the UK air quality strategy detailed as “possibly” or “probably” carcinogenic, detracting from the seriousness of their impact on neurological human health (2). This review proceeds to categorize outcomes into subgroups depending on the time of exposure to provide further insight into the demographics of the individuals most vulnerable to the pollution levels reported and to differentiate between the areas of cognitive function and neurodegeneration most impacted, elucidating the potential mechanisms of neurotoxicity. The observation that the most profound effect of PAH exposure culminates from the pre-natal period is in keeping with prior research, showing the fetal brain to be more vulnerable to environmental toxic insult than the adult. The increased permeability of the not yet fully formed blood–brain barrier combined with the rapid brain growth during the second trimester means the period of most intense construction and brain architecture is also the time the brain is most vulnerable to the passage of toxins and neurotoxicity (66). Overall, this review has systematically located, summarized, and meta-analyzed evidence about the potent neurotoxicity of direct or indirect exposure to PAHs across the human lifespan, highlighting the need for more well-designed epidemiological studies.

Limitations

Studies included in the analysis were limited to those written in the English language. Publication bias and selective reporting within studies cannot be discarded, nor can indexing issues, in which the search terms may have failed to identify relevant studies.

The study populations included only originated from nine countries, of which the UK was not one. Findings are therefore limited to the environments and seasonal variations in climate found in these countries, and no specific recommendations for the UK, where the present review was conducted, can be made. The studies included also involved the use of different subgroup samples from the same large cohort, due to the necessity and availability of a limited number of longitudinal study databases. Sampling bias cannot, therefore, be disregarded.

Other polluting aryl-hydrocarbons present in the air, in particulate matter (PM₂.₅) and diets, such as 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) and its congeners, dibenzofurans and dioxin-like polychlorinated biphenyls, have been reported to induce similar neurotoxicity and neurological disorders to PAHs. The concomitant exposure to these compounds, which are ubiquitously present as persistent organic pollutants, could have confounded the measured
TABLE 6 Studies with measured childhood PM$_{2.5}$ exposure on neurodegeneration.

| References                    | Sample size | Sample characteristics                                                                 | Male: female | Mean age (SD) | Age range | Comorbidities                  | Air pollution data acquisition method                      |
|-------------------------------|-------------|-----------------------------------------------------------------------------------------|--------------|---------------|-----------|-------------------------------|----------------------------------------------------------|
| Calderón-Garcidueñas et al. (30) | 1) 426 2) 81 | Children admitted to Mexico City hospital, resident in Mexico City Metropolitan area (MCMA) and other small cities in Mexico | 1) 256:161 2) 44:33 | 1) 13.36 (8.82) 2) 11.54 (5.1) | (Not reported) | Lymphoblastic leukemia | Environmental air sampling, for regulating levels above the USEPA standards |
| Calderón-Garcidueñas et al. (22) | 1) 73 2) 126 | Children admitted to Mexico City hospital, resident in Mexico City Metropolitan area (MCMA) and other small cities in Mexico | 1) 42:31 2) 59:70 | 1) 11.7 (5.14) 2) 17.49 (15.98) | (Not reported) | Lymphoblastic leukemia | Environmental air sampling |
| Calderón-Garcidueñas et al. (58) | 111         | Children within 5 miles of Mexico City Metropolitan Area (MCMA) or small control cities in Mexico (Zacatlán and Huachinango, Puebla; Zitaícuaro, Michoacán; Puerto Escondido, Oaxaca; Chalma, Veracruz; Tlaxcala, Tlaxcala), No ETS exposure, lived within 5 miles of an air monitoring station | 54:57        | 13.37 (4.2)   | (Not reported) | N/A                          | Environmental air sampling |

effects of PAHs reported in the studies reviewed. If the primary sources did not disentangle their effects, it is possible that some of the meta-analyzed results embody added effects of these aryl-hydrocarbons on ambient PAHs.

To adjust for heterogeneity, studies were stratified depending on the time point of exposure and outcome assessed; however, this did not account for heterogeneity between evaluators and instruments used, mainly due to the limited number of sources analyzed. In addition to this, the use of five different measures to quantify levels of PAH exposure, as well as the inclusion of quantification of PM$_{2.5}$ as a measure, resulted in heterogeneity in exposure measurement instruments and the inclusion of potential contaminating compounds within PM$_{2.5}$, which would confound results. Finally, there were insufficient data to calculate 95% CI for one study (32), and the request for numeric data from another study received no response (60); hence, the report of an effect on memory and cognitive disturbances was inferred from a figure with no confirmation from the raw data.

Future research

The role of biological sex in the neurotoxic effects of PAH exposure requires further investigation. Sex stratification of data concerning memory impairment in pre-natally exposed populations was contradictory. Further accounts of memory impairment following both childhood and adult exposure should be dichotomized to examine sensitivity between sexes. Pre-natal PAH exposure’s effect on motor development was an area of controversy. Additional research is required in this domain to eliminate ambiguity. Individual reports of a lack of association with motor performance and perceptual motor speed, respectively, were inadequate to clarify such controversy or draw any conclusions.

In addition to this, a more thorough examination of the timescale of PAH exposure is needed, utilizing a smaller scale to determine critical windows.

Stratification by pregnancy term elucidated differential full-scale IQ, inattentiveness, and memory performance results. No effect on the concentration of non-p-tau in CSF was reported following childhood exposure, however when looked at in a mixed cohort of childhood and adult exposure, as association in relation to age progression was reported, indicative of a critical window of exposure.

Furthermore, gene-environment interactions need further analysis, for example PAHs effect on the brain of genetically susceptible populations, such as carriers of the APOE4 allele.

Repeated future analysis on longitudinal cohorts is required to examine the impact of sustained high PAH exposure or subsequent markedly reduced exposure, the effects such fluctuations can have on cognitive function and neurodegeneration, and whether some adverse effects from pre-natal or early life exposure are recoverable.

Future research should identify and analyze the individual contributions and specific synergistic combinations of PAHs on neurological health. This would differentiate and determine the most neurotoxic PAHs and provide evidence for updates in policy, requiring the monitoring
of additional PAHs, rather than only B(a)P. Additional research into the threshold at which PAH is capable of exerting neurotoxic effects would inform policy, with scientific backing to implement a safe limit with regard to neurological health and update the limit of 0.25 ng.m$^{-3}$ B(a)P, which was set only with regard to carcinogenic properties. Furthermore, more studies are needed concerning populations in the UK, to account for the local environmental, climate, and seasonal variations capable of altering PAH's neurotoxic properties.

### Data availability statement

The datasets presented in this study can be found in online repositories. The name of the repository and
accession number can be found below: Edinburgh DataShare, https://datashare.ed.ac.uk/, https://doi.org/10.7488/ds/3031.

Author contributions

All authors listed have made a substantial, direct, and intellectual contribution to the work and approved it for publication.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fneur.2022.1052333/full#supplementary-material

Supplementary Figure 1

Anthropogenic sources (industrial, mobile, domestic and agricultural sources) of PAH release into the environment (figure made using: Biorender.com).

References

1. Cohen AJ, Brauer M, Burnett R, Anderson HR, Frostad J, Estep K, et al. Estimates and 25-year trends of the global burden of disease attributable to ambient air pollution: an analysis of data from the Global Burden of Diseases Study 2015. Lancet. (2017) 389:1907–18. doi: 10.1016/S0140-6736(17)30505-6
2. Department for Environment, Food and Rural Affairs. The Air Quality Strategy for England, Scotland, Wales and Northern Ireland. Norwich: HMSO (2011).
3. The European Parliament and The European Council of the European Union. Directive 2004/107/EC of the European Parliament and of the Council of 15 December 2004 relating to arsenic, cadmium, mercury, nickel and polycyclic aromatic hydrocarbons in ambient air. Brussels: The European Parliament and The Council of the European Union (2015).
4. Agency for Toxic Substances and Disease Registry (ATSDR). Toxicological profile for Polycyclic Aromatic Hydrocarbons (PAHs). Atlanta, GA: U.S. Department of Health and Human Services, Public Health Service (1995).
5. Menicini E, Bocca B. Polycyclic Aromatic hydrocarbons. In: Encyclopedia of Food Sciences and Nutrition. 2nd ed. Academic Press; Elsevier (2003). p. 4616–25. doi: 10.1016/B0-12-227055-X/00939-1
6. Hassanvand MS, Naddaf K, Faridi S, Nahizadeh R, Sowlat MH, Momennia F, et al. Characterization of PAHs and metals in indoor/outdoor PM10/PM2.5/PM1 in a retirement home and a school dormitory: Sci Total Environ. (2015) 527–8:1100–10. doi: 10.1016/j.scitotenv.2015.05.001
7. AGCII. Polycyclic Aromatic Hydrocarbons (PAHs) Biologic Exposure Indices (BEI) Cincinnati, Cincinnati, OH: American Conference of Governmental Industrial Hygienists (ACGIH) (2005).
8. Bach PB, Kelley MJ, Tate RC, McCrory DC. Screening for lung cancer: a review of the current literature. Chest. (2003) 123:1. doi: 10.1378/chest.123.1_supp1.72S
9. Boffetta P, Jounenkov a N, Gustavsson P. Cancer risk from occupational and environmental exposure to polycyclic aromatic hydrocarbons. Cancer Control. (1997) 8:3. doi: 10.1093/jahia/08-106997
10. Diggins DL, Harris KL, Rekhadevi PV, Ramesh A. Tumor microsomal metabolism of the food toxicant, benzo(a)pyrene, in ApcMin mouse model of colon cancer. Tumour Biol. (2012) 33:1–12. doi: 10.1007/s13277-012-0375-6
11. Olsson AC, Fevotte J, Fletcher T, Cassidy A, Mannette A, Zaridze D, et al. (2010). Occupational exposure to polycyclic aromatic hydrocarbons and lung cancer risk: a multicenter study in Europe. Occup Environ Med. (2010) 67:98–103. doi: 10.1136/oom.2009.046680
12. Kuo C-Y, Chien PS, Kuo W-C, Wei C-T, Rau J-Y. Comparison of polycyclic aromatic hydrocarbon emissions on gasoline- and diesel-dominated routes. Environ Monit Assess. (2012) 185:5749–61. doi: 10.1007/s10661-012-2981-6
13. Saunders CR, Das SK, Ramesh A, Shockley DC, Makkerje S. Benzo(a)pyrene-induced acute neurotoxicity in the F-344 rat: role of oxidative stress. J Appl Toxicol. (2006) 26:427–38. doi: 10.1002/jat.1157
14. Hartz AMS, Bauer B, Block ML, Hong J-S, Miller DS. Diesel exhaust particles induce oxidative stress, proinflammatory signaling, and P-glycoprotein up-regulation at the blood-brain barrier. FASEB J. (2008) 22:2723–33. doi: 10.1096/fj.08-106997
15. Abdel-Shafy H, Mansour MS. A review on polycyclic aromatic hydrocarbons. Environmental impact, effect on human health and remediation. Egypt J Petroleum. (2016) 25:107–23. doi: 10.1016/j.ejpe.2015.03.011
16. Kim KH, Ara JS, Kabir E, Brown RJ. A review of airborne polycyclic aromatic hydrocarbons (PAHs) and their human health effects. Environ Int. (2013) 60:71–80. doi: 10.1016/j.envint.2013.07.019
57. Calderón-Garcidueñas L, González-Maciel A, Reynoso-Robles R, Hammond J, Kulesza R, Lachmann I, et al. Quadruple abnormal protein aggregates in brainstem pathology and exogenous metal-rich magnetic nanoparticles (and engineered Ti-rich nanorods). The substantia nigrae is a very early target in young urbanites and the gastrointestinal tract a key brainstem portal. *Environ Res.* (2020) 191:1–17. doi: 10.1016/j.envres.2020.110139

58. Calderón-Garcidueñas L, Vojdani A, Blaurock-Busch E, Busch Y, Friedle A, Franco-Lira M, et al. Air pollution and children: neural and tight junction antibodies and combustion metals, the role of barrier breakdown and brain immunity in neurodegeneration. *J Alzheimers Dis.* (2015) 43:1037–58. doi: 10.3233/JAD-141365

59. Cho J, Sohn J, Noh J, Jang H, Kim W, Cho SK, et al. Association between exposure to polycyclic aromatic hydrocarbons and brain cortical thinning: the environmental pollution-induced neurological effects (EPINEF) study. *Sci Total Environ.* (2020) 737:1–9. doi: 10.1016/j.scitotenv.2020.140097

60. Ha M, Kwon H, Cheong HK, Lim S, Yoo SJ, Kim EJ, et al. Urinary metabolites before and after clean-up and subjective symptoms in volunteer participants in clean-up of the Hebei Spirit oil spill. *Sci Total Environ.* (2012) 429:167–73. doi: 10.1016/j.scitotenv.2012.04.036

61. Blazkova B, Pastorkova A, Solansky I, Veleminsky M Jr, Veleminsky M, Urbancova K, et al. Effect of polycyclic aromatic hydrocarbons exposure on cognitive development in 5 years old children. *Brain Sci.* (2020) 10:619–30. doi: 10.3390/brainsci10090619

62. Suter MK, Karr CJ, John-Stewart GC, Gomez LA, Moraa H, Nyatika D, et al. Implications of combined exposure to household air pollution and HIV on neurocognition in children. *Environ Res Public Health.* (2018) 15:163–76. doi: 10.3390/ijerph15010163

63. Saenem ND, Provost EB, Viaene MK, Vanpoucke C, Lefèvre W, Vrijens K, et al. Nawrot TS. Recent versus chronic exposure to particulate matter air pollution in association with neurobehavioral performance in a panel study of primary schoolchildren. *Environ Int.* (2016) 95:112–9. doi: 10.1016/j.envint.2016.07.014

64. Fu P, Guo X, Cheung FMH, Yung KKL. The association between PM_{2.5} exposure and neurological disorders: a systematic review and meta-analysis. *Sci Total Environ.* (2019) 655:1240–8. doi: 10.1016/j.scitotenv.2018.11.218

65. Olasehinde TA, Olaniran AO. Neurotoxicity of polycyclic aromatic hydrocarbons: a systematic mapping and review of neuropathological mechanisms. *Toxics.* (2022) 10:417. doi: 10.3390/toxics10080417

66. Lanphear BP. The impact of toxins on the developing brain. *Annu Rev Public Health.* (2015) 36:211–30. doi: 10.1146/annurev-publhealth-031912-114413