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Systematic Review

A systematic review of vitamin D status in populations worldwide

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Abstract

Vitamin D deficiency is associated with osteoporosis and is thought to increase the risk of cancer and CVD. Despite these numerous potential health effects, data on vitamin D status at the population level and within key subgroups are limited. The aims of the present study were to examine patterns of 25-hydroxyvitamin D (25(OH)D) levels worldwide and to assess differences by age, sex and region. In a systematic literature review using the Medline and EMBASE databases, we identified 195 studies conducted in forty-four countries involving more than 168 000 participants. Mean population-level 25(OH)D values varied considerably across the studies (range 4·9-136·2 nmol/l), with 37·3 % of the studies reporting mean values below 50 nmol/l. The highest 25(OH)D values were observed in North America. Although age-related differences were observed in the Asia/Pacific and Middle East/Africa regions, they were not observed elsewhere and sex-related differences were not observed in any region. Substantial heterogeneity between the studies precluded drawing conclusions on overall vitamin D status at the population level. Exploratory analyses, however, suggested that newborns and institutionalised elderly from several regions worldwide appeared to be at a generally higher risk of exhibiting lower 25(OH)D values. Substantial details on worldwide patterns of vitamin D status at the population level and within key subgroups are needed to inform public health policy development to reduce risk for potential health consequences of an inadequate vitamin D status.

Key words: Vitamin D: Populations: Public health



Vitamin D plays an important role in bone mineralisation and other metabolic processes in the human body such as Ca and phosphate homeostasis and skeletal growth^(1,2). Vitamin D deficiency, for example, causes rickets in children, leading to skeletal abnormalities, short stature, delayed development or failure to thrive⁽³⁾. In adults, low values of vitamin D are associated with osteomalacia, osteopenia, osteoporosis and subsequent risk of fractures⁽¹⁾. In addition to beneficial effects on musculoskeletal health, observational studies have suggested that low 25-hydroxyvitamin D (25(OH)D) values are associated with an increased risk for several extraskeletal diseases including cancer, infections, autoimmune diseases and CVD⁽⁴⁾. In light of the global ageing population⁽⁵⁾, an almost fourfold increase in osteoporotic hip fractures

since 1990⁽⁶⁾ and the possible risk of other chronic diseases, patterns of low 25(OH)D levels are of substantial public health interest.

Vitamin D status is traditionally measured through assays of 25(OH)D, the major circulating form of vitamin D⁽⁷⁾. Although 25(OH)D levels below 25 nmol/l have been associated with disorders of bone metabolism⁽⁸⁾ and are used to indicate severe vitamin D deficiency, the threshold for defining adequate stores of vitamin D in humans has not been established clearly⁽⁹⁾. The Institute of Medicine has suggested, for example, that approximately 97.5% of the population across all age groups meet their requirements for vitamin D, having serum 25(OH)D values higher than 50 nmol/l⁽¹⁰⁾. However, others consider 25(OH)D values of 75 nmol/l or higher to be adequate^(11,12).

Abbreviation: 25(OH)D, 25-hydroxyvitamin D.

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Given the absence of uniformly accepted definitions, previous reviews have reported substantial variations in the prevalence of vitamin D deficiency across countries throughout the world, with estimates ranging from 2 to 90% depending on the cut-off value and study population selected^(8,13-16). Insights from these earlier studies are limited, however, due to a focus on specific geographical regions, age or risk groups. Moreover, use of a binary approach to define the presence of vitamin D deficiency in some studies might have also obscured important relationships with chronic disease that might exist across a broader spectrum of values.

To provide a basis for future efforts to limit the health consequences of vitamin D deficiency and insufficiency worldwide, we conducted a systematic literature review of studies performed worldwide using continuous values for 25(OH)D to enable comparisons across studies and between different subgroups within the population. The specific objective of the present study, therefore, was to assess vitamin D status across a range of values at the population level and within key population subgroups defined by age, sex and region.

Methods

Literature search

We searched the Medline and EMBASE databases for original articles on vitamin D status in the general population. Keywords were chosen from the Medical Subject Headings terms and the EMTREE thesaurus, respectively, using the following search strategy: (vitamin D/D3 OR 25-hydroxyvitamin D/D3 OR 25(OH)D/D3 OR calcidiol) AND (epidemiologic studies OR population-based OR population OR survey OR representative OR cross-sectional OR observational) NOT (dihydroxycholecalciferols OR case reports OR case-control studies OR clinical trials OR reviews) AND humans. Search terms for vitamin D included the controlled term 'vitamin D' (including calcifediol and 25-hydroxycholecalciferol) and several free-text terms taking different notations of 25(OH)D into account.

Articles published in English between 1 January 1990 and 28 February 2011 (date of the final screen) were considered. We excluded articles published before 1990 because of a general shift in lifestyle, particularly in industrialised nations (e.g. spending less time outdoors), that might have affected mean population-level 25(OH)D values⁽¹⁷⁾. The final screen produced 2566 hits from both databases after excluding 449 exact duplicates identified using EndNote X6 (Thomson Reuters). Wherever possible, the methods used in the present review follow the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) statement (18).

Study selection

Studies were included in the present review if they met the following criteria defined a priori: (1) outcomes - report of mean or median plasma level for 25(OH)D; (2) study participants - randomly selected samples from the general population as well as subgroups defined by age, sex and specific areas within a country; (3) study designs - cross-sectional studies or baseline data from population-based cohorts. Studies were excluded if vitamin D status was estimated (e.g. through self-reported nutritional intake) or if data were available only on vitamin D2. We also did not consider studies using a binary indicator for vitamin D deficiency or insufficiency as the sole outcome measure, given differing thresholds used in the literature to define either state⁽⁵⁾. Furthermore, clinical samples or studies restricted to subgroups with specific characteristics (e.g. ethnicity, job and skin colour) were excluded, as they were not randomly selected from the general population.

All studies were independently screened and evaluated for selection by two of the authors (R. H. and A. F.). Inter-rater agreement was good to moderate, and disagreements were discussed and resolved by consensus in each case (abstract selection: $\kappa = 0.719$; full-text selection: $\kappa = 0.544$). Following the application of exclusion criteria to information contained in the study abstract, we reduced the 2566 screened records to 601 (Fig. 1); application of these criteria following review of each full-text article reduced the pool of potentially eligible articles to 272. Given the presence of multiple reports based on the same data, our final analytical sample comprised 195 unique studies. In several instances, multiple articles from single studies were retained for analysis as they provided separate 25(OH)D values for subgroups with the characteristics of interest (age, sex and region).

Data extraction, data elements and quality assessment

Each study was evaluated using a standardised data extraction form. In each case, we assessed a wide range of variables including vitamin D values, assays used and study characteristics as well as characteristics of the study population and method of recruitment. Data from most studies were represented in the dataset by a single entry for the total study population. Multiple subentries for a single study were included if data were presented by age, sex or region. All 25(OH)D values were expressed in nmol/l, following conversion from ng/ml (multiplied by a factor of 2·496) as necessary.

Based on the WHO recommendations, we classified geographical regions as follows: Latin America; North America; Europe; Asia/ Pacific; Middle East/Africa⁽¹⁹⁾. To determine age-related differences, we defined four age groups: newborns/infants (0-1 years); children/adolescents (>1-17 years); adults (>17-65 years); elderly (>65 years). In instances where details about age were not provided, we created a separate category ('other'). Where possible, we also distinguished elderly living in nursing homes (institutionalised elderly) from those living in the community.

We assessed study quality using data reported in each study on representativeness, validity and reliability. A study was considered representative if (1) this feature of the study was explicitly addressed in the corresponding full-text article or (2) any statement made by the authors suggested that the actual sample reflected the target population. A study was classified as non-representative if the corresponding full-text article contained information about an existing selection bias, which might also occur in a randomly selected sample (e.g. overestimation of females). Measurement validity was





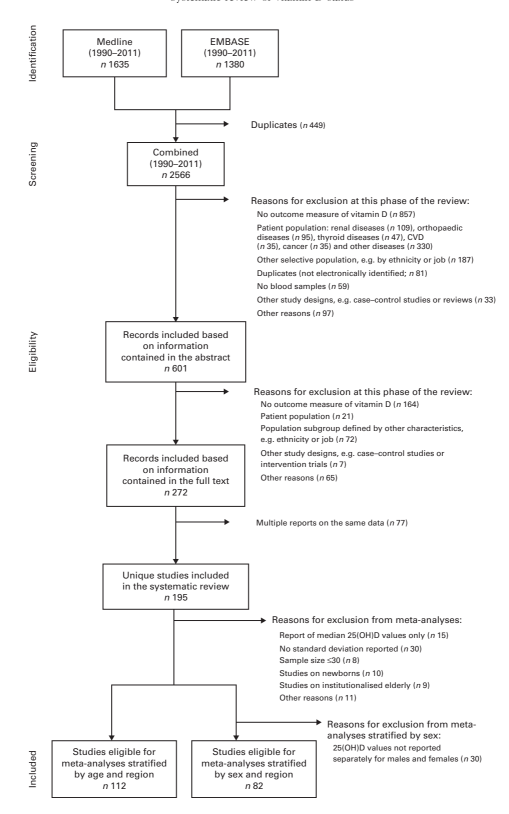


Fig. 1. Flow chart of the study selection (1990-2011). 25(OH)D, 25-Hydroxyvitamin D.

evaluated using information about the 25(OH)D measure (e.g. participation of the laboratory in the International Vitamin D Quality Assessment Scheme)⁽²⁰⁾. Finally, a study was classified as reliable if the intra- and inter-assay coefficients of variation

were below 10 and 15%, respectively. In instances where details about representativeness, validity or reliability were not provided, we created a separate category ('unknown') for each quality criterion.





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Table 1. Characteristics and main results from single studies on 25-hydroxyvitamin D (25(OH)D)*

Region and country	City/region within the country	Reference	n	Male (%)	Age group	Season	25(OH)D (nmol/l)	Reliability	Representativeness
Europe									
Austria									
71001110	Whole country	Koenig & Elmadfa ⁽³³⁾	1452	NA	0	NA	27.5	Unknown	Unknown
	Whole country	Kudlacek <i>et al.</i> ⁽³⁴⁾	1048	38.2	Ā	Winter	52.2	Unknown	No
Belgium	,,,,,,,,								
3 -	Brabant	Boonen et al. (35)	245	0.0	E	NA	56-4	Unknown	No
	Brussels	MacFarlane et al. (36)	126	31.0	Α	Winter	48.4	Unknown	No
	Brussels	Moreno-Reyes et al. (37)	401	50⋅1	Α	NA	35.0	Yes	No
	Northern Belgium	Richart et al. (38)	542	49.8	NA	NA	71.4†; 73.4‡	Unknown	Unknown
Czech Republic									
	Prague	Zofkova & Hill ⁽³⁹⁾	47	0.0	Α	NA	58-2	Unknown	No
Denmark									
	Copenhagen	Andersen et al. (40)	112	NA	C; E	Winter	24·4§; 47·8§	Yes	No
	Copenhagen	Brot et al. (41)	510	0.0	Α	NA	24.0§	Yes	No
	Faroe Islands	Dalgard <i>et al.</i> ⁽⁴²⁾	669	51.1	E	Mixed	47.6	Unknown	Unknown
	Odense	Frost et al. (43)	700	100.0	Α	Whole year	64.9	Unknown	No
	Aarhus	Rejnmark <i>et al.</i> ⁽⁴⁴⁾	315	0.0	Α	NA	57·0§	Unknown	No
	Aarhus	Rejnmark et al. (45)	2316	0.0	Α	Mixed	62·0§	Unknown	Yes
	Copenhagen	Rudnicki <i>et al.</i> ⁽⁴⁶⁾	125	42.4	Α	Whole year	25.5	Yes	Yes
Estonia									
	Vaike-Maarja	Kull <i>et al.</i> ⁽⁴⁷⁾	367	45.5	Α	Winter	43.7	Yes	Yes
Finland									
	Porvoo (region)	Andersen et al. (40)	120	NA	A; E	Winter	29·2§; 45·2§	Yes	No
	Whole country	Kauppi <i>et al.</i> ⁽⁴⁸⁾	6035	45.3	Α	NA	45.1†; 45.2‡	Yes	No
	Whole country	Lamberg-Allardt et al. (49)	328	38.4	Α	Mixed	45.0†; 47.0‡	Yes	Unknown
	Whole country	Matilla et al. (50)	4097	47.0	Α	Whole year	43.6	Yes	Unknown
	Whole country	Partti et al. (51)	6241	45.0	Α	Mixed	45.1	Unknown	No
	North Savo	Parviainen et al. (52)	776	53.9	Α	Mixed	34.0†; 35.0‡	Unknown	Unknown
	Turku	Piirainen et al (53)	82	NA	С	Mixed	54.7	Unknown	Unknown
	Helsinki	Viljakainen et al. (54)	64	0.0	С	Summer; winter	59.5; 37.3	Yes	Unknown
	Helsinki	Viljakainen <i>et al</i> . ⁽⁵⁵⁾	125	52.8	I	Winter	50⋅7	Yes	Unknown
France									
	Montpellier	Blain et al. (56)	248	0.0	Α	NA	64·1§	Yes	No
	Caen	Bougle et al. (57)	82	NA	I	NA	74.9	Unknown	No
	France	Chapuy et al. (58)	1569	48.8	Α	Winter	61.0	Yes	Unknown
	Burgundy	De Carvalho et al. (59)	164	42.7	Α	Whole year	74.4†; 52.8‡	Unknown	No
	Poitiers	Deplas et al. (60)	64	31⋅3	E	Spring	21.4	Unknown	No
	Whole country	Malvy <i>et al.</i> ⁽⁶¹⁾	1191	42.7	Α	Winter	79.5	Unknown	Unknown
Germany		(00)							
	Bonn	Braemswig et al. (62)	21	100-0	Α	Mixed	51.3	Unknown	Unknown
	Whole country	Hintzpeter et al. (63)	4030	43.7	0	NA	45·2§†; 44·7§‡	Yes	Yes
	Southern Germany	Scharla et al. (64)	415	50.4	A	Summer; winter	67.4; 42.4	Yes	Unknown
	Southern Germany	Woitge et al. (65)	41	36.6	0	Mixed	65-6	Unknown	No
_	Bonn	Zittermann et al. (66)	76	0.0	Α	Summer; winter	69-8; 30-3	Unknown	No
Greece		(67)					_		
	Athens	Nicolaidou <i>et al.</i> ⁽⁶⁷⁾	123	57.7	I_	Whole year	50-9§	Yes	Yes
	Athens	Papapetrou et al. (68)	279	17⋅2	E	Mixed	42.9	Unknown	No



Table 1. Continued

Region and ountry	City/region within the country	Reference	n	Male (%)	Age group	Season	25(OH)D (nmol/l)	Reliability	Representativenes
Iceland									
Toolaria	Reykjavik	Kristinsson et al. (69)	259	0.0	С	Winter	43.9	Yes	No
	Reykjavik	Sigurdsson et al. (70)	308	0.0	Ē	Mixed	53.1	Yes	NA
	Reykjavik	Steingrimsdottir <i>et al.</i> ⁽⁷¹⁾	944	52·0	Ā	Whole year	45.7	Yes	No
Ireland	rieykjavik	Stelligillisdottil et al.	344	32.0	^	vviiole year	45.7	165	INO
ITEIAITU	Cork (region)	Andersen et al. (40)	62	NA	C; E	Winter	41·3§; 43·7§	Yes	No
	Cork (region) Cork (city)	Hill et al. ⁽⁷²⁾	44	0.0	A A	Winter	54·5	Yes	Unknown
	Dublin	Keane <i>et al</i> ⁽⁷³⁾	116	NA	Ê	NA	37·1	Unknown	Unknown
Israel	Dubiiii	Realie et al	110	INA	_	INA	37.1	OTKHOWIT	OTIKHOWH
isiaei	Whole country	Oren et al. ⁽⁷⁴⁾	195	48.7	0	Whole year	57-2	Unknown	Yes
lant.	whole country	Oren et al.	195	40.7	U	vvriole year	57.2	Ulknown	res
Italy	Whole country	Adami et al. (75)	697	0.0	Е	Winter	37.9	Unknown	No
	Whole country	Carnevale et al. (76)	90	35.6		Winter	37·9 42·7		No
	Southern Italy	Romagnoli <i>et al.</i> ⁽⁷⁷⁾			A			Yes	
	Rome	Vezzoli <i>et al.</i> (78)	135	NA 50.0	A	Summer; winter	90.1; 45.9	Yes	No
NI atta antanada	Greve, Bagno a Ripoli	vezzoii <i>et ai.</i>	595	50.8	0	NA	61.2†; 48.2‡	Yes	Unknown
Netherlands	Dilah asasa di kasada	ALD -1-i4 -4(79)	05	40.0		NIA	04.04.77.04	I Indonesia	Halmanna
	Bilthoven, Utrecht	Al-Delaimy <i>et al.</i> ⁽⁷⁹⁾ Baynes <i>et al.</i> ⁽⁸⁰⁾	65	46.2	A	NA	91.2†; 77.2‡	Unknown	Unknown
	Zutphen	Baynes et al. (81)	142	100-0	E	Spring	42.0	Yes	No
	Rotterdam	Fang <i>et al.</i> ⁽⁸¹⁾	1317	NA	E	Whole year	65.5	Yes	No
	Whole country	Kuchuk et al. (82)	1319	48.7	E	Whole year	53-2	Yes	Yes
	Whole country	Löwik <i>et al.</i> ⁽⁸³⁾	529	50.7	E	NA	40.0†; 38.0‡	Unknown	No
	Hoorn	Pilz <i>et al.</i> ⁽⁸⁴⁾	614	NA	E	Whole year	56.5†; 50.8‡	Yes	No
	Amsterdam	Van Summeren et al. (85)	307	50.8	С	NA	69-6	Unknown	No
Norway		(20)							
	Skjervoy	Brustad et al.(86)	32	65-6	Α	NA	67-2	Unknown	No
	Northern Norway	Brustad et al. (87)	300	0.0	Α	Mixed	56.9	Yes	Unknown
	Tromso	Grimnes et al. (88)	6932	39.0	Α	NA	58.9	Yes	No
	Oslo	Meyer et al. (89)	869	42.8	Α	Mixed	74.8	No	No
Poland		442							
	Sadyba (Warsaw)	Andersen et al. (40)	126	NA	C; E	Winter	30·6§; 32·5§	Yes	No
	Warsaw	Napiorkowska et al. (90)	274	0.0	Е	Winter	33.7	Yes	Yes
Russia									
	NA	Sapir-Koren et al. (91)	122	0.0	E	NA	29.1	Unknown	No
Spain									
	Sabadell	Almirall <i>et al.</i> ⁽⁹²⁾	237	46.8	E	Winter	42.9	Unknown	No
	L'Hospitalet de Llobregat	Gomez et al. (93)	253	49.8	Α	Whole year	52.7†; 49.9‡	Unknown	Yes
	Betanzos	Moreiras <i>et al.</i> ⁽⁹⁴⁾	55	45.5	E	Spring	25.3	Unknown	Unknown
	Lleida	Muray et al. (95)	391	58.1	Α	Autumn	23.4†; 21.3‡	Unknown	No
	Murica	Perez-Llamas et al. (96)	86	33.7	Е	Mixed	50.1	Yes	Unknown
Sweden									
	Central Sweden	Burgaz <i>et al.</i> ⁽⁹⁷⁾	116	0.0	E	Winter	69.0	Yes	Unknown
	Uppsala, Västmanland	Burgaz <i>et al.</i> ⁽⁹⁸⁾	100	0.0	E	Winter	72.0	Unknown	No
	Malmo	Gerdhem <i>et al.</i> ⁽²⁸⁾	986	0.0	Ē	Whole year	95.0	Yes	No
	Uppsala	Hagström <i>et al.</i> ⁽⁹⁹⁾	958	100.0	Ē	NA	69.0	Unknown	Unknown
	Uppsala	Lind <i>et al.</i> ⁽¹⁰⁰⁾	34	100.0	Ā	NA	90.0	Unknown	No
	Stockholm	Melin et al. (101)	104	22.1	É	Spring	69.9†; 64.9‡	Yes	No
	Stockholm	Salminen et al. (102)	350	0.0	Ē	Whole year	91.0§	Yes	No

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Table 1. Continued

Region and country	City/region within the country	Reference	n	Male (%)	Age group	Season	25(OH)D (nmol/l)	Reliability	Representativeness
Switzerland									
	Vaud, Fribourg, Ticino	Burnand et al.(103)	3276	51.7	0	Mixed	50-0	Unknown	Yes
	Lausanne	Krieg <i>et al.</i> ⁽¹⁰⁴⁾	349	29.5	E	NA	26.5†; 23.2‡	Unknown	Unknown
	Basel	Theiler et al. (29)	505	57.4	E	Mixed	17·5† ; 18·2‡ ; 91·6†; 67·4‡	Yes	No
UK		(405)							
	Central, South, West England, Wales	Bates et al.(105)	924	NA	E	Mixed	51.9	Unknown	No
	East Kent	Carter et al. (106)	188	25.5	E	Mixed	31·2§	Unknown	No
	Northern Ireland	Cashman et al. (107)	1015	49.8	С	Mixed	61·1†§; 59·0‡§	Yes	Yes
	Great Britain	Davies et al.(108)	756	NA	С	Mixed	51.8	Unknown	Yes
	South England	Elia <i>et al.</i> ⁽¹⁰⁹⁾	1026	NA	E	NA	52.5	Unknown	No
	Isle of Ely	Forouhi <i>et al.</i> ⁽¹¹⁰⁾	524	40.8	Α	NA	60-2	Yes	Unknown
	Cambridge	Hegarty et al. (111)	96	49.0	E	Winter	23.1	Yes	Unknown
	Northern Ireland	Hill <i>et al.</i> ⁽¹¹²⁾	1015	49.8	С	Whole year	64.3	Yes	Yes
	England	Hirani & Primatesta ⁽¹¹³⁾	1297	40.3	Е	Whole year	40·0† ; 37·4‡ 58·3†; 49·4‡	Unknown	Yes
	Great Britain	Hypponen & Power ⁽¹¹⁴⁾	7437	50⋅1	Α	Summer; winter	60.3; 41.1	Yes	No
	Grampian	Macdonald et al. (115)	2905	0.0	Α	Mixed	53.9	Yes	No
	Aberdeen	Mavroeidi <i>et al.</i> ⁽¹¹⁶⁾	325	0.0	E	Mixed	53.3	No	No
	Isle of Ely	Wareham et al. (117)	1057	43.3	NA	Whole year	54.4†; 46.2‡	Yes	No
North America Canada	,					,	1, 1		
	Quebec	Barake et al.(118)	404	51.2	E	Mixed	74.0	Yes	No
	Nunavut	El Havek <i>et al.</i> ⁽¹¹⁹⁾	282	46.8	Ċ	Mixed	48-3§	No	Yes
	Whole country	Langlois <i>et al.</i> ⁽¹²⁰⁾	5306	48-4	O	Whole year	67.7	Yes	Yes
	St Theresa Point, Garden Hill	Lebrun et al. (121)	76	NA	Ī	Summer	26-2	Unknown	Unknown
	Toronto	Liu <i>et al.</i> ⁽¹²²⁾	155	49.7	Ē	Autumn	44.9	Unknown	Unknown
	Quebec	Mark <i>et al.</i> ⁽¹²³⁾	1753	50.3	C	Mixed	46.0	Yes	No
	Avalon Peninsula	Newhook et al. (124)	51	NA	Ĭ	Summer: winter	63-6; 48-6	Unknown	No
	Edmonton	Overton & Basu ⁽¹²⁵⁾	36	100-0	Ē	Summer	122.0	Unknown	No
	Calgary	Rucker et al. (126)	188	31.9	Ē	Winter	57.3	No	No
	Quebec	Sinotte et al. (127)	741	0.0	Ā	Winter	64.9	Yes	No
USA	440200	Cilibria of all			**		0.0	. 55	
00/.	NA	Alvarez <i>et al.</i> ⁽¹²⁸⁾	50	0.0	Α	Mixed	55-7	Unknown	No
	New York	Arunabh <i>et al.</i> ⁽¹²⁹⁾	410	0.0	A	Whole year	54.2	Yes	No
	Connecticut	Avery et al. (130)	114	NA	E	NA	113.1; 81.8	Yes	No
	Honolulu	Chai <i>et al</i> ⁽¹³¹⁾	182	0.0	Ā	NA	72.3	Unknown	Unknown
	Framingham	Cheng et al. (132)	3890	46.0	A	Whole year	92.9	No	No
	Boston	Dawson-Hughes et al. (133)	391	46.5	Ē	Whole year	82.4†; 68.9‡	Yes	Unknown
	Oakland	Dror et al. (134)	199	NA	Ī	Mixed	43.7	Unknown	Unknown
	Whole country	Looker et al. (135)	18462	47·2	0	Summer, winter	77.3; 67.2	No	Yes
	Framingham	Hannan <i>et al.</i> ⁽¹³⁶⁾	341	NA	Ē	NA	71.9	Yes	No
	Boston, Houston, West Lafayette	Hill et al. (137)	735	30.5	C	NA	66-2	Unknown	Unknown
	Whole country	lannuzzi-Sucich et al. (138)	337	42.1	E	NA	67.4†; 57.7‡	Yes	No
	Connecticut	llich <i>et al.</i> (139)	136	0.0	Ē	Whole year	52.8	Unknown	No
	Framingham	Jagues <i>et al.</i> ⁽¹⁴⁰⁾	759	38.2	Ē	NA	82.0†; 71.0‡	Yes	Unknown
	Northern Georgia	Johnson et al. (141)	317	20.2	Ē	Whole year	66.7	Yes	Unknown
	Rochester	Khosla <i>et al.</i> ⁽¹⁴²⁾	138	0.0	Ā	NA	77·6	Unknown	Unknown
	Bochester								



Table 1. Continued

California Eastern Nebraska Lappe et al. 145 1179 0.0 E Whole year Whole country Mansbach et al. 146 4558 49-6 C Whole year Whole country Mansbach et al. 146 4558 49-6 C Whole year Whole Country Mansbach et al. 146 66 46 46 E NA NA Parmington Parmingt	25(OH)D (nmol/l)	Reliability	Representativeness
Eastern Nebraska Lappe et al. (1465) 1179 0-0 E Whole year Whole country Mansbach et al. (1469) 4558 49-6 C Whole year Whole year Mirza et al. (1479) 40 0-0 A; E NA NA Marion Country Rock et al. (1490) 1042 39-4 O Mixed Mirza et al. (1490) 1042 39-4 O Mixed Mixed Greenwich Sabetta et al. (1490) 198 42-9 O Autumn Authors Sabetta et al. (1593) 1381 48-4 A NA NA Authors Stein et al. (1593) 1381 48-4 A NA NA Authors Stein et al. (1593) 122 0-0 C Summer Philadelphia Weng et al. (1593) 382 47-6 C Whole year Autumn Mixed Mi	75.1	Unknown	No
Whole country	71·8	Yes	No
Farmington Mirza et al. (147) 40 0.0 A; E NA	68.0	Unknown	Yes
Rancho Bernardo Reis et al. (1489) 1042 39-4 0 Mixed	74.9; 84.9	Yes	No
Marion County Rock et al. (149) 1042 39.4 0 Mixed Greenwich Sabetta et al. (150) 198 42.9 0 Autumn Framingham Shea et al. (150) 1381 48.4 A NA NA Athens Stein et al. (152) 168 0.0 C Whole year Sala/Pacific Weng et al. (153) 382 47.6 C Whole year Sala/Pacific Sydney Bowyer et al. (154) 382 47.6 C Whole year Sydney Brock et al. (156) 186 NA E NA NA E NA Dubbo Center et al. (157) 437 100.0 E NA NA Mixed	103.6	Yes	No
Greenwich Sabetta et al. (159) 198 42-9 0	31.9†; 29.3‡	Yes	Yes
Framingham	70.9	Unknown	Unknown
Athens	70.9 49.4	Unknown	No
Bangor Philadelphia Weng et al. (154) 382 47.6 C Whole year			
Philadelphia Weng et al. 154) 382 47-6 C Whole year 154 Sydney Sydney Brock et al. 155 186 NA E NA NA NA NA NA NA	93.8	Yes	No
Stal/Pacific Sydney	74.4	Yes	Unknown
Australia	69·9§	Yes	Yes
Sydney			
Sydney			
Dubbo Center et al. (157) 437 100-0 E NA Tasmania Ding et al. (159) 1002 NA A Mixed North-Western Adelaide Ngo et al. (159) 253 43-5 E NA Barwon Pasco et al. (160) 861 0-0 A Whole year Melbourne Stein et al. (160) 99 26-3 E Winter Sydney Zochling et al. (162) 584 21-2 E Mixed	60∙0§	Unknown	No
Tasmania Ding et al. (158) 1002 NA A Mixed	36.0; 33.0	Yes	No
North-Western Adelaide Ngo et al. (199) 253 43.5 E NA	70.7	Yes	No
North-Western Adelaide Ngo et al. (199) 253 43.5 E NA	52.8	Yes	Unknown
Melbourne Stein et al. (161) 99 26.3 E Mixed	72.2	Yes	No
Melbourne Stein et al.	70.0	Yes	No
China Chin	26·0§	Yes	No
China	21.4†; 16.9‡	Unknown	No
Linxian Abnet et al. (163) 720 42-2 A Spring Hong Kong Chan et al. (164) 53 0-0 E NA Linxian Chen et al. (165) 2018 54-0 A Spring Beijing Du et al. (166) 649 0-0 C Winter Shanxi Strand et al. (167) 250 52-4 C Spring Taipei Tsai et al. (168) 262 0-0 A Mixed Fiji Islands Myhole country Heere et al. (168) 511 0-0 A Winter India Agota Goswami et al. (170) 57 56-1 A Winter Tirupati Harinarayan et al. (171) 1146 21-2 A NA Lucknow Sachan et al. (172) 117 NA I Mixed Indonesia Jakarta, Bekasi Rinaldi et al. (173) 62 0-0 E Summer Japan Jakarta, Bekasi Setiati et al. (174) 74 0-0 E NA Japan NA Kuwabra et al. (176) 1094 41-7 E Winter Toyosaka Nakamura et al. (177) 160 0-0 E Summer Toyosaka Nakamura et al. (178) 1094 41-7 E Winter Toyosaka Nakamura et al. (178) 1094 41-7 E Winter Toyosaka Nakamura et al. (178) 117 0-0 E Summer Toyosaka Nakamura et al. (178) 117 0-0 E Summer Toyosaka Nakamura et al. (178) 117 0-0 E Summer Malaysia Kuala Lumpur Rahman et al. (180) 101 0-0 A NA		•	
Hong Kong	33.1	Yes	Unknown
Linxian Chen et al. (165) 2018 54-0 A Spring Beijing Du et al. (166) 649 0-0 C Winter Shanxi Strand et al. (167) 250 52-4 C Spring Taipei Tsai et al. (168) 262 0-0 A Mixed Fiji Islands Whole country Heere et al. (169) 511 0-0 A Winter India Agota Goswami et al. (170) 57 56-1 A Winter Tirupati Harinarayan et al. (171) 1146 21-2 A NA Lucknow Sachan et al. (172) 117 NA I Mixed Indonesia Jakarta, Bekasi Rinaldi et al. (173) 62 0-0 E Summer Jakarta, Bekasi Setiati et al. (174) 74 0-0 E NA Japan NA Kuwabra et al. (175) 50 30-0 E NA Tokyo Kwon et al. (176) 1094 41-7 E Winter Toyosaka Nakamura et al. (1778) 160 0-0 E Summer Toyosaka Nakamura et al. (178) 117 0-0 E Summer Tokyo Suzuki et al. (178) 117 0-0 E Summer Tokyo Suzuki et al. (178) 117 0-0 E Summer Tokyo Suzuki et al. (178) 117 0-0 E Summer Tokyo Suzuki et al. (178) 117 0-0 E Summer Tokyo Suzuki et al. (178) 117 0-0 E Summer Tokyo Suzuki et al. (178) 117 0-0 E Summer Tokyo Suzuki et al. (178) 117 0-0 E Summer Tokyo Suzuki et al. (178) 117 0-0 E Summer Tokyo Suzuki et al. (178) 117 0-0 E Summer Tokyo Suzuki et al. (179) 2957 32-1 E Autumn	57.7	Unknown	No
Beijing Strand et al. (166) 649 0.0 C Winter	31.7	Unknown	Unknown
Shanxi Taipei Tsai et al. (168) 250 52-4 C Spring	33.5	Yes	Yes
Fiji Islands Whole country Heere et al. (169) Agota	42.3†; 25.5‡	Unknown	Unknown
Fiji Islands Whole country Heere et al. (169) Agota Agota Agota Agota Agota Alirinarayan et al. (1710) Lucknow Sachan et al. (1711) Indonesia Indonesia Jakarta, Bekasi Jakarta, Bekasi Setiati et al. (1714) NA Indonesia NA Kuwabra et al. (1714) Tokyo Kwon et al. (175) Tokyo Kwon et al. (176) Tokyosaka Nakamura et al. (1776) Nakamura et al. (1776) Tokyo Tokyosaka Nakamura et al. (1777) Nakamura et al. (1780) Nakamura et al. (1778) Nakamura et al. (1778) Nakamura et al. (1779) Nakamura et al. (1780) Nakamura et al. (1780	• • •		
Myole country	76-6	Yes	No
India			
Agota Goswami et al. 170 57 56-1 A Winter	76.0	Unknown	Unknown
Tirupati			
Lucknow Sachan et al. (172) 117 NA I Mixed	36-4	Unknown	Unknown
Indonesia	46.3†; 38.7‡	Unknown	No
Jakarta, Bekasi Rinaldi et al. (173) 62 0.0 E NA	21.0	Yes	No
Jakarta, Bekasi Setiati et al.(174) 74 0.0 E NA Japan NA Kuwabra et al.(175) 50 30.0 E NA Tokyo Kwon et al.(176) 1094 41.7 E Winter Toyosaka Nakamura et al.(177) 160 0.0 E Summer Toyosaka Nakamura et al.(178) 117 0.0 E Summer Tokyo Suzuki et al.(179) 2957 32.1 E Autumn Malaysia Kuala Lumpur Rahman et al.(180) 101 0.0 A NA			
Japan NA Kuwabra et al. (175) 50 30·0 E NA Tokyo Kwon et al. (176) 1094 41·7 E Winter Toyosaka Nakamura et al. (177) 160 0·0 E Summer Toyosaka Nakamura et al. (178) 117 0·0 E Summer Tokyo Suzuki et al. (179) 2957 32·1 E Autumn Malaysia Kuala Lumpur Rahman et al. (180) 101 0·0 A NA	68-2	Unknown	Unknown
Japan NA Kuwabra et al. (175) 50 30·0 E NA Tokyo Kwon et al. (178) 1094 41·7 E Winter Toyosaka Nakamura et al. (177) 160 0·0 E Summer Toyosaka Nakamura et al. (178) 117 0·0 E Summer Tokyo Suzuki et al. (179) 2957 32·1 E Autumn Malaysia Kuala Lumpur Rahman et al. (180) 101 0·0 A NA	38.7	No	Yes
NA Kuwabra <i>et al.</i> ⁽¹⁷⁵⁾ 50 30·0 E NA Tokyo Kwon <i>et al.</i> ⁽¹⁷⁶⁾ 1094 41·7 E Winter Toyosaka Nakamura <i>et al.</i> ⁽¹⁷⁷⁾ 160 0·0 E Summer Toyosaka Nakamura <i>et al.</i> ⁽¹⁷⁸⁾ 117 0·0 E Summer Tokyo Suzuki <i>et al.</i> ⁽¹⁷⁹⁾ 2957 32·1 E Autumn Malaysia Kuala Lumpur Rahman <i>et al.</i> ⁽¹⁸⁰⁾ 101 0·0 A NA			
Tokyo Kwon et al. (176) 1094 41·7 E Winter Toyosaka Nakamura et al. (177) 160 0·0 E Summer Toyosaka Nakamura et al. (178) 117 0·0 E Summer Tokyo Suzuki et al. (179) 2957 32·1 E Autumn Malaysia Kuala Lumpur Rahman et al. (180) 101 0·0 A NA	27.7§	Unknown	Unknown
Toyosaka Nakamura et al. ⁽¹⁷⁷⁾ 160 0.0 E Summer Toyosaka Nakamura et al. ⁽¹⁷⁸⁾ 117 0.0 E Summer Tokyo Suzuki et al. ⁽¹⁷⁹⁾ 2957 32·1 E Autumn Malaysia Kuala Lumpur Rahman et al. ⁽¹⁸⁰⁾ 101 0·0 A NA	71.7†; 65.8‡	Unknown	No
Toyosaka Nakamura <i>et al.</i> ⁽¹⁷⁸⁾ 117 0·0 E Summer Tokyo Suzuki <i>et al.</i> ⁽¹⁷⁹⁾ 2957 32·1 E Autumn Malaysia Kuala Lumpur Rahman <i>et al.</i> ⁽¹⁸⁰⁾ 101 0·0 A NA	78.3	Yes	No
Tokyo Suzuki <i>et al.</i> ⁽¹⁷⁹⁾ 2957 32·1 E Autumn Malaysia Kuala Lumpur Rahman <i>et al.</i> ⁽¹⁸⁰⁾ 101 0·0 A NA	59·1	Yes	Yes
Malaysia Kuala Lumpur Rahman <i>et al.</i> ⁽¹⁸⁰⁾ 101 0·0 A NA	71.1†; 60.4‡	Unknown	No
. Kuala Lumpur Rahman <i>et al.</i> ⁽¹⁸⁰⁾ 101 0⋅0 A NA	7111,004	CHRIDWII	140
	44-4	Yes	No
	44.4	168	INO
Mongolia Ulaanbaatar Lander <i>et al.</i> ⁽¹⁸¹⁾ 98 72-4 C Autumn	24.1	Yes	No



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Table 1. Continued

Region and country	City/region within the country	Reference	n	Male (%)	Age group	Season	25(OH)D (nmol/l)	Reliability	Representativenes
	the country	Tiererenee		Wale (70)	rigo group	Ocuson	(1111071)	Tichability	1 topresentativenes
New Zealand		(183)							
	Auckland	Bolland <i>et al.</i> ⁽¹⁸²⁾	1984	19∙1	A; E	NA	84.0†; 51.0‡	Yes	No
	Auckland	Bolland et al. (183)	116	0.0	Α	NA	54.0	Unknown	Unknown
	Auckland	Bolland et al. (184)	100	50∙0	A; E	NA	91.0†; 51.0‡	Yes	No
	Wellington; Christchurch	Camargo et al. (185)	922	50.7	I	Whole year	44·0§	Yes	Unknown
	Auckland	Grant et al. (186)	353	47.6	I	Whole year	55.0	Yes	Unknown
	Dunedin	Houghton et al. (187)	193	57∙5	С	Mixed	52.0	Yes	Unknown
	Auckland	Ley <i>et al</i> . ⁽¹⁸⁸⁾	39	0.0	E	Winter	26-1	Unknown	No
	Auckland	Lucas <i>et al.</i> ⁽¹⁸⁹⁾	1606	0.0	E	Whole year	51.2	Unknown	No
	Whole country	Rockell et al. (190)	1585	50.5	С	Mixed	50.0	Yes	No
	Dunedin; Invercargill	Rockell et al. (191)	342	34.8	Α	Summer	85.0	Unknown	Unknown
	Auckland	Scragg et al. (192)	295	100.0	Α	Whole year	39.8	No	Yes
South Korea						,			
	Chungju	Kim <i>et al.</i> ⁽¹⁹³⁾	1330	38.0	E	Whole year	46-1	Unknown	No
	Seoul	Namgung et al. (194)	71	50.7	ı	Summer; winter	74.9; 26.7	Yes	Unknown
Thailand					-		,		
· · · · · · · · · · · · · · · · · · ·	NA	Chailurkit <i>et al.</i> ⁽¹⁹⁵⁾	158	48.7	0	NA	168-2†; 105-8‡	Unknown	Unknown
	Khon Kaen	Chailurkit <i>et al.</i> (196)	251	50.2	Ö	NA	128-3†; 93-6‡	No	Yes
	Bangkok	Chailurkit <i>et al.</i> ⁽¹⁹⁷⁾	229	47·2	Ö	NA	135.0†; 72.6‡	No	Unknown
	Bangkok	Chailurkit <i>et al.</i> (26)	446	0.0	Ē	NA	67.6	Yes	Unknown
	Khon Kaen	Soontrapa <i>et al.</i> ⁽¹⁹⁸⁾	65	0.0	Ē	Summer	83.2	No	Unknown
Vietnam	Kiloli Kaeli	Soontiapa et al.	03	0.0	L	Summer	00.2	NO	OHKHOWH
	Ho Chi Minh (city)	Ho-Pham et al. (199)	637	32.2	Α	Mixed	91.9†; 75.1‡	Yes	Yes
Middle East/ Africa									
Cameroon									
	Ntam	Njemini <i>et al.</i> ⁽²⁰⁰⁾	152	60.5	Е	NA	52.7	Unknown	No
Iran	· · · · · · · · · · · · · · · · · · ·	rijoriii ot a	102	000	_	101	02 /	Omarown	110
ii di i	Tehran	Bassir et al. (201)	44	NA	ı	Mixed	4.9	Unknown	Unknown
	Tehran	Dahifar <i>et al.</i> (202)	414	0.0	C	Mixed	74.9	Unknown	Unknown
	Tehran	Hashemipour <i>et al.</i> (203)	1210	59.1	Ö	NA	20·7§	Yes	No
	Tehran	Hossein-Nezhad <i>et al.</i> (204)	646	24.8	A	NA	31.3	Yes	Unknown
	Tehran	Hosseinpanah <i>et al.</i>	245	0.0	Ä	NA	73.0	Yes	Yes
	Zanjan	Kazemi <i>et al.</i>	61	NA	1	Mixed	16·7	Unknown	Unknown
	Shiraz	Masoompour <i>et al.</i> (207)			^		35.0		
		Masoompour et al. (208)	520	100.0	A	Winter		Yes	Yes
	Tehran	Mirsaeid Ghazi <i>et al.</i> ⁽²⁰⁸⁾	1171	41.8	0	Mixed	87.4†; 52.4‡	Yes	No
	Isfahan	Moussavi <i>et al.</i> ⁽²⁰⁹⁾	318	48.1	C	Winter	93.1†; 41.8‡	Yes	No
	Tabriz	Niafar <i>et al.</i> ⁽²¹⁰⁾	300	0.0	A	Mixed	35·4§	Yes	Unknown
	Tehran	Rabbani <i>et al.</i> ⁽²¹¹⁾	963	44.0	С	Winter	116.1†; 60.3‡	Yes	No
	Isfahan	Salek et al. (212)	88	NA	I	Summer	68-4	Yes	Unknown
Jordan		01 11 6 5 1 (00)		·					
	Northern Jordan	Gharaibeh & Stoecker ⁽²²⁾	186	27.4	Α	Summer	25.6	Unknown	Unknown
Lebanon		(010)							
	NA	Arabi <i>et al.</i> ⁽²¹³⁾	443	64.6	E	Spring	28.5	Unknown	Unknown
	Beirut, Bekaa	Gannage-Yared et al. (214)	316	31.3	Α	Winter	24.2	Yes	No





Table 1. Continued

Region and country	City/region within the country	Reference	и	Male (%)	Male (%) Age group Season	Season	25(OH)D (nmol/l)	Reliability	Representativeness
Nigeria	sop	Pfitzner <i>et al.</i> ⁽²¹⁵⁾	218	45.0	O	Mixed	8.99	Unknown	Unknown
South Africa	Cape Town	Charlton et al. ⁽²¹⁶⁾	173	48.0	ш		36.9	Unknown	No
Gambia Latin America	Whole country	Aspray <i>et al.</i> ⁽²¹⁷⁾	113	0.0	0	Ϋ́	27.7	o N	No
Argentina	Ushuaia	Oliveri <i>et al.</i> ⁽²¹⁸⁾	42	57.1	O	Winter	24.5	Unknown	No
Diazii	Sao Paulo Sao Paulo	Canto-Costa <i>et al.</i> ⁽²¹⁹⁾ Saraiva <i>et al.</i> ⁽²²⁰⁾	11 250	36.4 30.8	шш	NA Whole year	61.2 52.4	Yes	No Yes

In some cases, 25(OH)D , مصمة المانية عالياته المانية المانية المانية المانية المانية المانية المانية عالياته المانية الماني NA, not available; O, others; A, adults; E, elderly; C, children and adolescents; I, newborns/infants.

Data from three studies not indicating geographical region have been excluded^(221–223); data from a single study⁽⁴⁰⁾ providing country-specific many values were available by age, sex or region only. For some studies multiple regions are available by age, sex or region only. For some studies multiple regions. 125(OH)D mean values for men. 125(OH)D mean values for women. 325(OH)D median values. 125(OH)D mean values for institutionalised elderly.

Statistical analyses

Descriptive statistics were calculated for baseline characteristics of all the included studies. If mean 25(OH)D values were not reported in an article, we used median values (9.2% of the studies) in our descriptive analyses.

Meta-analyses were performed for subgroups stratified by age, sex and geographical region using random-effects models. Studies reporting median 25(OH)D values (n 15) or mean values without a corresponding standard deviation (n 30) were not included in this phase of the analyses (Fig. 1). In addition, our focus in the meta-analyses was limited to studies/subgroups with sample sizes greater than 30, given concerns about the precision of estimates. Studies on newborns $(n \ 10)$ and institutionalised elderly $(n \ 9)$ were also not included in the meta-analyses. For analyses stratified by sex, we also excluded studies that did not report separate 25(OH)D values for males and females (n 30).

Heterogeneity between the studies was assessed by visual inspection of forest plots and calculation of I^2 statistics. Because we found substantial heterogeneity across the studies, we decided to further explore potential explanatory factors. Therefore, we conducted heterogeneity analyses within each subgroup by accounting for a range of characteristics other than age and sex, which included season, assay type, distance from the equator⁽⁵⁾ and components of study quality. Studies were grouped by study characteristics (e.g. season and assay type) to assess whether heterogeneity was reduced as indicated by the I^2 statistics and the inspection of forest plots.

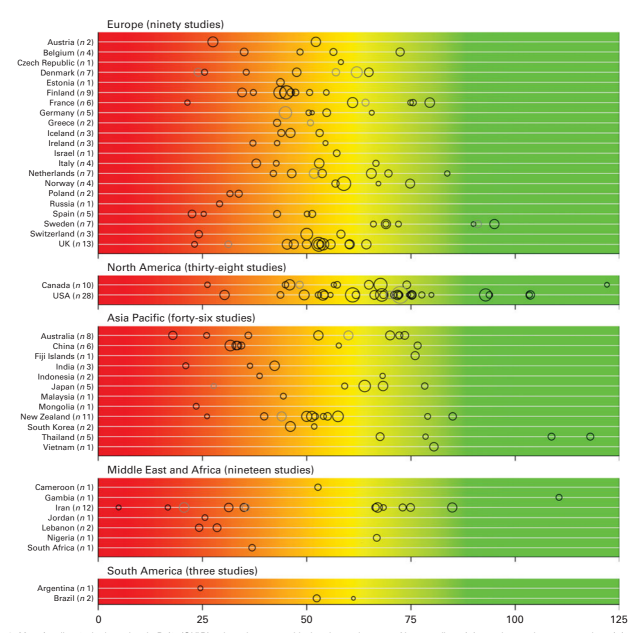
Supplementary analyses explored patterns of vitamin D status within specific subgroups (e.g. institutionalised elderly) and for selected associations reported in previous work. The purpose of these exploratory analyses was to support further research in this area by generating hypotheses that might be tested more thoroughly in future studies. All statistical analyses were conducted using STATA version 12.1 (StataCorp).

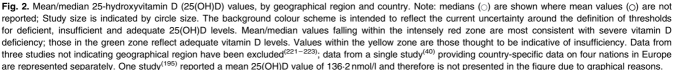
Results

Description of studies

Studies included in the present review (Table 1) contained data on a total of 168 389 participants from forty-four countries. The sample size of individual studies ranged from 11 to 18 462 participants with a median of 316 (interquartile range 117-861). While the majority of studies contained data on males and females, nine studies (4.7%) restricted their focus to males, while fifty-four studies (28·0%) contained data on only females. The overall proportions of males and females were 33.3 and 66.7%, respectively, and the mean age of the participants was 51.7 (sp 24.3) years. Most studies were conducted in Europe (45·1%), followed by the Asia/Pacific region (23.8%) and North America (19.7%). In terms of the country in which studies were conducted, most were carried out in the USA (n 28), followed by Iran (n 12), New Zealand (n 11) and Canada $(n\ 10)$.

The assays reported to measure 25(OH)D values included RIA (55.9%), competitive protein-binding assays (14.0%) and other methods such as chemiluminescence immunoassay and HPLC.





In terms of study quality, more than half of the studies (50·2%) were classified as non-representative of the target population and 14·9% qualified as representative according to the criteria defined previously. Evidence of representativeness could not be established in 34·9% of the studies due to missing information. Information on assay reliability was provided in 61·0% of the studies with 52·8% classified as providing reliable 25(OH)D measurements. Assay validity was reported in a minority of studies (9·7%).

Global vitamin D status

There was a significant variability in the estimates of 25(OH)D values across the studies with mean and median values ranging from 4·9 to 136·2 nmol/l and 20·7 to 91·0 nmol/l, respectively. We found that 88·1% of the samples presented in the present review had mean 25(OH)D values below 75 nmol/l, 37·3% had mean values below 50 nmol/l and 6·7% had mean values below 25 nmol/l. Fig. 2 provides an overview





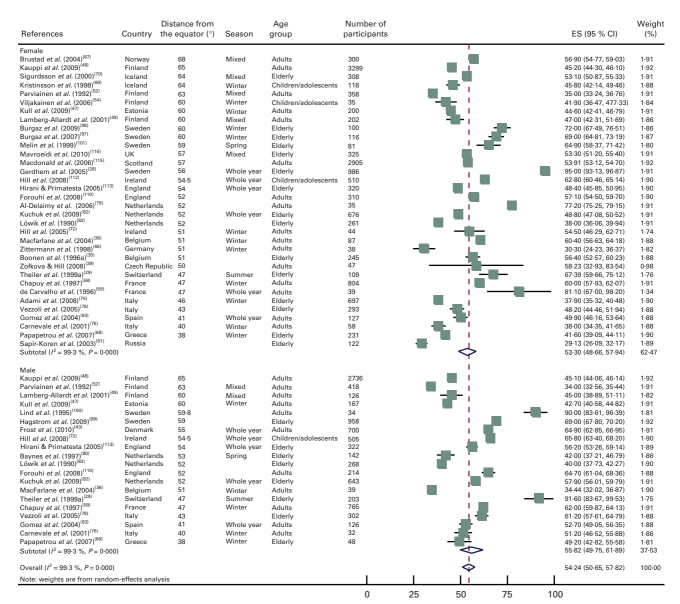


Fig. 3. Forest plot for Europe stratified by sex. ES, effect estimator. (A colour version of this figure can be found online at http://www.journals.cambridge.org/bjn)

of the distribution of country- and study-specific mean 25(OH)D values, stratified by region. In addition, a visualisation of the available data on a global map can be found elsewhere⁽²¹⁾.

Vitamin D status by age, sex and region

Due to a limited number of studies being identified from Latin America, it was not possible to perform meta-analyses for this region. Depending on the stratifying variable, I^2 values ranged from 84·5 to 99·7%, indicating substantial heterogeneity between the studies.

No significant age- or sex-related differences in 25(OH)D values were observed in the sample of eligible studies world-wide (data not shown). However, we observed differences by region with values being significantly higher in North America than in Europe or the Middle East/Africa region (Figs. 3–6). In an analysis stratified by age and region, we

did not find age-related differences for Europe and North America (Table 2). However, in the Asia/Pacific region, children/adolescents were found to have significantly lower 25(OH)D values than adults and elderly. In contrast, children/adolescents from the Middle East/Africa region had significantly higher values than the other two age groups. No significant sex-related differences were observed in any of the regions (Figs. 3–6). However, reports of 25(OH)D values in women tended to be lower, especially in the Asia/Pacific and Middle East/Africa regions.

Heterogeneity analyses

The substantial heterogeneity that we observed within the different geographical regions could not be explained by the characteristics of the study population or features of study quality. Grouping studies by age category and sex, assay type,



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References	Country	Distance from the equator (°)	Season	Age group	Number of participants		ES (95 % CI)	Weight (%)
Female						I I		
Sinotte et al. 2009 ⁽¹²⁷⁾	Canada	54	Winter	Adults	741	■i	64-90 (63-49, 66-31)	4.76
Khosla et al. 1997 ⁽¹⁴²⁾	USA	43		Adults	138	! - ■-	77-60 (71-74, 83-46)	4.48
Dawson-Hughes et al. 1997 ⁽¹³³⁾	USA	42	Whole year	Elderly	209		68-90 (64-55, 73-25)	4.61
Shea et al. 2009 ⁽²⁹⁰⁾	USA	42		Elderly	919		49-20 (47-95, 50-45)	4.76
Jaques et al. 1997 ⁽¹⁴⁰⁾	USA	42		Elderly	469	-	71.00 (68.38, 73.62)	4.71
Ilich et al. 2003 ⁽¹³⁹⁾	USA	41	Whole year	Adults	136		52.80 (50.65, 54.95)	4.73
Lappe <i>et al.</i> 2006 ⁽¹⁴⁵⁾	USA	41	Whole year	Elderly	1179		71-80 (70-64, 72-96)	4.76
lannuzzi-Sucich et al. 2002 ⁽¹³⁸⁾	USA	40		Elderly	195	- i	57-66 (54-82, 60-50)	4.70
Arunabh et al. 2003 ⁽¹²⁹⁾	USA	40	Whole year	Adults	410	■ 1	54-20 (50-84, 57-56)	4.67
Kremer et al. 2009 ⁽¹⁴⁴⁾	USA	37	Summer	Children/adolescents	90	 -	75-13 (68-43, 81-83)	4.40
Hill et al. (2010)(137)	USA	36		Children/adolescents	511		66-20 (63-85, 68-55)	4.72
Stein et al. (2006) ⁽¹⁵²⁾	USA	34	Whole year	Children/adolescents	168	i -	93.80 (89.55, 98.05)	4.62
Johnson et al. (2008)(141)	USA	32.5	Whole year	Elderly	200		67-90 (63-04, 72-76)	4.57
Chai et al (2010)(131)	Hawaii	21	,	Adults	182	·	72.30 (68.45, 76.15)	4.64
Alvarez et al. (2010)(128)	USA		Mixed	Adults	50	— ■ — i	55-66 (46-18, 65-14)	4.08
Subtotal ($I^2 = 98.8 \%$, $P = 0.000$)						$\stackrel{-}{\Leftrightarrow}$	66-57 (60-94, 72-20)	69-23
Male						1		
Overton & Basu (1999) ⁽¹²⁵⁾	Canada	53	Summer	Elderly	36	1	> 122·00 (106·32, 137·	-683-26
Jaques et al. (1997) ⁽¹⁴⁰⁾	USA	42		Elderly	290	-	82.00 (78.66, 85.34)	4.68
Shea et al. (2009)(290)	USA	42		Elderly	843		49.00 (47.81, 50.19)	4.78
Dawson-Hughes <i>et al.</i> (1997) ⁽¹³³⁾	USA	42	Whole year	Elderly	182	i - -	82-40 (77-20, 87-60)	4.54
Iannuzzi-Sucich et al. (2002)(138)	USA	40		Elderly	142	- III -	67-39 (64-03, 70-76)	4.67
Hill et al. (2010)(137)	USA	36		Children/adolescents	224	-	65.70 (62.83, 68.57)	4.70
Johnson et al. (2008)(141)	USA	32.5	Whole year	Elderly	37		60.50 (51.70, 69.30)	4.16
Subtotal ($I^2 = 99.0 \%$, $P = 0.000$)							74-44 (61-65, 87-24)	30.77
Overall ($I^2 = 98.9 \%$, $P = 0.000$)						♦	68-73 (63-71, 73-75)	100-00
Note: weights are from random-e	ffects analy	sis		1	1	1	1	

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Fig. 4. Forest plot for North America stratified by sex. ES, effect estimator. (A colour version of this figure can be found online at http://www.journals.cambridge.org/bjn)

season, distance from the equator or representativeness, for example, did not significantly reduce heterogeneity across the studies in our sample, as measured by the I^2 statistics.

Exploratory analyses

We found that mean 25(OH)D values for institutionalised elderly were lower than those for non-institutionalised elderly, especially in Europe and the Asia/Pacific region. Moreover, in specific subgroups in single countries within Europe, we observed differences, with Swedish elderly having higher 25(OH)D mean values than the elderly in other European countries. In addition, we found that newborns had lower 25(OH)D values than the other three age groups in several countries worldwide.

Discussion

Summary of the main findings

The published evidence on vitamin D status at the population level, as assessed by mean or median 25(OH)D values, is characterised by a high degree of variability across studies, countries and regions. Although no age- or sex-related significant differences in 25(OH)D values were observed across the sample of studies that we reviewed, we did observe differences by region with values being significantly higher in North America than in Europe or the Middle East/Africa region. In stratified analyses, significant age-related differences were observed in the Asia/Pacific and Middle East/ Africa regions, but not elsewhere. However, exploratory analyses suggested that newborns and institutionalised elderly were more likely to have lower reported 25(OH)D values in several regions worldwide. We found substantial heterogeneity between the studies in our sample from each geographical region that could not be explained in a detailed analysis.

Interpretation and comparison with previous studies

In contrast to previous reviews (5,13,14), we could not find differences in 25(OH)D values for children/adolescents, adults and elderly. However, in analyses stratified by geographical region, significant age-related differences could be observed for the Asia/Pacific region, with children/adolescents having lower 25(OH)D values than older groups. This might be primarily due to the low 25(OH)D values found for Chinese children/adolescents as reported in previous work⁽¹³⁾, who were observed to have low dietary Ca intake and limited sunlight exposure as possible reasons. In contrast, in the Middle East/Africa region, children/adolescents were found to have significantly higher 25(OH)D values than adults and elderly, a finding consistent with at least one previous study(8). One





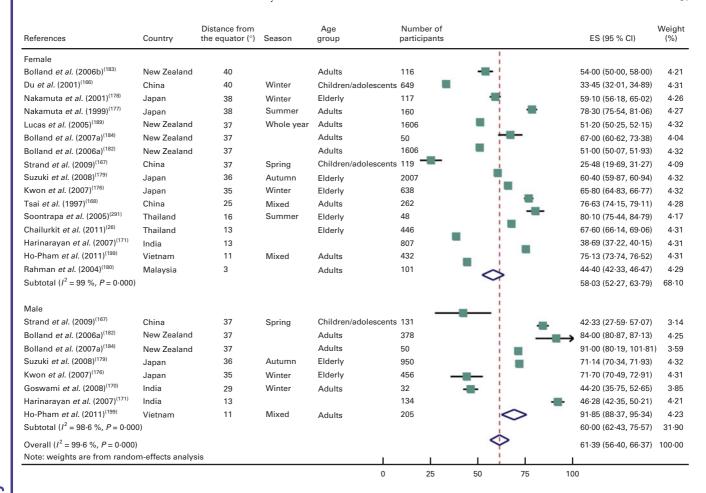


Fig. 5. Forest plot for the Asia/Pacific region stratified by sex. ES, effect estimator. (A colour version of this figure can be found online at http://www.journals. cambridge.org/bjn)

potential explanation for this pattern in the Middle East/Africa region could be that children/adolescents from this region generally spend more time outdoors compared with the other age groups (e.g. indoor working by the adult population) (22). However, others have also found age-related differences in other regions^(5,13,14), which could not be confirmed in the present meta-analyses. A reduction in differences and thus greater similarities across age groups might be attributable to lifestyle changes over the course of time in which younger individuals from industrialised countries spend more time indoors watching television, using computers and playing video games compared with older adults (23).

In contrast to previous reviews, we were also unable to find significant sex-related differences (8,13,16). On examining our data by region, however, we observed that females tended to have lower 25(OH)D values, especially in the Middle East/Africa and Asia/Pacific regions. Some have suggested that this finding may be related to cultural factors such as differences in clothing styles that may impede vitamin D conversion in the skin⁽²⁴⁾.

The highest mean 25(OH)D values were generally observed in North America, a finding that might be explained by the routine fortification of several foods (e.g. milk, juice and cereals) in the USA⁽²⁵⁾. The absence of significant differences between studies conducted in North America and those carried out in the Asia/Pacific region, however, may have been influenced by relatively high values found in Thailand, a country located near the equator with significant year-round sunlight exposure and higher daytime temperatures, resulting in the use of lighter-weight clothes, which afford less UV protection⁽²⁶⁾. Studies conducted in Japan and other Asian countries may have further contributed to somewhat higher regional values, resulting from diets rich in vitamin D foods such as oily fish(27).

Previous reviews (5,8,15) have reported an apparent northsouth gradient for 25(OH)D in Europe, with Scandinavian countries showing generally higher values than the Southern European countries. This finding is thought to result, in part, from population-based differences in skin pigmentation, diets rich in oily fish, the common use of cod-liver oil and a higher degree of vitamin D supplementation in Scandinavian countries (14,15). Although we did not find such a gradient in the present review, we observed generally higher 25(OH)D values in Swedish elderly than in those from other European countries. Some have suggested that this finding can be





References	Country	Distance from the equator (°)	Season	Age group	Number o			ES (95% CI)	Weight (%)
Female									
Rabbani et al. (2009) ⁽²¹¹⁾	Iran	36	Winter	Children/adolescents	539			60-34 (56-45, 64-23)	6-67
Hossein-Nezhad et al. (2009) ⁽²⁰⁴⁾	Iran	36		Adults	486			30-92 (29-31, 32-53)	6.75
Hosseinpanah et al. (2008) ⁽²⁰⁵⁾	Iran	35		Adults	245			73-00 (65-20, 80-80)	6-39
Charlton et al. (1996)(216)	South Afri	ica 34	Winter	Elderly	90	-		37-69 (35-06, 40-32)	6-72
Gannage-Yared et al. (2000) ⁽²¹⁴⁾	Lebanon	34	Winter	Adults	217			18-62 (16-70, 20-54)	6.74
Moussavi et al. (2005) ⁽²⁰⁹⁾	Iran	32	Winter	Children/adolescents	165	-		41-83 (38-65, 45-02)	6-70
Gharaibeh & Stoecker (2009)(22)	Jordan	31	Summer	Adults	93	-		25-60 (23-65, 27-55)	6.74
Dahifar et al. (2007) ⁽²⁰²⁾	Iran	30-5	Other	Children/adolescents	414	1		74-88 (71-08, 78-68)	6-67
Omrani <i>et al.</i> (2006) ⁽²⁸⁵⁾	Iran	29	Winter	Adults	676			28-90 (27-17, 30-63)	6.75
Subtotal ($I^2 = 990.3 \%$, $P = 0.000$)							>	43-24 (33-80, 52-67)	60-13
Male									
Rabbani et al. (2009) ⁽²¹¹⁾	Iran	36	Winter	Children/adolescents	424	1) 116·14 (111·00, 121·28)	6-60
Hossein-Nezhad et al. (2009) ⁽²⁰⁴⁾	Iran	36		Adults	160	-		32-57 (29-21, 35-93)	6-69
Gannage-Yared <i>et al.</i> (2000) ⁽²¹⁴⁾	Lebanon	34	Winter	Adults	96	-		35-74 (32-01, 39-48)	6-68
Charlton et al. (1996) ⁽²¹⁶⁾	South Afri	ica 34	Winter	Elderly	83	-		36-19 (33-67, 38-72)	6.72
Moussavi et al. (2005) ⁽²⁰⁹⁾	Iran	32	Winter	Children/adolescents	153	1		93.08 (85.66, 100.50)	6-43
Masoompour et al. (2008) ⁽²⁰⁷⁾	Iran	29	Winter	Adults	520	-		35-00 (33-54, 36-46)	6.75
Subtotal ($I^2 = 99.6 \%$, $P = 0.000$)						<		57-91 (39-19, 76-63)	39-87

Fig. 6. Forest plot for the Middle East/Africa region stratified by sex. ES, effect estimator. (A colour version of this figure can be found online at http://www. journals.cambridge.org/bjn)

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explained by the routine fortification of oil and low-fat milk products with vitamin D in Sweden⁽²⁸⁾.

Overall ($I^2 = 99.4 \%$, P = 0.000)

Note: weights are from random-effects analysis

In accordance with other reviews (5,8,15), our exploratory analyses also suggested that institutionalised elderly in Europe and the Asia/Pacific region had lower mean 25(OH)D values than the elderly living in the community. It is possible that such a finding may result from less time spent outdoors due to poorer health status (29), although similar findings in other groups of institutionalised individuals could be expected elsewhere. Further investigations of the patterns of vitamin D deficiency and insufficiency are needed in this vulnerable subgroup. Another interesting finding from our exploratory analyses was that newborns/infants were reported to have lower 25(OH)D values than the members of other age groups in several countries worldwide. Because newborn vitamin D status is mainly determined by maternal vitamin D status⁽³⁰⁾, this finding may be

75

49-05 (40-61, 57-48)

100

Table 2. Effect estimators (ES) from the meta-analyses stratified by age and region* (ES and 95 % confidence intervals)

Regions	I ² (%)	n (studies)	n (participants)	ES	95 % CI
Europe					_
Children/adolescents (>1-17 years)	99.5	6	1816	50.56	34.35, 66.77
Adults (>17-65 years)	99.4	35	28 844	52.98	45.01, 56.58
Elderly (>65 years)	99.4	30	10894	51.74	45.81, 57.66
North America					
Children/adolescents (>1-17 years)	98.5	3	993	78.35	59.44, 97.25
Adults (>17-65 years)	99.7	8	6201	71.83	57.71, 86.00
Elderly (>65 years)	99.3	15	5307	71.70	64.84, 78.57
Asia/Pacific					
Children/adolescents (>1-17 years)	85.4	3	899	31.89†	24.94, 38.84
Adults (>17-65 years)	99.5	13	3709	67.99	59.73, 76.25
Elderly (>65 years)	98.8	9	4965	66-16	62.16, 70.22
Middle East/Africa					
Children/adolescents (>1-17 years)	99-2	6	1913	75.41†	56.43, 94.38
Adults (>17-65 years)	98.5	6	2079	34.66	29.32, 40.01
Elderly (>65 years)	99-2	4	874	38-20	29.15, 47.25

^{*} Meta-analyses were not conducted for studies carried out in Latin America due to the limited number of eligible studies.



[†] Values were significantly different from those of the other age groups.



explained by generally inadequate vitamin D levels in pregnant women as suggested in previous work (31). Future research in these groups is needed to confirm these findings and test interventions aimed at interrupting this putative mechanism.

Strengths and limitations

To our knowledge, the present systematic review, conducted in accordance with the PRISMA statement (18), is among the first to focus on patterns of vitamin D status worldwide and in key population subgroups. We purposefully sought to identify studies with randomly selected samples from the general population to reduce sources of bias, which may otherwise obscure the public health importance of vitamin D status across the world. Use of continuous 25(OH)D values in our analyses is another important strength of the present study, given the inconsistent application of thresholds to indicate 25(OH)D deficiency, insufficiency and adequacy. A systematic search strategy based on two of the largest biomedical literature databases also reduced the probability of missing relevant articles. Besides the detailed data on 25(OH)D values among important subgroups by age, sex and region, the present review adds to the current understanding of vitamin D status in both developed and developing countries worldwide. We used the randomeffects model to account for the substantial heterogeneity that we observed across the studies. Between-study heterogeneity is common in systematic reviews, especially in observational epidemiology where unobserved characteristics at both the study and individual levels affect the outcomes of interest. The random-effects model adjusts for this heterogeneity by incorporating a between-study component of variance in the weights used for calculating the summary estimate (32).

It is important to consider the findings of the present review in the context of several potential limitations. First, we cannot fully exclude publication bias as studies reporting vitamin D deficiency might have been more likely to be published than those reporting mean or median levels within the normal range. Second, language bias may have affected the results, as we limited the present review to articles written in English. This may have accounted, for example, for the relative under-representation of studies conducted in Latin America in our sample. Efforts to identify and review studies published in languages other than English are needed in the future to gain a clear understanding of the full scope of vitamin D deficiency worldwide. Third, our strict inclusion criteria (e.g. inclusion of studies with randomly selected samples) might also explain the limited number of studies identified from some regions. However, previous reviews using more liberal inclusion criteria have also identified a limited number of studies conducted in these regions^(8,16). Fourth, recruitment strategies in the studies that we sampled may have focused to an extent on healthier populations, resulting in an overestimation of the prevalence of adequate vitamin D levels and a consequent minimisation of observable differences between the sexes or age-related subgroups. Fifth, we observed substantial heterogeneity between the studies in our sample that could not be explained by variables such as age, sex, season, distance from the equator, assay type or representativeness. Other unmeasured factors influencing vitamin D status (e.g. dietary intake, clothing style, time spent outdoors and use of sunscreen) may have contributed to the heterogeneity of results. Differences across the studies in study quality, adjustment for potential confounders and the definition of some characteristics or factors such as season may have contributed substantially to the heterogeneity that we observed. Finally, the precision of the estimates of vitamin D status in the subgroups of interest in the present review was probably affected by their relative under-representation in studies conducted in many regions of the world. High-quality population-based studies that assess and report all relevant data on 25(OH)D levels and central covariates including lifestyle factors to enable comparison of 25(OH)D values in the future, at least for population subgroups within the same country, have to be conducted.

Conclusion

Although we found a high degree of variability in reports of vitamin D status at the population level, more than one-third of the studies in the present systematic review reported mean 25(OH)D values below 50 nmol/l. Given the substantial heterogeneity of published evidence to date, further research on worldwide patterns of vitamin D deficiency at the population level and within key subgroups is needed to inform public health policy development to reduce risk for potential health consequences of an inadequate vitamin D status. The present review further suggests the importance of developing and implementing research designs that minimise potential sources of bias and consequently strengthen our understanding on vitamin D status in key subgroups worldwide.

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All authors declare that they have no conflicts of interest.





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