The InterLACE study: Design, data harmonization and characteristics across 20 studies on women’s health

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The InterLACE study: Design, Data Harmonization and Characteristics Across 20 Studies on
Women’s Health


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Abstract

Objectives: The International Collaboration for a Life Course Approach to Reproductive Health and Chronic Disease Events (InterLACE) project is a global research collaboration that aims to advance understanding of women’s reproductive health in relation to chronic disease risk by pooling individual participant data from several cohort and cross-sectional studies. The aim of this paper is to describe the characteristics of contributing studies and to present the distribution of demographic and reproductive factors and chronic disease outcomes in InterLACE.

Study design: InterLACE is an individual level pooled study of 20 observational studies (12 of which are longitudinal) from ten countries. Variables were harmonized across studies to create a new and systematic synthesis of life course data.

Main outcome measures: Harmonized data were derived in three domains: 1) socio-demographic and lifestyle factors, 2) female reproductive characteristics, and 3) chronic disease outcomes (cardiovascular disease (CVD) and diabetes).

Results: InterLACE pooled data from 229,054 mid-aged women. Overall, 76% of the women were Caucasian, 22% Japanese, and other ethnicity (of 300 or more participants) included Hispanic/Latin American (0.2%), Chinese (0.2%), Middle Eastern (0.3%), African/black (0.5%), and Other (1.0%). The median age at baseline was 47 years (Inter-quartile range (IQR): 41-53), and that at the last follow-up was 56 years (IQR: 48-64). Regarding reproductive characteristics, half of the women (49.8%) had their first menstruation (menarche) at 12-13 years of age. The distribution of menopausal status and the prevalence of chronic disease varied considerably among studies. At baseline, most women (57%) were pre- or peri-menopausal, 20% reported a natural menopause (range 0.8-55.6%) and remaining had surgery or were taking hormones. By the end of follow-up, the prevalence of CVD and diabetes were 7.2% (range 0.9-24.6%) and 5.1% (range 1.3-13.2%), respectively.

Conclusions: The scale and heterogeneity of InterLACE data provide an opportunity for strengthening evidence concerning the relationships between reproductive health through life and subsequent risks of chronic disease, including cross-cultural comparisons.

Keywords: baseline characteristics; reproductive health; chronic disease; life course research; cross-cultural comparison; harmonization
**Highlights**

- InterLACE is an international collaboration of 20 observational studies across 10 countries.
- Harmonized individual-level data on reproductive health and chronic disease are available from 230,000 women.
- The prevalence of diabetes and cardiovascular disease among mid-aged women were 5% and 7% at the end of study follow-up, respectively.
- InterLACE enables a detailed review of methodologies currently used in the field of women’s health.
1. Introduction

Since chronic diseases are typically characterized by long latency and complex causal pathways, the clear sex differences evident in their risks [1] highlight the need to understand the role of reproductive characteristics and sex hormones in non-communicable diseases (NCDs) across life. For instance, women with diabetes have a 3.5-fold increased risk of mortality from coronary heart disease, compared with 2-fold for men with diabetes [1]. Some aspects of female reproductive health act as markers for increased risk of NCDs in later life, in that they may signal an underlying predisposition or sub-clinical conditions [2-4]. Early menarche is associated with increased risk of type 2 diabetes mellitus (T2DM), cardiovascular disease (CVD) [5,6], and breast cancer [7]. Early menarche is also linked to poor reproductive health outcomes across life, such as irregular menstrual cycles [8], but with better bone health in later life [9,10]. Similarly, early menopause increases the risk of having chronic diseases in later life including T2DM and CVD [11,12], while the vasomotor symptoms and longer duration of menopausal transition also represent a period of increased metabolic and cardiovascular risks [13,14]. Various lifestyle, socioeconomic, and cultural factors also influence reproductive characteristics and chronic disease risk [15-17]. A more detailed understanding of the complex relationships between these modifiable factors and reproductive characteristics is needed to support targeted gender-specific preventive strategies for chronic diseases. Previous research based on individual studies has been constrained by issues such as small sample size, lack of control for comorbidities, and lack of sufficient information on the racial/ethnic and cultural diversity of the study samples.

The International Collaboration for a Life Course Approach to Reproductive Health and Chronic Disease, or InterLACE, aims to advance the evidence base for women’s health policy by developing a collaborative research program that takes a comprehensive life course perspective of women’s reproductive health in relation to chronic disease risk [18]. Established in June 2012, InterLACE has pooled individual-level observational data on reproductive health and chronic disease from almost 230,000 women from 20 observational studies, mostly on women’s health, across ten countries. InterLACE offers an integrated approach for a more detailed understanding of the determinants and characteristics of reproductive health across the life course in diverse populations [18]. A life course perspective emphasizes the differential effects of exposures and events at different stages of life [19], which in turn can be reflected in models that capture the different types of biological, psychological, and social mechanisms at work [20].

Findings from InterLACE can therefore provide insights into causal pathways for disease aetiology [21] and have implications for the timing and targeting of preventive health interventions [22]. This will enable a more detailed description of reproductive function and ageing by quantifying the markers of reproductive health through life, such as age at menarche, parity, and age at menopause in different populations. The project will determine the extent to which these markers and overall trajectories of lifetime reproductive health are associated with future chronic disease risks such as T2DM and CVD. Through InterLACE, the relationships of lifestyle, cultural factors, and reproductive health with subsequent risk of chronic disease will be identified. Recommendations for future study designs to facilitate rigorous cross-cultural comparisons across longitudinal studies will also be presented. The aim of this paper is to present the overall demographic and reproductive characteristics and to describe the prevalence of T2DM and CVD in InterLACE.
2. Methods

2.1 Study recruitment

Twenty observational studies, twelve of which are longitudinal, currently provide data for
InterLACE: Australian Longitudinal Study on Women’s Health (ALSWH) [23], Healthy Ageing of
Women Australia (HOW) [24], Melbourne Collaborative Cohort Study (MCCS) [25], Danish Nurse
Cohort Study (DNC) [26], Women's Lifestyle and Health Study (WLH) [27], Medical Research
Council (MRC) National Survey of Health and Development (NSHD) [28], National Child
Development Study (NCDS) [29], English Longitudinal Study of Ageing (ELSA) [30], UK Women's
Cohort Study (UKWCS) [31], Whitehall II study (WHITEHALL) [32], The Study of Women's Health
Across the Nation (SWAN) [33], Seattle Midlife Women’s Health Study (SMWHS) [34], Japan
Nurses’ Health Study (JNHS) [35], Japanese Midlife Women’s Health Study (JMWHS) [24], Hilo
Women’s Health Study (HILO) [36], San Francisco Midlife Women’s Health Study (SFMWHS) [37],
and The Decision at Menopause Study (DAMES-USA [38], Lebanon [39], Spain [40], Morocco [41]).
Participants in each study were recruited under Institutional Review Board protocols approved at each
research centre and provided informed consent. Details of the study design, recruitment, and research
aims for each study have been published elsewhere (see above for references). Brief descriptions of
the 20 studies are given in Table 1, with their geographic scope shown in Figure 1.

The majority of studies began between 1990 and early 2000, with the exception of NSHD (1946
British Birth Cohort) and NCDS (1958 British Birth Cohort), in which participants (male and female)
were recruited at birth. InterLACE used data from a sub-sample study of women’s health (n=1570)
from NSHD started in 1993 (and the baseline for InterLACE), when participants were aged 47 years,
with annual follow-up surveys until 2000 (age 54 years) to capture timing of menopause, menopausal
symptoms and menopausal hormone therapy (MHT) use [28]. Similarly, for NCDS we used data
from the women’s health survey in 2008 (n=5274) as the baseline when cohort members were aged
50 years and were followed up until 2013 for disease outcome.

The DNC and ELSA studies had multiple waves of recruitment. DNC first invited members of the
Danish Nurses Organisation to participate in 1993, with both a follow-up and recruitment of
additional nurses in 1999 [26]. ELSA commenced in 2002-03 (wave 1) with the original sample
recruited from households that had earlier participated in the Health Survey for England (HSE) in
1998, 1999, and 2001 (wave 0) [30]. New cohorts that were recruited from households that had
participated in HSE in 2001-04 and 2006 were added to the ELSA sample at wave 3 (2006-07) and
wave 4 (2008-09), respectively. The baseline years used in InterLACE for DNC and ELSA were
determined according to the year in which each participant was recruited.

The SWAN and SMWHS had different recruitment criteria at baseline. In SWAN, only women with
at least one menstrual period in the previous three months, without surgical removal of the uterus
and/or both ovaries, and without the current use of hormone therapy, were eligible. In SMWHS, only
women without surgical removal of uterus or ovaries were eligible to participate.

2.2 Study variables
InterLACE invited all individual studies to provide relevant data including a list of variables, survey questionnaires, data dictionaries/formats, and protocols or standard operating procedures. The data were requested from the three key domains:

1. **Socio-demographic and lifestyle factors**: age, birth year, race/ethnicity, marital and employment status, the level of education, body mass index (BMI), smoking status, alcohol consumption, physical activity, food and vegetable intakes, the consumption of soy products were provided if available. Marital status, employment, and lifestyle variables were also available at multiple time points in some longitudinal studies and were all preserved, although only baseline data are presented here. Use of these exposure variables will vary depending on the research questions.

2. **Female reproductive characteristics**: studies provided some or all of the following self-reported markers of reproductive health through life: age at menarche, age at first birth, number of pregnancies, parity, timing and duration of oral contraceptive pill (OCP) use, MHT use, age at natural menopause, hysterectomy/oophorectomy, menopausal status, and menopausal symptoms (e.g. vasomotor symptoms and psychological symptoms) [20]. Time-varying reproductive variables such as hormone use, surgery history, menopausal status, and menopausal symptoms were also available at multiple surveys in the longitudinal studies.

3. **Chronic disease outcomes**: data on CVD (stroke and heart diseases including general heart disease, heart attack, heart failure and angina) and diabetes (Type 1 and Type 2 diabetes) were collected from self-reported survey questionnaires and linkage with national registries (for DNC, WLH and SMWHS). Four studies (JMWHS, DAMES-USA, Lebanon, and Spain) did not have data available on CVD or diabetes.

### 2.3 Data harmonization

Once individual-level datasets were received, data were checked for outliers and inconsistencies, and if present, data providers were queried and the issue resolved. Harmonization rules and recoding instructions were created for each variable. When multiple studies had more detailed but similar information available, extra variables were created to encompass this alternative format and benefit from the increased granularity. In general, categorical variables were collapsed into the simplest level of detail to incorporate information from as many studies as possible. For example, education categories varied from study to study. It was categorised into ≤10 years, 11-12 years, and >12 years. Harmonized education category of less or equal to 10 years corresponds to less than high school or Certificate of Secondary Education (CSE) or General Certificate of Education Ordinary Level (GCE O-level) in the UK. Similarly, 11-12 years category corresponds to high school or GCE Advanced Level (A-level) in the UK, and >12 years corresponds to at least some college education including trade, certificate, vocational training, diploma, and university degree.

Harmonization of other specific variables such as race/ethnicity and menopausal status are presented in **Figures 2** and **Figure 3**. In detail, participants self-identified their specific race/ethnicity and/or population subgroup in ten studies from which ethnicity variable was defined. Of the remaining ten studies, ethnic groups were defined based on country of birth and language spoken at home (5 studies),
and where these were not available (DNC, JNHS, JMWHS, DAMES-Lebanon, and DAMES-Morocco), the country where the study was conducted was considered as a residency variable and used as a proxy for ethnicity [42]. In total, ten ethnic groups were defined: Caucasian-Australian/New Zealander, Caucasian-European, Caucasian-North American, Hispanic/Latin American, Asian-Japanese, Asian-Chinese, Asian-Other (South/Southeast Asian), Middle Eastern, African/Black, and Other (Native American, Pacific Islander, Caribbean, Hawaiian, and Mixed). We then collapsed Australian/New Zealander, European, and North American together as Caucasian, and combined Asian-Other and Other.

To harmonize menopausal status at baseline, we first reviewed 14 studies that either had predefined menopausal status (pre-, peri-, or post-menopause) or reasons for the cessation of menses. Among them, those reporting current use of hormone therapy (unless natural menopause specifically reported) and hysterectomy/oophorectomy were categorised separately. As a result, we have six categories of menopausal status: hysterectomy/oophorectomy, current MHT use, current OCP use, pre-menopause, peri-menopause, and natural menopause. For all other women, where predefined menopausal status was not available, we used related variables (hysterectomy/oophorectomy, current use of hormone, menstrual period in the last 12 months, menstrual period in the last 3 months, and irregular or changeable period) using a consistent rule (Figure 3) to assign them to one of the six groups defined above. In this way, each woman was provided with consistent and harmonized data on menopausal status at baseline. The same rules applied for the follow-up surveys. However once women had gone through natural menopause or surgery (hysterectomy/oophorectomy), their menopausal status remained throughout for any subsequent surveys. In addition to the harmonized menopausal status, more detailed information about the current and past use of MHT and OCP, hysterectomy, and unilateral/bilateral oophorectomy are available as separate variables. In this paper, we only present socio-demographic and reproductive characteristics at baseline, and show the cumulative prevalence of chronic disease outcomes over the study period. We used SAS 9.4 (SAS Institute, Inc., Cary, NC) for all data management and analysis.

3. Results

The InterLACE dataset pooled individual-level data from 229,054 participants. Of the twenty studies currently comprising InterLACE, nine are national cohorts from Australia, the USA, the UK, Japan, Sweden, Norway, and Denmark. The remaining state-based studies from specific cities or regions including San Francisco, Seattle, Hawaii, and Massachusetts in the USA; London, England; Melbourne and Queensland in Australia; Nagano, Japan; Beirut, Lebanon; Madrid, Spain; and Rabat, Morocco (Figure 1). Twelve studies provided longitudinal data with at least two waves of surveys and five years of follow-up, while eight studies provided only cross-sectional baseline data (Table 1). For the majority of studies, women’s average age at baseline was between 40 and early 50 years with an overall median of 47 years (IQR: 41-53 years), with the exceptions of HOW, MCCS, and ELSA where the women were older at baseline (median ranging from 55-58 years). JMWHS only provided categorical age (≤55 or >55 years), and almost half (48%) of the women were more than 55 years of age.
Table 2 presents the distribution of some key harmonized demographic and reproductive variables by studies at baseline. Of the seven categories of ethnicity, Caucasian (75.5%), Australian/New Zealander 12.6%, European 61.7%, North American 1.2%) were the most prevalent, followed by Japanese (22.4%, mainly living in Japan (98.9%) but also some living in the USA). The remaining minority racial/ethnic groups included Hispanic/Latin American, Chinese, Middle Eastern, African/Blacks, and Others, with a minimum of 300 participants in each group. Within studies, four (SWAN, SMWHS, HILO, and SFMWHS) had a combination of multi-racial/ethnic samples. The level of education varied greatly between studies. Some variations were due to original study designs (e.g. study of nurses). However, this could also be reflecting regional variation in education. For example, DAMES-Morocco had a very small percentage of women (4%) with >12 years of education, while most US studies had over 75% at that level. Meanwhile, >12 years of education was significantly lower in NSHD compared with other UK studies. In most studies, the percentage of unmarried women was less than 10%, except for WHITEHALL and JNHS, which both had more than 20% single women. In WLH, more than double the average percentage of women (38.4%) were single because marital status was recorded from mother’s birth registry, so for those who had not given birth this information was missing. The overall prevalence of obesity (BMI ≥30 kg/m²) was 10%. In four studies (ELSA, SWAN, SFMWHS, and DAMES-USA) nearly 30% of women were obese, while the corresponding figure for Japanese studies (JMWH and JNHS) was less than 2%.

Regarding reproductive factors, 40-60% of women reported that they had their first period (menarche) between the ages of 12 to 13 years. The percentage of women with earlier menarche (≤11 years) was around 20%, except for DNC and DAMES-Morocco where this was less than 10%. At baseline most women (57%) were still pre- or peri-menopausal, 20% reported natural menopause (range 0.8-55.6% among studies), 13% had hysterectomy or oophorectomy (range 1.7-29.6%), and the remaining 10% were taking either MHT or OCP. The distribution of vasomotor symptoms also varied considerably among studies, reflecting the range of age and menopausal status among studies. The studies with the oldest baseline age of late 50s (HOW, MCCS, ELSA, and JMWH) had the highest proportions of naturally menopausal women (range 43.5-55.6%) and high prevalence of vasomotor symptoms (30-50%). Conversely, studies with a younger baseline age of early 40s (WLH, SMWHS, and SFMWHS) had lower proportions of natural menopause (<3%) and lower prevalence of vasomotor symptoms (10-20%).

The prevalence of CVD and diabetes at baseline for cross-sectional studies and at the end of the follow-up period for the 12 longitudinal studies are provided in Table 3. Overall, the median age at last follow-up for disease outcome was 56 years (IQR: 48-64 years). The prevalence of CVD and diabetes were higher in longitudinal studies that followed participants into their 60s or 70s of age. The overall prevalence of CVD was 7.2%, but it ranged from 0.9-24.6% between studies with the lowest in JNHS (median age 41 years) and the highest in ELSA (median age 65 years). Of the total CVD cases, 2.0% were stroke and 5.8% were heart disease. There was little variation in the prevalence of stroke between studies, except for ELSA, which had more than double the prevalence (5.6%) of other studies. A wider variation was evident in the prevalence of heart disease across studies, which ranged from 0.6-22.4%. The overall prevalence of diabetes was 5.1%, with JNHS having the lowest (1.3%) prevalence and SWAN the highest (13.2%).

4. Discussion
With the pooled information from 230,000 mid-aged women across 20 cohort and cross-sectional studies, from ten countries, InterLACE has sufficient scale and heterogeneity to study the health of women in midlife. It provides a unique opportunity for advancing understanding of the relationships between reproductive characteristics and chronic diseases that are shown to have marked sex differences in their aetiology and prevalence. The study has assembled a broad spectrum of prospective data on mid-aged women, including socioeconomic status (education and marital status), lifestyle (BMI, smoking, and physical activities), reproductive factors (menarche, parity, and menopause), and disease outcomes (diabetes and CVD). It comprises a diverse range of race/ethnic groups (Caucasian, Asian, and Blacks) that enables inferences to be drawn regarding minority subgroups that would otherwise be underpowered in individual studies. This heterogeneity is important for detecting relationships that may not be apparent in homogeneous populations and increases the generalizability of the study findings.

The overall distribution of measures in InterLACE data are broadly consistent with that in the published literature, for example, most of the women had their first menstrual period between 12 and 13 years of age [43,44]. Similarly, the overall prevalence of obesity (10% at baseline) and diabetes (5% by final survey) among mid-aged women was comparable with the global prevalence of these conditions in the early 2000s [45,46].

The process of combining individual-level data from multiple cohorts and cross-sectional studies for InterLACE inevitably leads to a number of methodological challenges. The contributing studies varied in their sampling methods, inclusion and exclusion criteria, and modes of survey administration. For instance, women may respond differently to questions about their reproductive health if the survey is completed on-line or via a telephone interview compared with a self-completed paper-based questionnaire, which was the most frequently used data collection method. Retention of participants is an issue for all longitudinal studies. The contributing studies have different levels of sample attrition and missing data due to withdrawal, mortality, and other reasons for non-response at each wave of data collection. The studies also varied greatly in terms of likely representativeness of the sample with respect to the relevant national population; for example sampling from specific professional groups as illustrated by women in the civil service for the Whitehall II study, or women nurses for the DNC and JNHS studies. Variations in the prevalence of CVD across studies already serve to illustrate the effect of differences in the age range of the cohorts of women when they responded to the relevant survey questions. Future analyses of the data from InterLACE will need to identify and adjust for these potential sources of heterogeneity and clustering of information.

5. Conclusion

Despite the challenges, this study profile shows that InterLACE has the potential to build a more detailed understanding of the differential effects of timing, frequency or duration of reproductive characteristics on the risk of key chronic disorders. This will allow for the development of distinct profiles of reproductive characteristics throughout life. Because these profiles are likely to be associated with risk of chronic disease in later life, they have the potential to be developed as the basis for a more tailored approach for preventive health strategies when women discuss reproductive issues with health professionals. Moreover, such health service encounters may present an opportunity for timely and targeted interventions to reduce chronic disease risk [47] that can be
enhanced to individual needs through understanding the interactions between reproductive health profiles and modifiable risk factors for cardiovascular and metabolic conditions. Crucially, InterLACE also enables a detailed review of methodologies currently used in the field of menopausal symptom research. This will result in recommendations for study design, symptom measures, and reporting of results to improve international and cross-cultural comparisons. Standardization of methods will become increasingly important to enhance the value of studies of women’s health in low and middle-income countries and where currently there are manifest gaps in knowledge.

Further information is available on the InterLACE website http://interlace.org.au. The pooled data set is governed by a Collaborative Research Agreement among several institutions. Those interested in collaborating on the project can contact the scientific committee at interlace@uq.edu.au.

Acknowledgements

The data on which this research is based were drawn from several global observational studies including: Australian Longitudinal Study on Women’s Health (ALSWH), Healthy Ageing of Women Australia (HOW), Melbourne Collaborative Cohort Study (MCCS), Danish Nurse Cohort Study (DNC), Women's Lifestyle and Health Study (WLH), Medical Research Council National Survey of Health and Development (1946) (NSHD), National Child Development Study (1958) (NCDS), English Longitudinal Study of Ageing (ELSA), UK Women's Cohort Study (UKWCS), Whitehall II study (WHITEHALL), The Study of Women's Health Across the Nation (SWAN), Seattle Midlife Women’s Health Study (SMWHS), Japan Nurses’ Health Study (JNHS), Japanese Midlife Women’s Health Study (JMWHS), Hilo Women's Health Study (HILO), San Francisco Midlife Women’s Health Study (SFMWHS), and The Decision at Menopause Study (DAMES).

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Central Laboratory: University of Michigan, Ann Arbor – Daniel McConnell (Central Ligand Assay Satellite Services).


Steering Committee: Susan Johnson, Current Chair
Chris Gallagher, Former Chair

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Contributors

GDM conceived the study design and contributed to interpretation of the data and drafted the manuscript. LJ, HFC, NP harmonized the data and performed statistical analysis. AJD, DA contributed to interpretation of the data. NEA, SLC, EBG, DB, LLS, EB, JEC, VJB, DCG, GGG, FB, AG, KH, JSL, HM, DK, RC, RH, CMO, KAL, MKS, TY, NFW, ESM, MH, PD, SS, HOA, EW provided study data. All authors contributed to critical revision of the manuscript.
Conflicts of interest

The authors have no conflicts of interest to declare.

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Ethical approval

Each study in the InterLACE consortium has been undertaken with ethical approval from the relevant authorities and with the informed consent of participants.
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Table 1 Twenty studies contributing to the InterLACE dataset (n=229,054)

<table>
<thead>
<tr>
<th>Study (abbreviation)</th>
<th>Location</th>
<th>Baseline survey year</th>
<th>Baseline sample</th>
<th>Baseline age median (IQR)</th>
<th>No. of survey included</th>
<th>Latest survey year</th>
<th>Latest survey sample</th>
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<tbody>
<tr>
<td>Longitudinal data provided (n=175,749)</td>
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<td></td>
<td></td>
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</tr>
<tr>
<td>Australian Longitudinal Study on Women’s Health (ALSWH)</td>
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<td>1996</td>
<td>13,715</td>
<td>48 (46-49)</td>
<td>7</td>
<td>2013</td>
<td>9,151</td>
</tr>
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<td>Australia</td>
<td>2001</td>
<td>868</td>
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<td>3</td>
<td>2011</td>
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<tr>
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<td>1990-94</td>
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<td>55 (48-62)</td>
<td>3</td>
<td>2003-2006</td>
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<td>Denmark</td>
<td>1993/1999</td>
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<td>50 (47-58)</td>
<td>2</td>
<td>1999</td>
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<td>Sweden/Norway</td>
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<td>40 (35-45)</td>
<td>2</td>
<td>2003-2004</td>
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<td>UK</td>
<td>1993†</td>
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<td>47†</td>
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<td>UK</td>
<td>2008‡</td>
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<td>50‡</td>
<td>2</td>
<td>2013</td>
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<tr>
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<td>UK</td>
<td>2002-09</td>
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<td>58 (52-68)</td>
<td>5</td>
<td>2010-2011</td>
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<td>1996</td>
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<td>11</td>
<td>2006</td>
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<td>508</td>
<td>41 (38-44)</td>
<td>2</td>
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<td>2001-2007</td>
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<td>41 (35-47)</td>
<td></td>
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<tr>
<td>Japanese Midlife Women’s Health Study (JMWHS)</td>
<td>Japan</td>
<td>2002</td>
<td>847</td>
<td>N/A (45-60)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hilo Women's Health Study (HILO)</td>
<td>USA</td>
<td>2004-05</td>
<td>994</td>
<td>51 (46-56)</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>San Francisco Midlife Women’s Health Study (SFMWHS)</td>
<td>USA</td>
<td>1996</td>
<td>347</td>
<td>43 (42-45)</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>The Decision at Menopause Study (DAMES-USA)</td>
<td>USA</td>
<td>2001</td>
<td>293</td>
<td>50 (48-53)</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>The Decision at Menopause Study (DAMES-Lebanon)</td>
<td>Lebanon</td>
<td>1997</td>
<td>298</td>
<td>50 (48-53)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>The Decision at Menopause Study (DAMES-Spain)</td>
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<td>2002</td>
<td>300</td>
<td>50 (47-53)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>The Decision at Menopause Study (DAMES-Morocco)</td>
<td>Morocco</td>
<td>1998</td>
<td>299</td>
<td>49 (46-52)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Abbreviation: N/A, not applicable; IQR, interquartile range.

a NSHD (1946 British Birth Cohort) and NCDS (1958 British Birth Cohort) first collected information on women health in 1993 (aged 47) and in 2008 (aged 50), respectively, so we used 1993 and 2008 as the baseline year for the InterLACE.

b JMWHS provided age by category only, and 48% of women were aged more than 55 (age range: 45-60 years).

c The latest survey data contributed to the InterLACE dataset.
<table>
<thead>
<tr>
<th>Study</th>
<th>Race/Ethnicity (%)</th>
<th>Educationa (%)</th>
<th>Marital status (%)</th>
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<tbody>
<tr>
<td></td>
<td>n</td>
<td>Caucasian Latino</td>
<td>Asian Japanese Chinese</td>
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<tr>
<td>Overall</td>
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<td>75.5</td>
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<tr>
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<td>13,715</td>
<td>96.1</td>
<td>0.3</td>
</tr>
<tr>
<td>HOW</td>
<td>868</td>
<td>96.5</td>
<td>N/A</td>
</tr>
<tr>
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<td>100</td>
<td>N/A</td>
</tr>
<tr>
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<td>28,731</td>
<td>100</td>
<td>N/A</td>
</tr>
<tr>
<td>WLH</td>
<td>49,259</td>
<td>100</td>
<td>N/A</td>
</tr>
<tr>
<td>NSHD</td>
<td>1,570</td>
<td>100</td>
<td>N/A</td>
</tr>
<tr>
<td>NCDS</td>
<td>5,274</td>
<td>98.0</td>
<td>N/A</td>
</tr>
<tr>
<td>ELSA</td>
<td>9,118</td>
<td>96.4</td>
<td>N/A</td>
</tr>
<tr>
<td>UKWCS</td>
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<td>98.7</td>
<td>N/A</td>
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<td>WHITEHALL</td>
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<td>84.2</td>
<td>N/A</td>
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<td>46.9</td>
<td>8.7</td>
</tr>
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<td>77.2</td>
<td>1.2</td>
</tr>
<tr>
<td>Cross-sectional data</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>JNHS</td>
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<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>JWMSH</td>
<td>847</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>HILO</td>
<td>994</td>
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<td>0.9</td>
</tr>
<tr>
<td>SFMWSH</td>
<td>347</td>
<td>46.4</td>
<td>27.4</td>
</tr>
<tr>
<td>DAMES-USA</td>
<td>293</td>
<td>94.2</td>
<td>1.0</td>
</tr>
<tr>
<td>DAMES-Lebanon</td>
<td>298</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>DAMES-Spain</td>
<td>300</td>
<td>95.3</td>
<td>3.7</td>
</tr>
<tr>
<td>DAMES-Morocco</td>
<td>299</td>
<td>N/A</td>
<td>N/A</td>
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</table>

Table 2: Baseline demographic and reproductive variables for the 20 studies
<table>
<thead>
<tr>
<th>Study</th>
<th>Normal &lt;25 kg/m²</th>
<th>Overweight 25-29.9 kg/m²</th>
<th>Obese ≥30 kg/m²</th>
<th>Age at menarche (n)</th>
<th>Menopausal status (%)</th>
<th>Vasomotor symptoms a (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>kg/m²</td>
<td>kg/m²</td>
<td>≤11 years</td>
<td>12-13 years</td>
<td>≥14 years</td>
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<tr>
<td>Overall</td>
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<td>66.9</td>
<td>23.2</td>
<td>10.0</td>
<td>214,759</td>
<td>16.9</td>
</tr>
<tr>
<td>Longitudinal data</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
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<td>28.9</td>
<td>18.6</td>
<td>11,396</td>
<td>18.8</td>
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<td>821</td>
<td>43.2</td>
<td>32.0</td>
<td>24.7</td>
<td>508d</td>
<td>19.5</td>
</tr>
<tr>
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<td>36.2</td>
<td>21.9</td>
<td>24,389</td>
<td>16.5</td>
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<tr>
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<td>22.8</td>
<td>5.6</td>
<td>28,477</td>
<td>7.9</td>
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<tr>
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<td>21.8</td>
<td>5.8</td>
<td>48,544</td>
<td>12.9</td>
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<tr>
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<td>60.7</td>
<td>25.5</td>
<td>13.8</td>
<td>1,242</td>
<td>16.2</td>
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<tr>
<td>NCDS</td>
<td>4,158</td>
<td>44.4</td>
<td>33.0</td>
<td>22.6</td>
<td>4,227</td>
<td>16.5</td>
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<tr>
<td>ELSA</td>
<td>7,485</td>
<td>34.4</td>
<td>37.6</td>
<td>28.0</td>
<td>6,314d</td>
<td>20.9</td>
</tr>
<tr>
<td>UKWCS</td>
<td>33,990</td>
<td>64.8</td>
<td>25.4</td>
<td>9.8</td>
<td>34,596</td>
<td>22.1</td>
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<tr>
<td>WHITEHALL</td>
<td>3,411</td>
<td>61.1</td>
<td>27.9</td>
<td>11.0</td>
<td>N/A</td>
<td>N/A</td>
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<tr>
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<td>26.9</td>
<td>33.0</td>
<td>3,267</td>
<td>24.2</td>
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<tr>
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<td>55.4</td>
<td>25.8</td>
<td>18.7</td>
<td>507</td>
<td>22.9</td>
</tr>
<tr>
<td>Cross-sectional data</td>
<td></td>
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<tr>
<td>JNHW</td>
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<td>11.0</td>
<td>1.8</td>
<td>49,175</td>
<td>21.0</td>
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<tr>
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<td>13.1</td>
<td>1.2</td>
<td>N/A</td>
<td>N/A</td>
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<tr>
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<td>955</td>
<td>46.9</td>
<td>29.7</td>
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<td>972</td>
<td>25.4</td>
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<td>32.3</td>
<td>31.3</td>
<td>N/A</td>
<td>N/A</td>
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<tr>
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<td>43.7</td>
<td>29.0</td>
<td>27.3</td>
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<td>22.3</td>
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<td>DAMES-Lebanon</td>
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<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>298</td>
<td>21.1</td>
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<td>33.0</td>
<td>8.0</td>
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<td>20.9</td>
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<td>N/A</td>
<td>N/A</td>
<td>259</td>
<td>10.0</td>
</tr>
</tbody>
</table>

**Abbreviation:** N/A, not applicable; MHT, menopause hormone therapy; OCP, oral contraceptive pill.

**a** Education ≤10 years corresponds to less than high school (equivalent to CSE or GCE O level in the UK), 11-12 years to high school (equivalent to GCE A level in the UK), and >12 years to at least some college (including trade, certificate, vocational training, diploma, and university degree).

**b** In the WLH study, marital status was only recorded from mothers' birth registry hence the data were missing for all women who did not give birth.
Body mass index data were reported as body weight appearance by category only (e.g. normal, overweight, obese), instead of measured or self-reported weight and height.

In the HOW study, age at menarche was only collected from survey 2 in 2006; in the ELSA study, age at menarche was only collected at wave 3 and wave 4 hence the data were missing for those women who lost to follow-up.

Had surgery category included hysterectomy or oophorectomy.

The baseline eligibility criteria for the SWAN study were: at least one menstrual period in the previous three months, without surgical removal of the uterus and/or both ovaries, and without the current use of hormone therapy. The baseline eligibility for the SMWHS study was without surgical removal of uterus or ovaries.

In the DAMES studies, women on MHT use were categorised as post-menopause.

Vasomotor symptoms were asked whether participants had experienced the symptoms in different time periods prior to baseline: in the last 12 months (ALSWH, NSHD, and NCDS), in the past month (DAMES studies), in the last one/two weeks (SFMWHS, SWAN, and HILO), and in the past 24 hours/at the moment (HOW, WHITEHALL, SMWHS, and JMWHS).
Table 3 The prevalence of chronic diseases at the end of study follow-up for the 20 studies

<table>
<thead>
<tr>
<th>Study</th>
<th>Age at last follow-up median (IQR)</th>
<th>Cardiovascular disease</th>
<th></th>
<th>Diabetes</th>
<th>Type 1 or Type 2</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Stroke and/or heart disease (%)</td>
<td>n</td>
<td>Stroke (%)</td>
<td>Heart diseases (%)</td>
</tr>
<tr>
<td>Overall</td>
<td>56 (48-64)</td>
<td>218,082</td>
<td>7.2</td>
<td>217,608</td>
<td>2.0</td>
</tr>
<tr>
<td>Longitudinal data&lt;sup&gt;a&lt;/sup&gt;</td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ALSWH</td>
<td>63 (60-65)</td>
<td>13,714</td>
<td>12.3</td>
<td>13,714</td>
<td>2.9</td>
</tr>
<tr>
<td>HOW</td>
<td>63 (60-66)</td>
<td>522</td>
<td>13.2</td>
<td>515</td>
<td>2.3</td>
</tr>
<tr>
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<td>24,467</td>
<td>10.3</td>
<td>24,467</td>
<td>2.9</td>
</tr>
<tr>
<td>DNC</td>
<td>64 (50-73)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>28,640</td>
<td>10.9</td>
<td>28,592</td>
<td>2.9</td>
</tr>
<tr>
<td>WLH</td>
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<td>49,149</td>
<td>6.0</td>
<td>49,021</td>
<td>2.2</td>
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<tr>
<td>NSHD</td>
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<td>1,526</td>
<td>13.6</td>
<td>1,518</td>
<td>0.8</td>
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<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
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<td>24.6</td>
<td>9,115</td>
<td>5.6</td>
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</tr>
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<td>7.8</td>
<td>3,300</td>
<td>3.1</td>
</tr>
<tr>
<td>SMWHS</td>
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<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
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<tr>
<td>Cross-sectional data&lt;sup&gt;a&lt;/sup&gt;</td>
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<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>JNHS</td>
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<td>49,658</td>
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<td>N/A</td>
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<td>N/A</td>
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<tr>
<td>SFMWHS</td>
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<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>DAMES-USA</td>
<td>50 (48-53)</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>DAMES-Lebanon</td>
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<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>DAMES-Spain</td>
<td>50 (47-53)</td>
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<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>DAMES-Morocco</td>
<td>49 (46-52)</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>

N/A, not applicable; IQR, interquartile range.

<sup>a</sup> Longitudinal studies provided the cumulative prevalence of chronic diseases over the study follow-up period. Once women reported they had CVD or diabetes, their disease status carried forward at subsequent surveys. Cross-sectional studies only provided the prevalence of disease at baseline.
b DNC, WLH, and SMWHS provided diseases outcome data from survey questionnaires and also from hospital registries (DNC: 1993-2013, WLH: 1991-2010, SMWHS: 1990-2013). NSHD also provided disease outcome data from the latest 2010 survey, when cohort members were aged 64 years.

c Heart diseases included general heart disease, heart attack, heart failure and angina.
Figure legends

**Figure 1** Locations of the 20 studies contributing to the InterLACE study

There are ten participating countries: Australia, Denmark, Sweden, Norway, UK, USA, Japan, Lebanon, Spain, and Morocco.

**Figure 2** Example of data harmonization to obtain common categories for race/ethnicity

**Figure 3** Example of data harmonization to obtain common categories for menopausal status.

Abbreviations: MHT, menopause hormone therapy; OCP, oral contraceptive pill.
**Figure 1** Locations of the 20 studies contributing to the InterLACE study

There are ten participating countries: Australia, Denmark, Sweden, Norway, UK, USA, Japan, Lebanon, Spain, and Morocco.
**Race/ethnicity**

10 studies had self-reported race/ethnicity
(MCCS, NCDS, ELSA, UKWCS, WHITEHALL, SWAN, SMWH, HILO, SFMWHS, DAMES-USA)

10 studies did not have race/ethnicity variable

5 studies had data on related variables

1. Country of birth* (ALSWH, HOW, NSHD, DAMES-Spain) or country of residency in childhood (WLH)
2. Language spoken at home (ALSWH, HOW)

*Note:
   a. Women who born in South Africa and had migrated to Australia or Spain were categorised as Caucasian.
   b. Women born in South America and Central America countries were categorised as Hispanic/Latin American.
   c. Morocco and Egypt were categorised as Middle Eastern countries.
   d. Women born in China, Hong Kong, and Taiwan were categorised as Chinese, while women born in Singapore/Malaysia were categorised as Asian-other.

5 studies only had data on country of study

Country of residency (DNC, JNHS, JMWH, DAMES-Lebanon, DAMES-Morocco)

5 studies had insufficient information

5 studies only had data on country of study

Country of residency (DNC, JNHS, JMWH, DAMES-Lebanon, DAMES-Morocco)

Race/Ethnicity 10 categories

1. Caucasian-Australian/New Zealander
2. Caucasian-European
3. Caucasian-North American
4. Hispanic/Latin American
5. Asian-Japanese
6. Asian-Chinese
7. Asian-Other (South/Southeast Asian)
8. Middle Eastern
9. African/Black
10. Other (Native American, Pacific Islander, Caribbean, Hawaiian, mixed)

Race/Ethnicity 7 categories

1. Caucasian (Australian/New Zealander, European, North American)
2. Hispanic/Latin American
3. Asian-Japanese
4. Asian-Chinese
5. Middle Eastern
6. African/Black
7. Other (Asian-Other, Native American, Pacific Islander, Caribbean, Hawaiian, mixed)

**Figure 2** Example of data harmonization to obtain common categories for race/ethnicity
Figure 3 Example of data harmonization to obtain common categories for menopausal status

Abbreviations: MHT, menopause hormone therapy; OCP, oral contraceptive pill