

REVIEW

Particulate Air Pollution and Osteoporosis: A Systematic Review

Kok-Lun Pang Sophia Ogechi Ekeuku Kok-Yong Chin

Department of Pharmacology, Faculty of Medicine, Universiti Kebangsaan Malaysia, Cheras, 56000, Kuala Lumpur, Malaysia

Abstract: Air pollution is associated with inflammation and oxidative stress, which predispose to several chronic diseases in human. Emerging evidence suggests that the severity and progression of osteoporosis are directly associated with inflammation induced by air pollutants like particulate matter (PM). This systematic review examined the relationship between PM and bone health or fractures. A comprehensive literature search was conducted from January until February 2021 using the PubMed, Scopus, Web of Science, Google Scholar and Cochrane Library databases. Human cross-sectional, cohort and case-control studies were considered. Of the 1500 papers identified, 14 articles were included based on the inclusion and exclusion criteria. The air pollution index investigated by most studies were PM_{2.5} and PM₁₀. Current studies demonstrated inconsistent associations between PM and osteoporosis risk or fractures, which may partly due to the heterogeneity in subjects' characteristics, study design and analysis. In conclusion, there is an inconclusive relationship between osteoporosis risk and fracture and PM exposures which require further validation.

Keywords: particulate matter, PM₁, PM_{2.5}, PM₁₀, bone mineral density, fracture

Introduction

Air pollution is a critical environmental and health issue in both developing and developed countries. According to the World Health Organization (WHO) statistics in 2016, around 91% of the world's population was living with poor air quality. Air pollution is closely associated with the incidence of pulmonary and non-pulmonary diseases, including metabolic disorders, cardiovascular diseases, central nervous system diseases and cancer.²⁻⁷ Recently studies also showed that air pollution predisposed the public to a higher risk of breast cancer and childhood leukaemia, apart from lung cancer.^{8,9} Besides, air pollution is estimated to contribute to 7 million deaths worldwide in 2016.¹

Air pollutants can be categorised into gaseous or solid type. The common examples of gaseous pollutant are ammonia, nitrogen dioxide (NO2), carbon monoxide (CO), sulphur dioxide (SO₂), tropospheric or ground-level ozone (O₃) and volatile organic compounds. 7,10,11 Particulate matter (PM) is the sum of heterogeneous solid air pollutants, comprising water, dust and particles. The composition of PM is highly diverse, which is usually made up of acids, water droplets, elemental carbon (black carbon), organic carbon, polycyclic aromatic hydrocarbons (PAHs), metal dust, geographical mineral dust, and nitrate or sulphate compounds. 10,12-14 The classification of PM is based on its aerodynamic diameter but not its composition, wherein the particles with diameter <10 μm are grouped as PM₁₀, <2.5 μm as PM_{2.5}

Correspondence: Kok-Yong Chin Department of Pharmacology, Faculty of Medicine, Universiti Kebangsaan Malaysia, Level 17, Preclinical Building, Jalan Yaacob Latif, Bandar Tun Razak, Cheras, Kuala Lumpur, 56000, Malaysia Tel +603 9145 9573

Email chinkokyong@ppukm.ukm.edu.my

Received: 19 April 2021

Accepted: 3 June 2021 Published: 24 June 2021 and <1 μ m as PM₁.¹⁰ Ultrafine particles with aerodynamic diameter <0.2 μ m^{15–17} or ≤0.1 μ m^{18,19} are also being investigated but not as common as other PM species, probably due to the limitation in detection technology. These PMs are produced mainly from human activities, such as vehicle emission, coal or biomass combustion and high-temperature industrial works (manufacturing, mining, and agricultural activities).^{10,11,20}

The negative effects of air pollution are partly attributed to PM. 21 Short-term exposure to PM could lead to respiratory discomfort, airway inflammation, lung damages and cardiovascular disorders. 22-24 Chronic exposure to PM is strongly associated with cardiopulmonary diseases, neurological disorders, cancer formation and increased mortality.25-28 PM₁₀ is deposited mainly on the head or nose area, with a slight deposition in the upper respiratory tracts. PM₁ and PM_{2.5} can reach the deeper lung area, including alveoli and terminal bronchioles. 15,29-31 Subsequently, these fine particles could cross the alveolar barrier, enter the systemic circulation and reach several extrapulmonary organs. 15 Mechanistically, PM could induce oxidative and inflammatory damages on respiratory tracts via mitogen-activated protein kinase and Toll-like receptor signalling pathway. 32-34 Additionally, PM_{2.5} and its component, PAHs, also possess genotoxic, mutagenic and clastogenic effects, contributing to its cancer induction properties.^{35–37}

Osteoporosis is a chronic age-related disease of the skeletal system associated with changes in endocrine. metabolic and mechanical factors. 38,39 According to the National Health and Nutrition Examination Survey 2013–2014, nearly 6–11% of adults age \geq 50 years in the United States were osteoporotic. 40,41 Osteoporosis affects mainly the elderly in both sexes, but women have a 4-time higher risk due to lower peak bone mass and the rapid decline of bone mass during menopause. 42 Fragility or atraumatic fractures are the major contributors to osteoporosis-related comorbidity and mortality.⁴³ Bone mass, measured as bone mineral content (BMC) or bone mineral density (BMD), is the surrogate indicators of bone strength. Osteoporosis is defined as a BMD value 2.5 standard deviations or more below the average value for young adult (T-score ≤ -2.5) at the spine, hip or midradius. 44,45 WHO45 and the International Osteoporosis Foundation⁴⁶ recommended using dual-energy X-ray absorptiometry (DXA) to measure the BMD for the diagnosis of osteoporosis. Quantitative ultrasound (QUS) is an alternative bone health screening technology. It is non-invasive, radiation-free and highly portable, and correlated well with DXA measurement. 47,48

Some of the fixed and modifiable risk factors of osteoporosis include sex (female), old age, ethnicity, low body mass index (BMI), menopause, low physical activity, malnutrition, use of glucocorticoid, smoking, alcohol consumption and chronic diseases like diabetes and chronic kidney disease (CKD). 49 The previously neglected role of pollution, like air pollution, as a risk factor of osteoporosis, is gaining attention in the recent 5 years. ^{50,51} Several molecular mechanisms were postulated in explaining the association between PM and osteoporosis risk/fracture (reviewed in Prada et al. 52). Several preclinical and epidemiological studies reported the pro-inflammatory properties of PM by increasing the inflammatory cells and acute response protein level and inducing inflammatory-related diseases like airway inflammation, cardiovascular diseases and arthritis.^{2,53-62} The upregulated inflammatory cytokines, including tumour necrosis factor-α, interleukin-1β, interleukin-6 and granulocyte-macrophage colony-stimulating factor, are osteoclastogenic and could stimulate bone resorption. 63-66 Moreover, PM2 5 and PM10 exposures were significantly associated with serum receptor activator of nuclear factor-kappa B ligand level, suggesting their osteoclastogenic properties.⁵³

Furthermore, PM exposure has been linked with vitamin D deficiency (reviewed in Afsar et al⁶⁷). PM exposure was positively associated with kidney diseases and negatively associated with kidney function in converting the inactive 25-hydroxyvitamin D to biologically active 1,25dihydroxyvitamin D. 68-70 Additionally, PM components like metal dust are nephrotoxic. 52 Moreover, PM also reduces the cutaneous vitamin D biosynthesis in populations with normal kidney function⁷¹ by reducing the surface solar⁷² and ultraviolet radiation.⁷³ Epidemiology studies reported lower serum vitamin D levels among healthy women, adolescents and children from the polluted area. 74-76 Additionally. PM components like PAHs were also reported to increase vitamin D catabolism.77 Nevertheless, the causal relationship between PM and vitamin D level is not yet confirmed.

To the best of our knowledge, a systematic review that summarises the relationship between PM and bone health or fractures is not available. Therefore, this systematic review aims to summarise the relationship between PM and bone health or fracture in the human population.

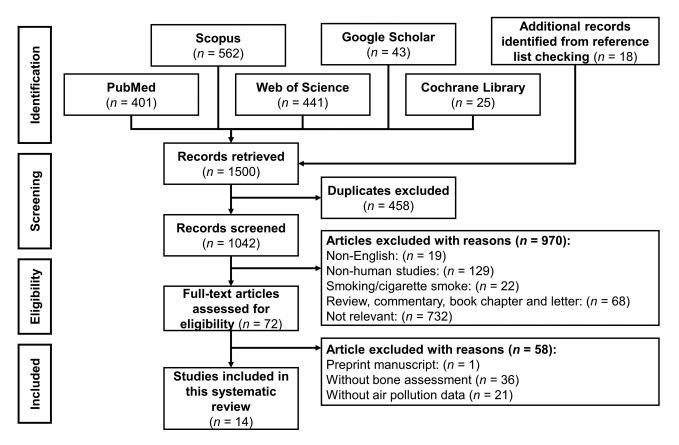


Figure 1 PRISMA flow chart of the systematic literature search.

Notes: PRISMA figure adapted from Moher D, Liberati A, Tetzlaff J, et al. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. BMJ. 2009;339:b2535. 78 Creative Commons.

Materials and Methods

Literature Search Strategies

This systematic review was conducted according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines and checklist. We conducted an electronic search using five databases, including PubMed, Scopus, Web of Science, Google Scholar and Cochrane Library from January to February 2021. The following search string was used: (1) (osteoporosis OR bone OR fracture OR "X-ray absorptiometry") AND (2) ("air pollution" OR "particulate matter" OR PM2.5 OR PM10). A manual search was performed to retrieve additional records from the reference list of included studies or review papers. The detailed search strategy is provided in Supplementary Table S2. The PRISMA checklist is included as Supplementary Table S1.

Eligibility Criteria and Study Selection

We included cross-sectional, case-control, longitudinal/prospective and retrospective cohorts that reported the relationship between air pollutants, primarily particulate matter, and bone health, osteoporotic risk or fractures published within 30 years, from 1990 to 2021. We excluded studies that were (1) only available in abstract form; (2) not written in English; (3) books, book chapters, reviews, meta-analysis, conference/proceeding papers, letter to editor, and commentary; (4) pollutants or PM from smoking, cigarette smoke and tobacco; (5) no PM measurement; or (6) without bone mass assessment. The PRISMA flow chart that summarises the records identification, screening, eligibility, and inclusion of articles, are shown in Figure 1.

Study Extraction

Two reviewers (K.-L.P. and S.O.E) independently extracted the data from each article into an extraction table, firstly by referring to the title and abstract, followed by a full-text screening. Discussions with the third reviewer (K.-Y.C.) were held if there was any disagreement in the inclusion of an article. Table 1 shows the data retrieve from the articles, including the name of the first author, year of publication, year of subject recruitment/ study period, study location, study design, number of subjects, PM assessment, bone health, osteoporosis or fracture assessment and outcomes.

 Table I
 The Design and Major Findings of the Included Studies

	JBI Score	ω	٢	9	ω
Cross-Sectional Studies	Outcomes	Total body BMD was negatively associated with PPh _{2.5} [β = -47, 95% Cl= (-77, -17] and PM _{1.0} [β = -28, 95% Cl= (-48, -8)] after BM _{1.5} smoking, physical activity and education adjustment Risk of low total body BMD was positively associated with PM _{2.5} [OR= 1.33, 95% Cl= (1.05, 1.70)] and PM _{1.0} levels [OR= 1.28, 95% Cl= (1.00, 1.63)] after multivariate adjustment	The forearm fracture was not significantly associated with $PM_{2.5}$ and PM_{10} regardless of age and sex (all p>0.05) Distal forearm BMD was not associated with $PM_{2.5}$ and PM_{10} among subjects aged 59–60 years old and 75–76 years old regardless of sex after adjustment of education, smoking, years of smoking, physical activity and years after menopause (all p>0.05)	The total body BMD (but not hip BMD) was negatively associated with PM ₁₀ without covariate adjustment	Osteoporosis risk was not significantly associated with PM ₁₀ level [OR= 1.012, 95% CI= (0.949, 1.079)], after adjusted for age, sex, BMI, current smoking, drinking, inhaled corticosteroids usage, 6-minute walk distance, COPD severity and smooth functions of visit date and yearly temperature
	Bone Health, Osteoporosis or Fracture Assessment	Total body BMD by DXA Low BMD = BMD Z-score ≤ -1	Mean distal forearm BMD by DXA Self-reported forearm (wrist/lower arm) fracture	BMDs of total body, lumbar spine or femoral neck by DXA	The hip, femoral neck and lumbar spine BMD by DXA Followed WHO definition of osteopenia and osteoporosis
	PM Assessment	Annual mean PM _{2.5} and PM ₁₀ levels from the Norwegian Institute of Air Research, AirQUIS system	Annual mean PM _{2.5} and PM ₁₀ levels from outdoor air pollution exposure estimated by EPISODE dispersion model, the Norwegian Institute of Air Research	Annual mean PM ₁₀ from EPA Bihor	An annual mean concentration of PM ₁₀ calculated from the daily data from monitoring stations operated by the Taiwan Environmental Protection Administration
	Sample Size	590 men aged 75–76 years old	5976 subjects with men (n= 2674) and women (n=3302) aged 59–60 or 75–76 years old	IO5 subjects aged 62.2 ± 3.98 years old	70 retired workers with men (n=68) and women (n=2) aged: 75.2 ± 5.5 years old
	Study Design	Cross- sectional study	Cross- sectional study	Cross- sectional study	Gross- sectional study
	Study Location	Oslo, Norway	Oslo, Norway	Oradea, Romania	New Taipei City, Taiwan
	Study Period	2000–2001	2000–2001	Jan – Dec 2009	l Jan 2010–31 Dec 2012
	Author	Alvær et al, 2007 ⁸¹	Alver et al 2010 ⁸²	Cevei and Stoicanescu 2010 ⁸³	Lee et al 2014 ⁸⁴

ω	7
PM _{2.5} was not associated with the total body (β=-0.05, p=0.75) and pelvic BMD (β=-0.13, p=0.76) after adjusting for age, sex, weight, and height Total body and pelvic BMD were negatively associated with residential distance from the nearest freeways (≤ 500 m; all p< 0.05) after adjusting for age, sex, weight, height, body fat percentage, menopausal status, gestational diabetes, physical activity, ambient pollutants, contextual variables, daily calcium and vitamin D intakes. Total body BMD (but not pelvic BMD) was negatively associated with residential distance from a major road (75–150 m; all p< 0.05) after multivariate adjustment.	PM _{2.5} and PM ₁₀ exposures were significantly higher among subjects with normal BMD (all p<0.05). NS association between BMD T-score and PM _{2.5} [β= -0.002; 95% CI (-0.006, 0.002); p= 0.311] and PM ₁₀ [β= 0.001; 95% CI (-0.002, 0.004); p= 0.491] after adjusting for age, sex, smoking history, diabetes, hypertension, BMI, systolic blood pressure, diastolic blood pressure, fasting glucose, triglyceride, total cholesterol, high-density lipoprotein-cholesterol, how-density lipoprotein-cholesterol, haemoglobin, estimated glomerular filtration rate, uric acid, and regular exercise.
Total body and pelvic BMD by DXA	Bone health of the non-dominant foot by calcaneal QUS Followed WHO definition of osteopenia and osteoporosis
Annual mean ambient PM _{2.5} level from the US Environmental Protection Agency's Air Quality System	Annual mean concentrations of PM _{2.5} and PM ₁₀ levels from the Taiwan Air Quality Monitoring Database
I 173 Mexican American with men (n=324) and women (n=849) aged 34.4 years old	4595 subjects with men (n=2118) and women (n=2477) aged 49.7 ± 10.7 years old
Cross- sectional study	Cross- sectional study
California, United State	Taiwan
2002–2008	2012 -April 2014
Chen et al 2015 ⁸⁵	Lin et al 2020 ⁸⁶

(Continued)

Table I (Continued).

	JBI Score	7	ω
Cross-Sectional Studies	Outcomes	Osteoporosis risk were positively associated with PP4, [56.5–57.7 µg/m³: OR= 1.068, 95% CI= (1.357, 1.907); >57.7 µg/m³: OR= 2.075, 95% CI= (1.724, 2.497)]; PP4, ₂₅ [>73.2 µg/m³: OR= 2.280, 95% CI= (1.899, 2.738)] and PP4, [1.83–13.0 µg/m³: OR= 1.770, 95% CI= (1.492, 2.100); >133.0 µg/m³: OR= 1.770, 95% CI= (1.492, 2.100); >133.0 µg/m³: OR= 1.929, 95% CI= (1.602, 2.322)] after adjusting for age, sex, education level, marital status, smoking, drinking, physical activity, dietary habits, and PP4,, PP4, ₂₅ and PP40 (for each 1 µg/m³ increase) were associated with a 14.9%, 14.6% and 7.3% higher risk of osteoporosis. An estimated 20.29% (PP4), 2.3.20% (PP4) ₂₅ and 24.36% (PP4 ₁₀) osteoporosis cases could be prevented by reducting the exposed PPs below their respective first quartile limit.	PM _{2.5} (for every 3 µg/m³ increase) was negatively associated with lumbar spine BMC [mean difference= −0.57 g, 95% Cl= (−1.06, −0.07)] after adjusting for the bone area, DXA type, age, sex, sex-by-age interaction, percentage lean, percentage fat body mass, fruit intake, vegetable intake, calcium intake, physical activity, smoking, household cooking fuel, occupation, education and standard of living index. NS association between PM _{2.5} and lumbar spine BMC, hip BMC and hip BMD. Subgroup analysis revealed that PM _{2.5} was negatively associated with hip and spine BMC and hip (but not spine) BMD for those aged ≥ 40 years old.
	Bone Health, Osteoporosis or Fracture Assessment	Bone health of the non-dominant foot by QUS Followed WHO definition of osteoporosis	BMC, and BMD at left hip and lumbar spine LI-L4 were measured via DXA
	PM Assessment	3-year average PM1, PM2,5 and PM10 levels were estimated using machine learning algorithms (random forests model) with satellite remote sensing, land use information, and meteorological data	Annual mean exposures of PM _{2,5} was estimated from sampling data and landuse regression models
	Sample Size	8033 subjects with men (n=3001) and women (n= 5032) aged 55.8 ± 10.8 years old	3717 subjects with men (n= 2006) and women (n= 1711) aged 35.7 ± 14 years old
	Study Design	Cross-sectional study	Cross-sectional study
	Study Location	Five rural regions (Suiping county, Vizzhou county, Xinxiang county, Tongxu county and Yima county) of Henan province, China	Hyderabad, South India
	Study Period	July 2015- Sept 2017 (based on Henan Rural Cohort study)	2009–2012
	Author	Qiao et al 2020 ⁹⁷	Ranzani et al 2020 ⁹⁸

ω		JBI score	ω	9	ω
Osteoporotic fractures were positively associated with PN _{2.5} [aHR= 1.10, 95% CI= (1.01), 1.23) for 26.8–29.2 µg/m³, aHR= 1.13, 95% CI= (1.02, 1.24) for 29.2–34.6 µg/m³ among women aged > 50 years, after adjusting for age, household income, Charlson Comorbidity Index and region. NS association between osteoporotic fracture and PM _{2.5–10} and PM ₁₀ Similar positive associations were found between PM _{2.5–10} and PM ₁₀ Similar positive associations were found between PM _{2.5–10} and on-spine fractures [aHR= 1.17, 95% CI= (1.00, 1.38) for 29.2–34.6 µg/m³ and non-spine fractures [aHR= 1.16, 95% CI= (1.01, 1.33) for 29.2 µg/m³, aHR= 1.16, 95% CI= (1.01, 1.33) The sensitive analysis identified a similar positive association with PM _{2.5} after excluding subjects diagnosed with osteoporotic fractures within I year and 2 years (all p=0.008).		Outcomes	NS association between hip fracture and PM _{2.5} [IRR= 1.01, 95% Cl= (0.92-1.09), p = 0.955] and PM _{1.0} [IRR= 1.01, 95% Cl= (0.99-1.03), p = 0.144], regardless of sex or age group (> or ≤ 75 years old; all p> 0.05) after adjusting a natural spline function of time, season and mean air temperature	NS association between osteoporotic hip fracture IRR and $PM_{2.5}$ [r= -0.114 , $p>0.05$] regardless of sex (all $p>0.05$) without covariate adjustment	PM ₁₀ was positively associated with hip fracture incidence [OR= 1.02; 95% CI (1.01, 1.03); p<0.05] after adjusting with age, sex, BMI, annual income, residence area, smoking, alcohol drinking, exercise, Charlson Comorbidity Index and/or an underlying disease
Osteoporotic fracture data from the National Health Insurance Service database, South Korea		Bone health, osteoporosis and fracture assessment	Hip fracture record from the Hospital Universitario Fundacion Alcorcon database	Osteoporotic hip fracture record from the Ministry of Health's Department of Health Statistics and Information, Chile	Hip fracture record from National Health Insurance Service database, South Korea
Annual mean PM _{2.5} , PM _{2.5-10} and PM ₁₀ levels from the Air Korea database based on residential address estimation	Retrospective or prospective cohorts	PM assessment	Daily mean levels of PM _{2,5} and PM ₁₀ from the Air Pollution Monitoring Network, Community of Madrid Environmental Local Government and Regional Planning Department	Annual mean concentration of PM _{2.5} from the National Air Quality Information System, Chile	Annual mean concentration per area of PM ₁₀ from the National Ambient Air Information System, Korean Ministry of the Environment
44.602 women aged > 50 years old	Retro	Sample size	4271 subjects with men (n=925) and women (n=3346) aged 83.8 ± 8.9 years old	8322 elderly aged > 65 years old	178.147 subjects with men (n= 98.749) and women (n=79.398) aged 49.5 ± 12 years old
Gross-sectional study		Study type/ design	Retrospective cohort study	Retrospective cohort study	Retrospective cohort study
Seoul, Incheon and Busan, South Korea		Study location	Alcorcón, Spain	Chile	South Korea
2002–2013		Study period	l Jan 2000–31 Dis 2015	2017	2010–2015
Sung et al 2020 ⁸⁹		Author	Mazzucchelli et al 2018 ⁹⁰	Ormeño Illanes and Quevedo Langenegger 201991	Oh et al 2020 ⁹²

Table I (Continued).

	JBI Score	ω	ω
Cross-Sectional Studies	Outcomes	PM _{2.5} was positively associated with bone fractures [RR= 1.041; 95% CI (1.030, 1.051); p=0.000 I] with a nearly linear relationship after adjusting for sociodemographic variables (age, sex, race, education and income), geographical characteristics, obesity, number of days with freezing temperatures and calendar year. A positive association between PM _{2.5} and bone fracture was observed in women [RR= 1.046; 95% CI (1.036, 1.056); p=0.0002] and men [RR= 1.037; 95% CI (1.027, 1.047);	PM _{2.5} was not significantly associated with BMDs of the femoral neck, total hip, lumbar spines, ultradistal radius and one-third distal radius (all p>0.0.5) after adjusting for age, race, height, weight, smeking, household income, physical activity, caffeine consumption, serum vitamin D levels and/or C-reactive protein. Serum parathyroid hormone (but not serum calcium and vitamin D) was negatively associated with PM _{2.5} [β = -7.39; 95% CI (-14.17, -0.6.1); p<0.05] after multivariate adjustment.
	Bone Health, Osteoporosis or Fracture Assessment	Annual osteoporotic-related bone fractures data (hip, wrist, spine and pelvis) from hospital data	BMDs of the femoral neck, total hip, lumbar spine [L1–L4], distal radius and ultradistal radius by DXA (during baseline and follow-up) Serum parathyroid hormone, calcium (at baseline), and vitamin D [25(OH)D ₂₃]
	PM Assessment	Annual mean PM _{2,5} concentrations estimated using a validated spatiotemporal prediction model	Annual PM _{3.5} concentrations estimated using a validated spatiotemporal prediction model
	Sample Size	763,630 subjects with men (n=314,525) and women (n=449,105) aged ≥ 65 years old	692 African, Latin, and European American male residents aged 30–79 years old with 8 years of follow-up
	Study Design	Retrospective cohort study	Prospective cohort study
	Study Location	Northeast-mid-Atlantic US states	Greater Boston, MA, USA
	Study Period	2003–2010	Nov 2002- Oct 2012
	Author	Prada et al 2017 ⁹³	

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Maternal trabecular bone health was negatively associated with PM _{2.5} exposure (for each 10 µgm³ increase) during first [β=-0.18; 95% CI (-0.35, -0.01); p< 0.05] and third trimester exposure [β=-0.18; 95% CI (-0.36, -0.01); p< 0.05] after adjusting for maternal age, BMI, socioeconomic status, education, parity time since conception, natural trajectory of bone strength changes over time and BM1, socioeconomic at other time and PM _{2.5} concentrations at other time and PM _{2.5} concentrations at other gugm³ increase) during the first month of post-partum [β=-0.20; 95% CI (-0.39, -0.01); p< 0.05]. PM _{2.5} exposure during the 60 days preconception was positively associated with the trabecular and cortical bone health, during the trabecular and cortical bone health, during the trabecular and socioeconomic status, education, parity and PM _{2.5} concentrations at other periods. The first and second trimester PM _{2.5} exposures were initially negatively associated with the bone health of 1 to 6 months post-partum (all p<0.05). NS association for PM _{2.5} exposure during the first month post-partum was negatively associated with trabecular (p<0.05). NS association for PM _{2.5} exposure during the third trimester for both bone health of 1 to 6 months post-partum (p<0.001).
Bone health (SOS T-score) of radius (trabecular) and the proximal phalanx (cortical) of the middle finger by using QUS scan
Daily mean ambient PM _{2.5} was estimated from PM _{2.5} monitoring stations data by using a spatialtemporal model
941 pregnant women aged 27.3 ± 5.5 years old with 3 years of follow-up years of follow-up
Prospective cohort study
Mexico City, Mexico
3rd July 2007–21 st Feb 2011
Wu et al 2020 ⁹⁴

Abbreviations: aHR, adjusted hazard ratio; β , adjusted regression coefficient; BMC, bone mineral content; BMD, bone mineral density; BMI, body mass index; CI, confidence interval; COPD, chronic obstructive pulmonary disease; DXA, dual-energy X-ray absorptiometry; IRR, incidence rate ratio; JBI, Joanna Briggs Institute; LI, first lumbar spine; L4, fourth lumbar spine; NS, not significant; OR, odds ratio; PM, with aerodynamic diameter = 1.5 mm; PM_{1.5}, PM with aerodynamic diameter = 1.5 mm; PM_{1.5}, PM with aerodynamic diameter of pun; QUS, quantitative ultrasound; r, Pearson correlation coefficient; RR, risk ratio; SOS, speed-of-sound; WHO, World Health Organization.

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Quality Assessment

Two reviewers (K.-L.P. and S.O.E.) independently evaluated the article quality using the Joanna Briggs Institute (JBI) critical appraisal checklist. 79,80 Two different checklists were used for cross-sectional and cohort studies covering "Sampling", "Exposure", "Confounding factors", "Outcomes" and "Statistical analysis" domains, with a maximum score of 8 (cross-sectional study) or 11 (cohort study). Every item was rated as Yes (score of 1), No or Unclear (score of 0). Non-applicable item is excluded from the overall scoring. Cross-sectional or retrospective cohort studies with an overall score ≥6 or prospective cohort studies with ≥ 8 were considered as high-quality articles. Any disagreement was resolved by discussion among three reviewers. The overall score was listed in the evidence table (Table 1) and detailed scoring was shown in Supplementary Table S3 and S4.

Result

Search Results and Study Selection

We identified 1500 articles, of which 401 were obtained from PubMed, 562 from Scopus, 441 from the Web of Science, 42 from Google Scholar, 25 from Cochrane Library and 18 additional articles from the reference list of included articles and reviews. A total of 1042 unique records were identified after excluding 458 duplicates. A total of 970 articles were excluded based on the inclusion and exclusion criteria where 19 articles were not written in English, 68 articles were not primary articles, 129 articles were non-human studies, 22 articles studied cigarette smoking and 732 articles were irrelevant to the topic. A total of 72 articles fulfilling the criteria were assessed for eligibility. After examining the full-text, we excluded 58 articles, of which 1 article was a preprint manuscript, 36 articles did not perform bone health assessment and 21 articles did not measure the PM level. Finally, 14 articles were included in this systematic review.

Study Characteristics

The included studies were published between 2007 and 2020, wherein 9 articles were cross-sectional studies, ^{81–89} 4 were retrospective studies ^{90–93} and 2 were prospective studies. ^{93,94} All the articles are considered high quality according to JBI critical appraisal checklist as shown in <u>Table S3 and S4</u>. However, some studies have validity, ^{81,82,86,87,94} confounding factor ^{83,91} and subjects follow-up issues. ^{93,94} Four studies were conducted in North and South America (the United

States, ^{85,93} Chile⁹¹ and Mexico⁹⁴); 4 studies were conducted in Europe (Norway, ^{81,82} Romania⁸³ and Spain⁹⁰), and the remaining 6 studies were conducted in Asia (Taiwan, ^{84,86} Henan province of China, ⁸⁷ India⁸⁸ and South Korea^{89,92}). The total number of participants was 1,024,864, wherein 68,861 were from cross-sectional studies and 956,003 were from cohort studies. Two cohort studies had a sample size >100,000 participants, ^{92,93} whereas 8 studies had a sample size of 1000–100,000 participants, ^{82,85–91} and the remaining 5 studies enrolled <1000 patients. ^{81,83,84,93,94} The participants were mainly elderly, ^{81–84,89–91,93} followed by middleaged adults, ^{86,87,89,92,93} young adults ^{85,88,93} and pregnant women. ⁹⁴

One study investigated PM₁, ⁸⁷ 11 studies ^{81,82,85–91,93,94} investigated PM_{2.5}, 55,56,59-65,67,68 9 studies investigated $PM_{10}^{81-84,86,87,89,90,92}$ and 1 study investigated PM with aerodynamic diameter between 2.5 and 10 μ m (PM_{2.5-10}).⁸⁹ Most of the studies reported the annual mean concentration of PM, except studies by Mazzucchelli et al⁹⁰ and Wu et al⁹⁴ that used the daily mean PM levels (averaged across the study period). WHO air quality guidelines stated that the annual mean $PM_{2.5}$ and PM_{10} levels should not exceed 10 and 20 μg/m³ respectively. 95 The permissible limit for PM₁ has not been established by WHO or other organisations. Most of the study locations had PM level exceeding the permissible limit with an annual mean PM_{2.5} levels ≤10 μg/ $m^{3,81,82,91}$ 10–20 $\mu g/m^{3,85,86,91,93}$ or >20 $\mu g/m^{3,87-89,91}$ and the annual mean PM_{10} levels $\leq 20 \mu g/m^3$, 81,82 or $>20 \mu g/m^3$ m³.8³,8⁴,8⁶,8⁷,8⁹,9² Mazzucchelli et al⁹⁰ reported a daily mean $PM_{2.5}$ and PM_{10} of 9.52–12.34 $\mu g/m^3$ and 23.47– 31.02 µg/m³ respectively, while Wu et al⁹⁴ reported a daily mean PM_{2.5} of 22.3–23.5 μ g/m³.

The bone health status was reported either as fracture incidence. 82,89-93 BMC and BMD assessed via DXA^{81-85,88,93} or QUS method.^{86,87,94} BMDs of total body, 81,83,85 pelvic, 85 hip, 84,88,93 femoral neck, 83,84,93 lumbar spine, 83,84,88,93 distal forearm, 82 distal radius 93 and ultradistal radius⁹³ were measured. Besides, Lin et al.86 Qiao et al87 and Wu et al94 measured bone health via QUS method without DXA validation. Four studies^{83,84,86,87} adopted the WHO definition of osteopenia and/or osteoporosis to classify the subjects.-57,58,60,61 Besides, Alvær et al⁸¹ defined subjects with Z-score ≤ -1 as having low BMD, which does not comply with the existing recommendation. Adjusted regression coefficient (β), 81,82,85,86,93,94 odd ratio (OR), 81,82,84,87 mean differences, 86,88 or Pearson correlation coefficient (r)83 were used in these studies to

demonstrate the association between PM and bone mass. For the association between PM and bone fracture, β , 82 adjusted hazard ratio, 89 incidence rate ratio (IRR), 90 r, 91 OR 92 and risk ratio 93 were used.

Relationship Between PM Exposure and Bone Health or Osteoporosis Risk

The relationship between PM₁ and bone health was scarce. Only a study by Qiao et al⁸⁷ demonstrated that osteoporosis risk was positively associated with PM₁ exposure after multivariate adjustment. Logistic regression analysis also showed that every 1 µg/m³ increase in PM₁ was associated with a 14.9% increased risk of osteoporosis. 87 Besides, an estimated 20.29% of PM₁-related osteoporosis cases could be prevented if PM_1 exposure was $<55.2 \mu g/m^3.87$ Subgroup analysis revealed that the association between PM₁ and risk of osteoporosis were significantly higher among non-alcoholic drinkers, but it was not affected by age, sex, smoking, vegetable or fruit consumption and physical activity.⁸⁷ However, this study classified osteoporosis based on OUS assessment of non-dominant foot without DXA validation, which could introduce misclassification bias in the study.

The relationship of PM_{2.5} and/or PM₁₀ with bone mass or osteoporosis risk was heterogeneous in other reports, whereby they revealed an insignificant 82,84-86,93 or negative association. 81,83,87,88,94 A cross-sectional study (Oslo Health study) on 1039 subjects aged 59-60 and 75-76 years old by Alver et al⁸² demonstrated that the distal forearm BMD was not significantly associated with PM_{2.5} and PM₁₀ exposures regardless of age and sex after multivariate adjustment. Similarly, a cross-sectional study by Chen et al⁸⁵ also reported that ambient PM_{2.5} exposure was not significantly associated with total body and pelvic BMDs among 1173 Mexican American women with an average age of 34.4 years after covariate adjustment. Lee et al⁸⁴ reported that osteoporosis risk was not significantly associated with PM₁₀ exposure among 70 COPD patients aged 75.2 ± 5.5 years after multivariate adjustment. Similarly, Prada et al⁹³ also reported that PM_{2.5} exposure was not significantly associated with femoral neck and ultradistal radius BMDs in a population-based prospective cohort study with 692 men aged 30-79 years old after 8-year of follow-up. Another recent cross-sectional study by Lin et al⁸⁶ on 4595 Taiwanese $(49.7 \pm 10.7 \text{ years old})$ also revealed that PM_{2.5} and PM₁₀ exposures were not significantly associated with QUS

readings of the non-dominant foot. In their study, the bone health of subjects remained normal even they were exposed to higher levels of PM_{2.5}.86

Other studies demonstrated that PM exposure is a significant risk factor for osteoporosis. The cross-sectional Oslo Health study (590 men aged 75-76 years) demonstrated that total body BMD was negatively associated with PM_{2.5} and PM₁₀ after multivariate adjustment.⁸¹ Besides, the risk of low total body BMD (Z-score <-1) was positively associated with PM_{2.5} and PM₁₀ exposures.⁸¹ Similarly, a cross-sectional study involving 105 Romanians by Cevei and Stoicanescu⁸³ reported that the total body BMD (but not hip) was negatively associated with PM₁₀ exposure without covariate adjustment. Additionally, a similar negative association of PM_{2.5} and PM₁₀ exposures with bone health was reported in two recent cross-sectional studies by Qiao et al⁸⁷ and Ranzani et al.⁸⁸ The osteoporosis risk was positively associated with PM_{2.5} and PM₁₀ exposures among 8033 Chinese $(55.8 \pm 10.8 \text{ years old})$ from the rural area after multivariate adjustment. 87 Subsequent logistic regression analysis also demonstrated that every 1 µg/m³ increase in PM_{2.5} and PM₁₀ were associated with a respective 14.6% and 7.3% increase in osteoporosis risk. 87 Besides, an estimated 23.20% and 24.36% of PM_{2.5} and PM₁₀-related osteoporosis cases could be prevented if PM25 and PM10 exposure were less than 70.5 and 125.8 µg/m³ respectively.⁸⁷ Subgroup analysis also demonstrated similar associations between osteoporosis risk and PM_{2.5} and PM₁₀ among non-alcoholic drinkers or subjects with low physical activity.87 Ranzani et al88 reported that PM2.5 was associated with lumbar spine BMC but not with left hip BMC, left hip BMD and lumbar spine BMD among 3717 Indian $(35.7 \pm 14 \text{ years old})$ after multivariate adjustment. Subgroup analysis revealed PM_{2.5} exposure was also negatively associated with left hip BMC and BMD, as well as lumbar spine BMC among subjects aged ≥40 years old.⁸⁸

Another recent prospective cohort study on 941 Mexican pregnant women with 3-year of follow-up by Wu et al⁹⁴ also reported that maternal trabecular bone health was negatively associated with PM_{2.5} exposure during first- and third-trimester exposure. Besides, maternal cortical bone health was also negatively associated with PM_{2.5} exposure during the first trimester.⁹⁴ A timespecific subgroup analysis revealed that these associations were biphasic across the time of exposure. PM_{2.5} exposure during 60-day preconception was positively associated with maternal bone health during mid-to-late gestation

but turned into a negative association during 1 to 6 months post-partum period. 94 PM_{2.5} exposures during the firstand second-trimester were initially negatively associated with trabecular and cortical bone health during mid-to-late gestation but then positively associated with bone health during 1 to 6 months post-partum. 94 Higher PM_{2.5} exposure during the third trimester and first month post-partum predicted a slower post-partum bone health recovery. 94 Wu et al considered the radius and proximal phalanx of the middle finger to represent trabecular and cortical bones respectively. We believed that it is a misnomer as QUS cannot differentiate between trabecular and cortical bone, particularly at the radius, which consists of both trabecular and cortical bones.

Relationship of PM Exposure and Bone **Fractures**

Similar to bone health, the relationship between PM exposure with bone fracture was also heterogeneous, wherein insignificant^{82,89–91} or positive associations^{89,92,93} have been reported. A cross-sectional study from Alver et al⁸² reported that self-reported forearm fracture was not significantly associated with PM_{2.5} and PM₁₀ exposure among 5976 elderly regardless of age and sex. Besides, a retrospective cohort study by Mazzucchelli et al⁹⁰ reported PM_{2.5} and PM₁₀ were not significantly associated with hip fracture among 4271 elderly aged 83.8 ± 8.9 years after univariate or multivariate adjustment. Parallelly, another retrospective study by Ormeño Illanesalso and Quevedo Langenegger⁹¹ on 8322 Chilean people aged ≥65 years also reported that the association between PM2.5 and osteoporotic hip fracture was not significant regardless of sex. Subjects from Magallanes (the lowest PM_{2.5} region in Chile) and Aysén region (the highest PM_{2.5} region) showed similar IRR for bone fractures. 91 Nevertheless, the results of this study were not adjusted for confounding factors.91

On the other hand, a recent cross-sectional study from Sung et al⁸⁹ on 44,602 South Korean women aged >50 years old revealed that PM25 exposure but not PM25-10 and PM₁₀ was positively associated with osteoporotic fractures, including both spine and non-spine fractures. A similar positive association was also reported after excluding the subjects with an osteoporotic fracture in the recent 1 to 2 years. 89 Additionally, retrospective cohort studies by Prada et al⁹³ demonstrated a positive association between osteoporotic-related bone fracture and PM2.5 exposure among 763,630 residents aged \geq 65 years old from Northeast-mid-Atlantic US states, which also present in men and women subgroups.⁹³ A nearly linear relationship between PM25 exposure and bone fracture rate was reported across the range between 3 and 22 µg/m³ PM_{2.5} level. 93 Besides, a recent retrospective cohort study on 178,147 South Korean aged 49.5 ± 12 years old by Oh et al⁹² also demonstrated that PM₁₀ was positively associated with hip fracture incidence after multivariate adjustment. Nevertheless, this study was limited by the relatively low hip fracture incidence (n= 919) compared to healthy control (n= 177,228).⁹²

Discussion

The relationships between PM exposure, including PM₁, PM_{2.5}, PM_{2.5-10} and PM₁₀, with bone health and fracture incidence are not conclusive based on current evidence. PM_{2.5} and/or PM₁₀ exposures were demonstrated to be associated with bone fracture positively^{89,92,93} or not significantly. 82,89-91 Similarly, the association between PM_{2.5} and/or PM₁₀ with bone mass was negative^{81,83,87,88,94} significant.82,84-86,93 or not Interestingly, Wu et al demonstrated a time-specific biphasic association between PM_{2.5} exposure and bone health measured by QUS among pregnant women. On the other hand, relevant findings on PM₁ are scarce as only one study demonstrates a positive association with osteoporosis.⁸⁷ The relationship between PM₁ and bone fracture is yet to be determined. Besides, only one study reported a non-significant association between PM_{2.5-10} and osteoporotic fractures, 89 and its relationship with bone mass is yet to be determined.

The inconsistent findings in the relationship between PM and bone mass/fracture may be partly due to the heterogeneous sample size. Five included studies with sample sizes of less than 1000 patients may not accurately represent the studied population and introduce bias in interpretation. 81,83,84,93,94 Besides, an appropriate adjustment for confounding factors is essential to avoid confounding effects and false interpretation of causality. For instance, Alver et al82 reported that distal forearm BMD was negatively associated with PM2.5 and PM10 levels among men aged 75-76 years old. However, these associations were not significant after adjustment for education, smoking, years of smoking, physical activity and years after menopause. Similarly, Ranzani et al⁸⁸ also reported that the negative association between PM_{2.5} exposure and lumbar spine BMD became insignificant after

adjusting for additional covariates. Additionally, several included studies did not perform covariate adjustment, 83,91 casting some doubts on the validity of the results. Furthermore, critical covariates such as sunlight exposure, vitamin D level, dietary pattern and inflammatory status were not considered in most of the included studies, contributing to the inconsistency of findings.

Additionally, the detection methods of PM may partly contribute to the inconsistent findings in the relationship between PM and bone mass/fracture. PM metric like particle mass is commonly measured using gravimetric and optical methods. 96 However, these detection technologies vary in terms of practicability (cost, size, noisiness and mobility), precision, accuracy and sensitivity/detection limits.96 Studies included in the current review assessed the PM data, but the underlying detection technologies were not disclosed. Additionally, PM statistical modelling is commonly used to estimate and predict indoor air quality and individual exposure to PM because direct individual PM exposure measurement is technically impractical or difficult to be performed.⁹⁷ There are several factors needed to be considered in developing and calibrating the PM levels, including geographical location, meteorological/climate conditions and aerosol optical depth. 97,98 Most studies employed a spatiotemporal prediction model by adjusting the subjects' geographical or residential location. However, some studies did not describe how individual PM exposure estimation was performed or PM modelling was developed. 83,84,89,91,92 Some studies employed the previously reported prediction model^{81,82} or self-developed spatiotemporal prediction model without disclosing the cross-validation values. 85,86,90 In several studies, the PM model was adjusted/calibrated for climate, weather and/or traffic conditions. 85,87,93,94 However, some studies like Lee et al and Mazzucchelli et al measured the climate and weather conditions but did not include them in PM model calibration.^{84,90} The remaining studies did not disclose the PM modeling calibration. It is noteworthy that variations in PM modelling may contribute to inaccurate individual PM exposure estimation, leading to inconsistency in the findings between PM exposure and bone health.

Osteoporosis is diagnosed based on the BMD T-score of any major common bone fracture sites, such as at the spine, hip or mid-radius. However, T-score discordance at different bone sites is not an unusual observation, probably due to the non-homogeneous process of bone loss. 103,104 Increasing the number of bone sites scanned will increase

the chance of discordance and detecting osteoporosis. 100 Thus, the number of bone sites examined could influence the relationship between PM exposure and osteoporosis risk. As evidence, Cevei and Stoicanescu⁸³ and Ranzani et al⁸⁸ demonstrated inconsistent associations between PM and bone mass at different sites. Moreover, several studies^{81,83,85} employed the total body BMD as the skeletal outcome of interest, which is less sensitive than regional BMDs. 105 Although total body BMD correlates with regional BMDs, 103 this value is not used to diagnose osteoporosis per WHO recommendation. Besides, some studies used QUS to define the bone health of the subjects. 86,87,94 DXA and QUS adopt different technology in identifying bone health, so their results are not interchangeable. 48 The WHO classification system to diagnose osteoporosis based on BMD T-score cannot be used for OUS. 106 Although OUS indices correlate with bone mass and several bone microarchitectural indices, 48 they cannot be used directly to infer bone strength as in the studies of Wu et al. 94 A biomechanical assessment, like the three-point bending flexural test, ¹⁰⁷ can indicate bone strength directly. ¹⁰⁸ However, the destructive nature of this test prohibits its use among live subjects. Reference point indentation or micro-indentation test is an alternative method to estimate bone strength in vivo directly. 109 Additionally, PM species are generally present together^{82,87} and coexist with other air pollutants like nitrogen monoxide, NO2 and SO2. 87,90 These air pollutants were reported positively correlated with osteoporosis risk.^{86,99} Besides, there are synergistic effects between CO-nitrogen oxide and SO₂-NO₂, which could further reduce BMD.⁸⁶ Therefore, it is impossible to attribute the skeletal effects to a single PM species or single air pollutants. Additionally, residential proximity to the nearest freeway (≤500 m) but not PM_{2.5}, NO₂ and O₃ exposure was negatively associated with total body and pelvic BMD after multivariate adjustment, including the pollutants levels.85 This observation suggests that other air pollutants like PAHs or black carbon from vehicle exhaust emissions might also contribute to BMD reduction.85,88

Previous studies observed a higher bone mass and fewer bone fractures among subjects from rural areas than urban areas. 110-121 Several factors such as occupation, lifestyle, physical activity, dietary pattern and traffic accident are attributed to this observation. It would be interesting to ask whether air quality could contribute to the difference in bone health between rural and urban populations. However, studies included in this review showed that it might be erroneous to presume rural areas are less polluted. For instance, Qiao et al⁸⁷

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demonstrated 6 to 7 times high annual mean PM_{2.5} and PM₁₀ levels in the rural area than the WHO air quality standard. High PM levels in the rural area could be contributed by increasing numbers of factory and biomass usage or burning fuel activities. ^{53,59,87,122} Besides, biomass cooking in the rural area had been demonstrated to produce significantly higher PM_{2.5} and/or PM₁₀ levels compared with liquid petroleum gas. ^{53,59}

This systematic review, like others, has its limitations. This review did not include unpublished, grey literature and proceeding articles without complete data. Besides, we limited those studies in the recent 30 years as PM-related research began receiving attention from 1990 onwards. However, it is still possible that we might miss out on some important studies. We tried to minimise this limitation by referring to the reference lists of included articles. Moreover, we did not perform a meta-analysis due to the heterogeneity of the study design, outcomes and analysis. The included studies consisted of cross-sectional, retrospective and prospective cohort studies adopting various statistical strategies, the definition of bone health and PM types, hindering meta-analysis from being conducted.

Conclusion

The current literature suggests an inconclusive association between PM exposures and osteoporosis risk and/or fracture, potentially due to the heterogeneity in subject characteristics, study design, sample size, outcome measurement and covariate adjustment during analysis among various studies. Further validation in human studies is required to validate the positive association between PM_{2.5} and/or PM₁₀ and osteoporosis risk or fracture. Furthermore, most of the studies emphasised on PM_{2.5} and PM₁₀ with a limited number of studies on PM₁. It is crucial to investigate the potential relationship between PM₁ and other ultrafine particles with bone health/fracture.

Acknowledgments

The authors thank the Universiti Kebangsaan Malaysia for the Research University Grant under grant number GUP-2020-021. K.-L.P. and S.O.E are post-doctoral researchers funded by Universiti Kebangsaan Malaysia through RGA-1 and FPR-1 grants.

Disclosure

The authors report no conflicts of interest in this work.

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