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BMJ Open Variation in the estimated prevalence of multimorbidity: systematic review and meta-analysis of 193 international studies

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ABSTRACT

Objective (1) To estimate the pooled prevalence of multimorbidity in all age groups, globally. (2) To examine how measurement of multimorbidity impacted the estimated prevalence.

Methods In this systematic review and meta-analysis, we conducted searches in nine bibliographic databases (PsycINFO, Embase, Global Health, Medline, Scopus, Web of Science, Cochrane Library, CINAHL and ProQuest Dissertations and Theses Global) for prevalence studies published between database inception and 21 January 2020. Studies reporting the prevalence of multimorbidity (in all age groups and in community, primary care, care home and hospital settings) were included. Studies with an index condition or those that did not include people with no long-term conditions in the denominator were excluded. Retrieved studies were independently reviewed by two reviewers, and relevant data were extracted using predesigned pro forma. We used meta-analysis to pool the estimated prevalence of multimorbidity across studies, and used random-effects meta-regression and subgroup analysis to examine the association of heterogeneous prevalence estimates with study and measure characteristics.

Results 13807 titles were screened, of which 193 met inclusion criteria for meta-analysis. The pooled prevalence of multimorbidity was 42.4% (95% Cl 38.9% to 46.0%) with high heterogeneity ($l^2 > 99\%$). In adjusted meta-regression models, participant mean age and the number of conditions included in a measure accounted for 47.8% of heterogeneity in effect sizes. The estimated prevalence of multimorbidity was significantly higher in studies with older adults and those that included larger numbers of conditions. There was no significant difference in estimated prevalence between low-income or middle-income countries (36.8%) and high-income countries (44.3%), or between self-report (40.0%) and administrative/clinical databases (52.7%).

Conclusions The pooled prevalence of multimorbidity was significantly higher in older populations and when studies included a larger number of baseline conditions. The findings suggest that, to improve study comparability and quality of reporting, future studies should use

Strengths and limitations of this study

- This study used meta-regression to examine the variation of estimated prevalence of multimorbidity and how measure and study characteristics influenced prevalence estimates.
- The use of multiple imputation in this study minimised biased estimates caused by missing values and unbalanced classes and enhanced statistical accuracy.
- The inclusion of studies with various measure and study characteristics enabled a better understanding of the contributing factors of the heterogeneity of multimorbidity prevalence.
- Due to inconsistent reporting of multimorbidity prevalence and data unavailability, the estimated multimorbidity prevalence stratified by sex, ethnicity and socioeconomic status could not be explored in this study.

a common core conditions set for multimorbidity measurement and report multimorbidity prevalence stratified by sociodemographics. **PROSPERO registration number** CRD42020172409.

INTRODUCTION

Population ageing is a worldwide phenomenon, with WHO estimating that the proportion of the global population aged 60 years and older will double from 12% to 22% between 2015 and 2050.¹ A key implication of population ageing is that increasing numbers of people will be living with multimorbidity. Multimorbidity, commonly defined as the co-occurrence of two or more long-term conditions,² adversely affects people's risk of death, health-related quality of life, functional ability and mental well-being.^{3 4} Multimorbdiity affects all groups of society, but is known

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to be more common in older people, in women and in those from low socioeconomic backgrounds, particularly in high-income countries.⁵⁻⁷ In low-income and middleincome countries (LMICs), people living in urban areas, on the other hand, were found to have a higher rate of multimorbidity prevalence.⁸ Multimorbidity poses major challenges to the delivery of care in health systems internationally, which are often focused on the management of single diseases and lack appropriate coordination and continuity of care across different sectors.^{9 10} Disparities in health and health and social care could be found at any stage along the continuum of chronic diseases, from prevention to the management of diseases. To understand these disparities among multimorbid populations, it requires consistently monitoring the populations (eg, incidence, prevalence, health impact, risk factors and delivery of care) defined by race and ethnicity, gender, age, socioeconomic status, physical environment and geographic factors.

Previous systematic reviews have identified issues in the measurement of multimorbidity, related to the choice of chronic conditions counted in measures, the categorisation of conditions and diseases and the counting or weighting method used.¹¹⁻¹³ Although weighted measures are often used when the purpose of measurement is to predict future outcomes, a simple count of conditions remains the most commonly used method for the measurement of multimorbidity, and is optimal for estimating multimorbidity prevalence.^{13 14} However, the estimated prevalence of multimorbidity varies widely in the literature ranging from 3.5% to 100%,¹⁵ likely reflecting a combination of varying measures and varying populations studied.¹⁶ Much of the research up to now has not quantitatively investigated the variation in multimorbidity prevalence and its influencing factors in much detail. Understanding the links between prevalence estimates and measurement approaches can better inform and support future development of multimorbidity measurement guidelines. Therefore, this review aimed to examine the pooled prevalence of multimorbidity in all age groups, globally and how measurement of multimorbidity impacted the estimated prevalence.

Research questions

- What is the pooled prevalence of multimorbidity and does it differ between different age groups?
- What are the factors that influenced the variation in prevalence estimates across studies?

METHODS

The systematic review and meta-analysis reported here is part of a larger review which aimed to examine (1) how multimorbidity has been constructed, (2) measured by international studies and (3) variation in the estimated prevalence of multimorbidity across studies. Analysis in relation to the first two registered objectives has been reported,¹³ and this paper reports the third registered objective. The PROSPERO registration number for this paper is therefore the same as for the first published paper from this work. 13

Inclusion and exclusion criteria

The eligibility criteria for this review were defined based on the CoCoPop framework-condition, context and population.¹⁷ The condition included in this review is prevalence of multimorbidity. The majority of studies defined multimorbidity as the co-existence of two or more chronic conditions, and used the cut-off to estimate its prevalence in a population of interest. We therefore included studies that used this definition for examining multimorbidity prevalence across international studies. For this analysis, we included studies carried out in the community, primary care, care home and hospitals and those estimating the prevalence of multimorbidity in the population studied. Studies that did not include a relevant denominator population-for example, only examining patients with an index condition or excluding patients who did not have multimorbidity-were excluded. Qualitative research, studies not published in English and conference abstracts were also excluded.

Search strategy

The search strategy for this review was developed in collaboration with a specialist medical librarian (online supplemental table S1). Key terms relevant to multimorbidity and measurement were combined using Boolean logic to identify studies that met the inclusion criteria. We included medical subject headings to provide a sensitive search for relevant literature. Databases included in the search were Ovid interface (PsycINFO, Embase, Global Health, Medline), Scopus, Web of Science, Cochrane Library, EBSCO interface (CINAHL Plus) and ProQuest Dissertations and Theses Global, from inception to 21 January 2020 (we are not aware of any large recently published studies since that date). In addition to the database searches, our secondary search strategy included hand-searching reference lists of retrieved articles and tracked citations to maximise the yield.

Study screening and selection

Articles retrieved from databases were organised using EndNote X9 bibliographic software and Excel, and then were imported to Covidence for screening.¹⁸ Titles, abstracts and full-texts of retrieved articles were screened against the eligibility criteria by two reviewers. Throughout the review process, any disagreement that arose was resolved through discussion between the two reviewers (IS-SH and PH), and through the involvement of a third reviewer (BG) if necessary. The study selection process is summarised in figure 1.

Data extraction

We extracted data on the characteristics of the included studies using predesigned data extraction pro forma. The extracted data included (1) authors, (2) publication year, (3) study purpose, (4) method, (5) country, (6), continent, (7) country income (classified as 'high' and

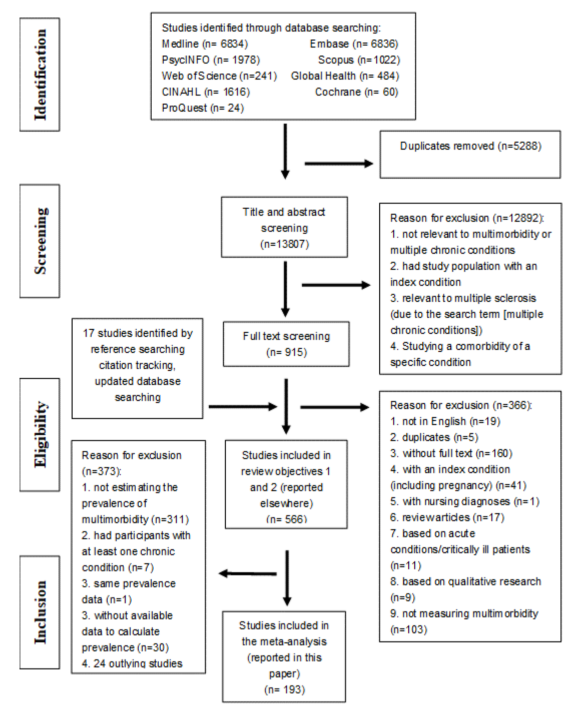


Figure 1 Preferred Reporting Items for Systematic Reviews and Meta-Analyses flow diagram.

'low or medium' (combined because of small numbers) allocated based on the World Bank Group at the time of review¹⁹), (8) study participants, (9) mean age, (10) sample size, (11) number of conditions, (12) setting, (13) data collection method/data source, (14) number of multimorbidity cases and (15) proportion of multimorbidity (calculated based on item 10 and 14). Data on the estimated prevalence stratified by sex, ethnicity and socioeconomic status were fragmented and unavailable in many studies, and thus these could not be retrieved for analyses.

Risk of bias assessment

We used the Effective Public Health Practice Project quality assessment tool for quantitative studies to assess the risk of bias and the quality of each of the included studies, in terms of (1) selection bias, (2) study design, (3) confounders, (4) blinding, (5) data collection method, (6) withdrawals and dropouts.²⁰ We assessed also publication bias (rated high if there was selective reporting within studies) and conflict of interest (rated unclear if conflict of interest declaration was not reported). Each study was rated and assigned an overall risk of bias as

'high', 'moderate' or 'low' (see the details in online supplemental appendix p. 26).

Data analysis

Descriptive statistics were used to summarise study characteristics. Since distributions were skewed, median and IQR were used to measure the central tendency and examine variability of variables such as mean age and number of conditions. Categorical (eg, continent, study population and data source) and ordinal data (eg, country income and risk of bias) were examined using frequency tables. To investigate the association between continuous/count predictor (mean age/number of conditions) and categorical predictors, univariate generalised linear models were used. We summarised the prevalence of multimorbidity using metaprop.^{21 22} The presence of effect size heterogeneity was examined using the Q statistic and I². Significant heterogeneity was identified, so we used subgroup analysis and meta-regression with random-effects models to identify potential moderating factors.

Outlying studies were identified using studentised residuals, leave-one-out analysis and Mahalanobis distance. Studies with studentised residuals that were larger than two or three and those that contributed to heterogeneity in leave-one-out analyses were scrutinised.²³ Mahalanobis distance was used for pattern recognition and multivariate outlier detection.²⁴ Study effect sizes were graphically displayed to identify outlying studies and explore subgroup effects (online supplemental figure S1). In initial analysis of heterogeneity and outliers, 24 studies were found to make a significant contribution to the high level of observed heterogeneity in multimorbidity prevalence and significant changes in the summary effect size. The 24 studies were excluded for one or more of the following reasons: (1) their contribution to high levels of heterogeneity in the leave-one-out test, (2) being identified as an outlying value in the studentised residuals test $(z-score \geq 2)$, (3) their Mahalanobis distance exceeding the χ^2 critical value at a 0.01 significance level, (4) infrequent values in compositional categorical data (eg, only one study examined prevalence in children). The process of identifying outliers, the rationale for exclusion of each study and the characteristics of outlying studies are documented in online supplemental figure S2 ane table S2 and online supplemental table S3. Sensitivity analysis was performed to explore the impact of excluding the 24 studies in meta-analysis.

There was missingness in two predictors, with 37% missingness in the 'mean age' of the study population variable (some of which reported it categorically, and thus were treated as missing data) and 6% missingness in the 'number of conditions' included in the multimorbidity measure variable. Previous research has shown that complete case removal (removing missing data in a data set) in meta-regression could lead to biased coefficient estimates of predictors (varied widely from complete-data estimates), whereas multiple imputation was found to perform well at generating estimates that were close

to complete-data estimates.²⁵ Therefore, in this review, multiple imputation with 60 imputed datasets and 10 iterations was conducted where random forest was used to impute missing data.^{26 27} Following multiple imputation, fraction of missing information was computed to quantify the impact of missing data, which ranged from 0.05 to 0.3 indicating that the uncertainty in the values imputed for missing data is small/moderate.²⁸

A random-effects regression tree approach with 10-fold cross-validation was used to identify subgroups (cut-offs) of the 'mean age' and 'number of conditions' variables with differential effect sizes.²⁹ Given considerable variation in the effect sizes, we conducted meta-regression with the restricted maximum likelihood (REML) estimator to examine the possible sources of heterogeneity in effect sizes.^{21 22 30} As the variable 'multimorbidity prevalence' did not follow the normal distribution (positively skewed), we applied logit transformation to the variable for analyses and converted the logits back to ORs (e^{logit}) and proportions $(p=e^{logit}/e^{logit}+1)$ for reporting. For model selection, we refitted the models using maximum likelihood and then conducted a log-likelihood test to compare the fit of models.³¹ A permutation test with 1000 permuted datasets was conducted to validate the robustness of the final model by rearranging and shuffling the order of the data and re-calculating p values to check whether there is type 1 error.³² Subgroup analysis with the REML method was used to estimate the pooled multimorbidity prevalence of subgroups of each variable (age, the number of conditions included in a measure, setting, data source, continent, country income, study risk of bias). Forest-like plots were used to display the effect sizes of included studies.³³ The presence of publication bias was assessed using Egger's test, which did not find evidence of publication bias.³⁴ All statistical tests were performed using R V.4.0.4.

Patients and public involvement

No patients were involved in the development of the research question, outcome measures, study design and implementation. Nonetheless, we have previously discussed preliminary review findings and issues relevant to multimorbidity measurement with our patient and public involvement group. We plan to disseminate the review findings to researchers, clinicians, policy makers and public audiences through news media, social media and seminars.

RESULTS

After screening 13807 titles and abstracts, 217 studies were identified which estimated the prevalence of multimorbidity using a cut-off of 'two or more' conditions. Following the removal of 24 outlying studies, 193 studies were included in the meta-analysis (table 1, online supplemental table S4). Of the 193 studies, 64 studies were from Europe, 47 from North America, 44 from Asia, 11 from Australasia, 12 from South America and 4 from Africa (table 1 and figure 2).

Table 1	Summary of study characteristics (online	
supplem	ental table S8 shows the definition of variable	es)

supplemental table 58 shows the	· · · · · ·
Name of variable	Descriptive statistics (n=193)
Prevalence of multimorbidity (%)	Range: 2.7–95.6 Pooled prevalence with the REML estimator: 42.4 (95% Cl 38.9 to 46.0)
Mean age of study population (year)	Range of mean age: 32.2–83.8 Median of mean age: 62.6 (Q1, Q3: 50.1, 72.4)
No. of conditions (count)	Range: 3–60 Median: 13 (Q1, Q3: 9, 19)
Country income (count, %)	
High income	145 (75.1%)
Low income or middle income	48 (24.9%)
Continent (count, %)	
Europe	64 (33.2%)
North America	47 (24.4%)
Asia	44 (22.8%)
Australasia	11 (5.7%)
South America	12 (6.2%)
Africa	4 (2.1%)
Multiple continents	11 (5.7%)
Study population (count, %)	
Only older people	63 (32.6%)
Middle-aged and older	46 (23.8%)
All adults	84 (43.5%)
Setting (count, %)	
Community	147 (76.2%)
Primary care	32 (16.6%)
Hospital	14 (7.3%)
Source (count, %)	
Self-report	150 (77.7%)
Database	43 (22.3%)
Risk of bias assessment (count, %)	
Low	9 (4.7%)
Moderate	162 (83.9%)
High	22 (11.4%)

The percentages were rounded so they do not add to 100%. REML, restricted maximum likelihood.

Seventy-five per cent of studies were from high-income countries (n=145) and 24.9% from LMICs (1 from low-income, 8 from lower middle-income, 29 from upper middle-income and 10 from multiple LMICs). The majority of studies (n=147) estimated the prevalence of multimorbidity in community settings, followed by primary care (n=32) and hospital setting (n=14).

Prevalence data were collected through either self-report (n=150) or medical records and administrative databases (n=43). In a univariate linear regression (online supplemental table S5), we found that studies from Europe, database studies and studies conducted in hospital settings were more likely to measure multimorbidity in an older population and included a larger number of conditions in a multimorbidity measure, compared with those from other continents, self-report studies and studies conducted in primary care and community settings. In respect to risk of bias in included studies (online supplemental table S6 and figure S3), 11.4% were rated as high risk of bias, 83.9% as moderate risk of bias and 4.7% as low risk of bias.

The pooled estimate of multimorbidity prevalence across the 193 studies was 42.4% (95% CI 38.9% to 46.0%), τ^2 is 1.0 (95% CI 0.9 to 1.3) with high heterogeneity ($I^2 > 99\%$) and meta-regression was therefore used to examine study characteristics associated with heterogeneity. Mean age (F=89.8, p<0.0001, R²=31.7%) and number of conditions (F=39.2, p<0.0001, R²=16.7%) were the strongest univariate predictors and positively associated with the estimated prevalence of multimorbidity (figure 3). Meta-regression tree analysis (online supplemental figure S4) partitioned the mean age variable into three homogeneous subgroups (aged <59, 59-73 and ≥74 years) and the number of conditions variable into four homogeneous subgroups (<9, 9–19, 20–43, \geq 44). The categorical 'mean age' and 'number of conditions' variables explained 35.9% and 19.5% of the heterogeneity in effect sizes, respectively (larger than the original numerical variables). Therefore, the categorical variables identified from the regression trees for meta-analyses were used for meta-regression.

In univariate meta-regression, primary care studies (pooled multimorbidity prevalence 50.5%, OR 1.6, 95% CI 1.1 to 2.3) and hospital-based studies (pooled multimorbidity prevalence 59.6%, OR 2.3, 95% CI 1.3 to 4.0) had significantly higher rates of multimorbidity than community-based studies (39.1%) (table 2). Multimorbidity prevalence was significantly higher in database studies (pooled multimorbidity prevalence 52.7%, OR 1.7, 95% CI 1.2 to 2.4) than self-reported studies (pooled multimorbidity prevalence 40.0%). In the mean age categorical variable, the pooled prevalence estimates of the three subgroups were statistically significantly different from one another, and considerably higher in studies with mean participant age ≥ 74 years (pooled multimorbidity prevalence 67.0%, OR 5.2, 95% CI 3.8 to 7.2) and mean participant age 59-73 years (pooled multimorbidity prevalence 47.6%, OR 2.3, 95% CI 1.8 to 3.0) than those with mean participant age <59 years (pooled multimorbidity prevalence 28.0%) (table 2 and figure 4). Similar patterns were also found in the number of conditions variable where studies including \geq 44 conditions in measurement (pooled multimorbidity prevalence 87.6%, OR 16.5, 95% CI 6.4 to 42.6), 20-43 conditions (pooled multimorbidity prevalence 52.1%, OR 2.5, 95% CI 1.7 to 3.7)

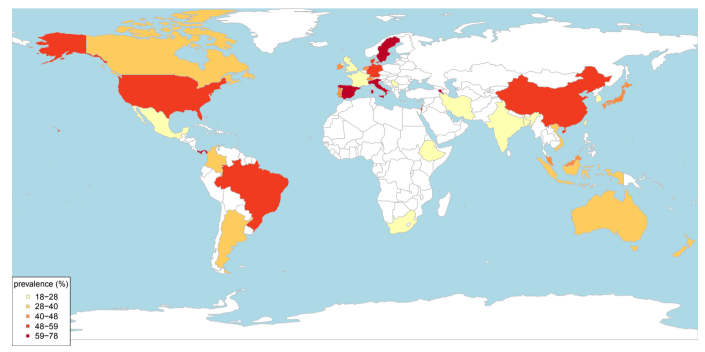


Figure 2 Country of origin of the included studies estimating the prevalence of multimorbidity (except studies from multiple countries).

and 9–19 conditions (pooled multimorbidity prevalence 43.7%, OR 1.8, 95% CI 1.3 to 2.5) yielded higher prevalence estimates than studies including <9 conditions in measurement (pooled multimorbidity prevalence 30.1%) with a dose-response relationship. The estimated prevalence of multimorbidity was 44.3% in high-income countries compared with 36.8% in LMICs, but the difference was not statistically significantly different (OR 1.4, 95% CI 1.0 to 1.9). In study risk of bias, no statistically significant difference in pooled prevalence of multimorbidity was

found between studies with low, moderate and high risk of bias.

In the adjusted meta-regression model, compared with studies where participant mean age was <59 years, multimorbidity prevalence remained significantly higher in studies with mean participant age 59–73 years (OR 2.2, 95% CI 1.7 to 2.8) and in studies with mean participant age \geq 74 years (OR 4.4, 95% CI 3.3 to 5.9). Compared with measures including <9 conditions, multimorbidity prevalence was higher in measures including \geq 44 conditions

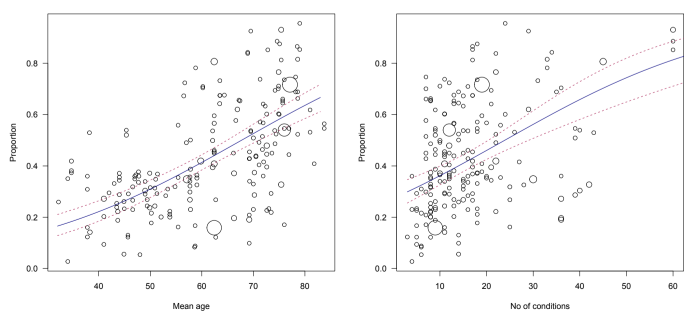


Figure 3 Relationship between the prevalence of multimorbidity and mean age or number of conditions (the area of points is proportional to inverse variances).

Table 2 Output of meta-a				
	Pooled prevalence of multimorbidity of each subgroup (%, 95% CI)	Meta-regression Unadjusted OR (95% CI)	Meta-regression Adjusted OR (95% CI) R ² 54.3%	FMI
Group of mean age (years)		R ² 35.9%		
<59	28.0 (24.9 to 31.5)	Ref	Ref	Ref
59–73	47.6 (42.5 to 52.8)	2.3 (1.8 to 3.0)***	2.2 (1.7 to 2.8)***	0.3
≥74	67.0 (60.4 to 72.9)	5.2 (3.8 to 7.2)***	4.4 (3.3 to 5.9)***	0.2
No. of conditions		R ² 19.5%		
<9	30.1 (24.9 to 35.7)	Ref	Ref	Ref
9–19	43.7 (39.5 to 48.0)	1.8 (1.3 to 2.5)***	1.8 (1.4 to 2.3)***	0.1
20–43	52.1 (43.8 to 60.3)	2.5 (1.7 to 3.7)***	2.3 (1.6 to 3.2)***	0.2
≥44	87.6 (81.3 to 92.0)	16.5 (6.4 to 42.6)***	8.2 (3.8 to 17.5)***	0.06
Setting		R ² 5.1%		
Community	39.1 (35.5 to 42.8)	Ref	Ref	Ref
Primary care	50.5 (39.6 to 61.3)	1.6 (1.1 to 2.3)*	1.6 (1.1 to 2.3)**	0.2
Hospital	59.6 (45.6 to 72.2)	2.3 (1.3 to 4.0)**	1.5 (1.0 to 2.4)	0.2
Source		R ² 4.0%		
Self-report	40.0 (36.2 to 43.8)	Ref	Ref	Ref
Database	52.7 (45.2 to 60.1)	1.7 (1.2 to 2.4)**	0.7 (0.5 to 1.0)	0.2
Continent		R ² 6.8%		
North America	50.4 (43.6 to 57.3)	Ref	Ref	Ref
Europe	44.8 (38.2 to 51.5)	0.8 (0.5 to 1.2)	0.5 (0.4 to 0.7)***	0.1
Australasia	35.8 (29.5 to 42.5)	0.5 (0.3 to 1.1)	0.5 (0.3 to 0.8)**	0.08
Asia	35.3 (29.3 to 42.0)	0.5 (0.4 to 0.8)**	0.6 (0.4 to 0.8)***	0.1
South America	47.5 (31.2 to 64.4)	0.9 (0.5 to 1.7)	0.8 (0.5 to 1.3)	0.1
Africa	13.8 (4.5 to 32.8)	0.2 (0.06 to 0.4)***	0.3 (0.1 to 0.6)**	0.1
Multiple continents	38.4 (29.1 to 48.6)	0.6 (0.3 to 1.2)	0.7 (0.4 to 1.1)	0.1
Country income		R ² 1.2%		
Low-income or middle- income	36.8 (29.7 to 44.4)	Ref		
High-income	44.3 (40.3 to 48.4)	1.4 (1.0 to 1.9)		
Study risk of bias		R ² 0.0%		
Low risk	33.3 (20.2 to 49.6)	Ref		
Moderate risk	42.4 (38.6 to 46.3)	1.5 (0.7 to 3.0)		
High risk	46.4 (34.1 to 59.1)	1.7 (0.8 to 3.9)		
Publication year		1.0 (1.0 to 1.0)		
*P<0.05 **p<0.01 ***p<0.001				

*P<0.05, **p<0.01, ***p<0.001,.

.FMI, fraction of missing information; Ref, reference category.

(OR 8.2, 95% CI 3.8 to 17.5), 20–43 conditions (OR 2.3, 95% CI 1.6 to 3.2) and 9–19 conditions (OR 1.8, 95% CI 1.4 to 2.3). In respect to study settings, the pooled prevalence was significantly higher in primary care settings compared with community settings (OR 1.6, 95% CI 1.1 to 2.3). Compared with studies from North America, prevalence was lower in studies from Europe (OR 0.5, 95% CI 0.4 to 0.7), Australasia (OR 0.5, 95% CI 0.3 to 0.8), Asia (OR 0.6, 95% CI 0.4 to 0.8) or Africa (OR 0.3 95% CI 0.1

to 0.6). No significant difference in prevalence estimates between self-report and routine database studies was evident after controlling for study and measure characteristics. The model explained 54.3% of the heterogeneity in multimorbidity prevalence, with the mean age and number of conditions variables providing most explanatory power (47.8% of the heterogeneity).

Sensitivity analysis including the 24 outlying studies (online supplemental table S7) was similar to primary

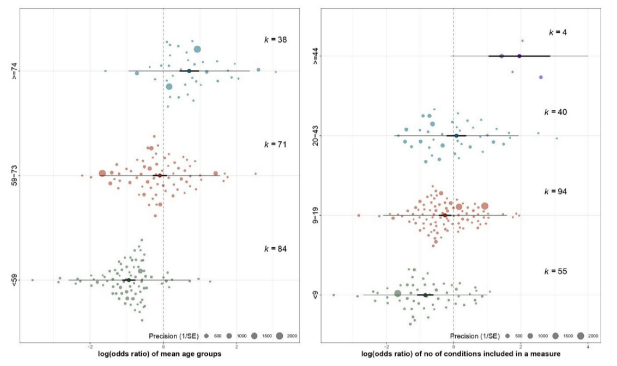


Figure 4 The distribution of prevalence estimates within the subgroups of mean age and number of conditions (forest-like plot for a large review).

analysis except for 'number of conditions' variable. The mean participant age and number of conditions variables remained the strongest predictors of multimorbidity prevalence in sensitivity analysis. However, the estimated prevalence in sensitivity analysis (including outlying studies) was much lower in studies including \geq 44 conditions in a multimorbidity measure (pooled multimorbidity prevalence 54.5, OR 2.8, 95% CI 1.5 to 5.4) compared with primary analysis excluding outlying studies (pooled multimorbidity prevalence 54.6). The difference in estimates was mainly attributed to the three outlying studies that included 146, 147 and 259 conditions in a measure respectively but yielded relatively low mean multimorbidity prevalence (mean prevalence 54.3%).^{35–37}

DISCUSSION

The overall estimate of multimorbidity prevalence in adults across all the included studies was 42.4% (95% CI 38.9% to 46.0%), but with very high heterogeneity. More than half of the observed heterogeneity was explained by study mean participant age and the number of conditions included in the multimorbidity measure, with older age and larger number of conditions strongly associated with a higher prevalence of multimorbidity. The difference in estimated prevalence was small between self-reported and administrative/clinical databases, and between study settings. No significant difference was found between studies from LMICs and high-income countries, but North American studies had higher estimated prevalence

and African studies had the lowest estimated prevalence than other continents.

Three prior systematic reviews examined the prevalence of multimorbidity across studies.³⁸⁻⁴⁰ Fortin et al⁸⁸ and Violan et al⁴⁰ conducted a narrative review and found various operationalisations of multimorbidity and a large variation in the prevalence of multimorbidity, particularly in studies with older adult populations or those with low socioeconomic status.^{38 40} Nguyen *et al*³⁹ meta-analysed the prevalence of multimorbidity across 70 studies from community settings and found that the pooled estimated prevalence was 33.1% with high levels of heterogeneity $(I^2 > 99\%)$.³⁹ The pooled prevalence of multimorbidity in the study by Nguyen et al is lower than in this study, likely because we have included studies from primary care and hospital settings (the pooled prevalence of multimorbidity in community-based studies in this analysis was 39.5%). Nguyen *et at*³⁹ did not carry out a metaregression, but in narrative analysis comment that the prevalence of multimorbidity appeared higher in older adults and women.³⁹ Our review findings are consistent with previous literature finding that age is most important determinant of multimorbidity.^{5 38 39 41} While we did not find a significant difference between LMICs and highincome countries, Nguyen et al in their review showed a statistically significantly higher pooled prevalence in high-income countries (the pooled prevalence from 18 studies was 37% compared with 36.8% in this review of 145 studies) than LMICs (the pooled prevalence from 31 studies was 29% compared with 44.3% in this review of 48 studies). This difference in findings may be due to the

inclusion in our review of a larger number of studies from high-income or upper middle-income countries. The low number of included studies from low-income countries in this review could be explained by less attention paid to this relatively new research field (multimorbidity) in lowincome countries and our literature search restricted to English language (proficient language of reviewers). The estimated prevalence of multimorbidity in North America was higher compared with other continents in this study despite older study populations and larger numbers of conditions found in studies from Europe. A possible explanation for the higher prevalence in North America is that private or insurance-based healthcare systems are more likely to code conditions since it affects remuneration, as well as cultural differences in relation to overdiagnosis and medicalisation.⁴² On the other hand, the lower estimated multimorbidity prevalence in African studies could be attributed to the predominance of infectious diseases and inadequate access to medical care including diagnostic services.⁴³

The strengths of this review are searches conducted in multiple databases, the large number of studies identified and the use of meta-analytic approaches to examine factors associated with heterogeneity of estimated multimorbidity prevalence. We examined and handled outlying studies and missing data (multiple imputation) with rigour and excluded studies that did not take into account 'healthy' populations (populations with no long-term conditions) to minimise biased estimates of multimorbidity prevalence. This review has limitations. Sensitivity analysis including all studies had similar findings with one exception, namely that sensitivity analysis found a weaker (but still statistically significant) association with the number of conditions included in the multimorbidity measure than primary analysis. Although we examined associations with study characteristics including mean participant age, a limitation is the lack of information in the reviewed studies on prevalence estimates stratified by participant characteristics including sex, ethnicity and socioeconomic status. An additional uncontrolled factor is how studies measured multimorbidity in terms of the type (as opposed to the number) of the conditions included in measures, which varied substantially across studies with too much heterogeneity to model.¹³ The exclusion of non-English studies in this review may also limit the generalisability of the research findings. Last but not least, measurement of multimorbidity is a relatively new research field and its labelling has been used variably. Thus, it is likely that not all relevant studies were identified and included in this review, but we were rigorous in our application of inclusion/exclusion criteria and did not favour adding known papers that did not appear in the search or where excluded through the process.

In spite of the methodological limitations, this review adds to our understanding of how study and measure characteristics can influence the estimated prevalence of multimorbidity. Mean age of the study population and the number of conditions included in the multimorbidity measure were the major factors associated with varying estimated prevalence of multimorbidity. A key implication is that comparing prevalence between studies requires more stratified estimates of multimorbidity prevalence. We therefore strongly recommend that as well as overall prevalence, future studies should clearly report multimorbidity prevalence stratified by age, in 5-year age bands to ensure granularity and by sex at a minimum, and ideally by ethnicity and socioeconomic status. This will allow readers to capture a more holistic picture of multimorbidity prevalence in the population studied, and allow better comparison of prevalence in different populations, and accurate pooled estimates of prevalence in reviews.

Additionally, the number of conditions included in a measure is strongly associated with estimated multimorbidity prevalence. It would be ideal if studies additionally reported prevalence using a common core set of conditions agreed by consensus. Parallel reporting of the bespoke set chosen for the context and purpose, and a core set would improve comparability of prevalence estimates, and help identify the additional value of any bespoke multimorbidity measures. The lack of any significant difference in estimated prevalence between self-report and clinical/administrative databases in this review suggests that provided careful attention is paid to the number and type of conditions included in measures, exactly how data are collected may be less important.

To conclude, in recent years, there has been an increasing interest in the epidemiology of multimorbidity internationally. This review finds that population characteristics and measurement content are the major factors that influenced prevalence estimates of multimorbidity. Studies with older populations and larger numbers of conditions yielded a higher estimate of multimorbidity prevalence. However, heterogeneity between studies has made comparison of multimorbidity prevalence across studies difficult. To improve comparability and quality of reporting, this review suggests that future studies should use common core condition set for the measurement of multimorbidity and clearly report the prevalence of multimorbidity stratified by sociodemographics.

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Supplementary appendix

Supplement to: Ho ISS, Azcoaga-Lorenzo A, Akbari A, et al. Variation in the estimated prevalence of multimorbidty: systematic review and meta-analysis of 193 studies.

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Table S1: Search strategy

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Table S2: Summary of the characteristics of outlying studies

Name of variable	Outlying studies (n=24)	All studies (n=217)
Prevalence of multimorbidity (%)	Range: 7.3 to 89.1	Range: 2.7-95.6
	Pooled prevalence with the REML	Pooled prevalence with the REML
	estimator: 31.0 (21.6-42.2)	estimator: 41.1 (37.7-44.6)
Mean age of study population (year)	Range of mean age: 39.6 to 82.2	Range of mean age: 32.2 to 83.8
	Median of mean age: 56.6 (Q1, Q3: 52.3,	Median of mean age: 62.4 (Q1,Q3:
	66.4)	50.2,72.0)
No of conditions (count)	Range: 7 to 259	Range: 3 to 259
	Median: 34 (Q1, Q3: 19.5, 54.5)	Median: 14.0 (Q1, Q3: 9, 21)
Country income (count, %)		
High income	21 (87.5%)	166 (76.5%)
Low- or Middle-income	3 (11.5%)	51 (23.5%)
Continent (count, %)		
Europe	6 (25.0%)	70 (32.3%)
North America	7 (29.2%)	54 (24.9%)
Asia	7 (29.2%)	51 (23.5%)
Australasia	3 (12.5%)	14 (6.5%)
Multiple continents	1 (4.2%)	12 (5.5%)
South America		12 (5.5%)
Africa		4 (1.8%)
Study population (count, %)		
Only older people	2 (8.3%)	65 (30.0%)
Middle-aged and older	1 (4.2%)	47 (21.7%)
All adults	15 (62.5%)	99 (45.6%)
Only children	1 (4.2%)	1 (0.5%)
All age population	5 (20.8%)	5 (2.3%)
Setting (count, %)		
Community	12 (50.0%)	159 (73.3%)
Primary care	7 (29.2%)	39 (18.0%)
Hospital	4 (16.7%)	18 (8.3%)
Care home	1 (4.2%)	1 (0.5%)
Source (count, %)		
Self-report	8 (33.3%)	158 (72.8%)
Database	16 (66.6%)	59 (27.2%)
Risk of bias assessment (count, %)		
Low	4 (16.7%)	13 (6.0%)
Moderate	19 (79.2%)	181 (83.4%)
High	1 (4.2%)	23 (10.6%)

IQR: Interquartile range. SD: Standard deviation. The percentages were rounded so they do not add up to 100%.

Table S3: Characteristics of 24 outlying studies

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias	Rationale for exclusion
¹ Stanley et al (2018)	New Zealand	Australasia	High	Hospitals	All adults	Not reported	3489747	Medical records and administrative database	61	275706	0.08	Moderate	Contributing to high levels of heterogeneity of effect sizes (Leave-one-out analysis) and the studentized residual of this study is more than 2 standard deviations away from its expected value.
² Lenzi et al (2016)	Italy	Europe	High	Hospitals	All adults	66.4	3759836	Medical records and administrative database	26	574208	0.15	Moderate	Contributing to high levels of heterogeneity of effect sizes (Leave-one-out analysis)
³ Hu et al (2019)	Taiwan	Asia	High	Community	All adults	Not reported	1429527	Medical records and administrative database	20	939485	0.66	Low	Contributing to high levels of heterogeneity of effect sizes (Leave-one-out analysis)
⁴ Gawron et al (2020)	USA	North America	High	Hospitals	All adults but not older people	Not reported	741612	Medical records and administrative database	Not reported	53824	0.07	Moderate	Contributing to high levels of heterogeneity of effect sizes (Leave-one-out analysis) and the studentized residual of this study is more than 2 standard deviations away from its expected value.
⁵ Low et al (2019)	Singapore	Asia	High	Community	All adults	39.6	1181024	Self-report	48	309428	0.26	Low	Contributing to high levels of heterogeneity of effect sizes (Leave-one-out analysis)
⁶ Wang et al (2014)	China	Asia	Low or middle	Community	Whole population	Not reported	162464	Self-report	40	17987	0.11	Low	Contributing to high levels of heterogeneity of effect sizes (Leave-one-out analysis)
 ⁷ Gaulin et al (2019) 	Canada	North America	High	Hospitals	All adults	51.2	1316832	Medical records and administrative database	34	416282	0.32	Moderate	Contributing to high levels of heterogeneity of effect sizes (Leave-one-out analysis)

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias	Rationale for exclusion
⁸ Violan et al (2014)	Spain	Europe	High	Primary care	All adults	47.4	1356761	Medical records and administrative database	146	645818	0.48	Moderate	Contributing to high levels of heterogeneity of effect sizes (Leave-one-out analysis)
 ⁹ Nicholson et al (2019) 	Canada	North America	High	Primary care	All adults	52.3	367743	Medical records and administrative database	20	195838	0.53	High	Contributing to high levels of heterogeneity of effect sizes (Leave-one-out analysis)
¹⁰ Bao et al (2019)	China	Asia	Low or middle	Community	Middle aged and older	61.36	18137	Self-report	19	3773	0.21	Moderate	Contributing to high levels of heterogeneity of effect sizes (Leave-one-out analysis)
¹¹ Fortin et al (2005)	Canada	North America	High	Primary care	All adults	56.55	980	Medical records and administrative database	14	873	0.89	Moderate	The studentized residual of this study is more than 2 standard deviations away from its expected value.
¹² Prazeres et al (2015)	Portugal	Europe	High	Primary care	All adults	56.3	1993	Medical records and administrative database	147	1449	0.73	Moderate	Its Mahalanobis distance exceeds the chi-squared critical value at a 0.01 significance level (multivariate outlier detection)
¹³ Lawson et al (2013)	UK	Europe	High	Community	All adults	72.7	7054	Medical records and administrative database	40	1243	0.18	Moderate	Irregular patterns found in compositional data (in scatter plot and Mahalanobis distance test)- low prevalence in studies with high mean participant age and a larger number of conditions
¹⁴ Sullivan et al (2012)	USA	North America	High	Community	All adults	Not reported	47178	Medical records and administrative database	259	19666	0.42	Moderate	Its Mahalanobis distance exceeds the chi-squared critical value at a 0.01 significance level (multivariate outlier detection)

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias	Rationale for exclusion
¹⁵ Peng et al (2020)	China	Asia	Low or middle	Community	Only older people	71.6	1321	Self-report	15	589	0.45	Moderate	Contributing to high levels of heterogeneity of effect sizes (in leave-one-out analysis)
¹⁶ Excoffier et al (2018)	Switzerland	Europe	High	Primary care	All adults	56.5	2904	Medical records and administrative database	75	1513	0.52	Moderate	Its Mahalanobis distance exceeds the chi-squared critical value at a 0.01 significance level (multivariate outlier detection)
¹⁷ Chung et al (2015)	Hong Kong	Asia	High	Community	All adults	Not reported	25780	Self-report	46	3227	0.13	Moderate	Its Mahalanobis distance exceeds the chi-squared critical value at a 0.01 significance level (multivariate outlier detection)
¹⁸ Ki et al (2017)	South Korea	Asia	High	Community	All adults	57.05	19942	Medical records and administrative database	66	5979	0.30	Moderate	Its Mahalanobis distance exceeds the chi-squared critical value at a 0.01 significance level (multivariate outlier detection)
¹⁹ Bobo et al (2016)	USA	North America	High	Community	Whole population	Not reported	138858	Self-report	19	33682	0.24	Moderate	Infrequent values in compositional categorical data (few studies focused on whole population)
²⁰ Randall et al (2018)	Australia	Australasia	High	Community	Whole population	Not reported	5437018	Self-report	30	660449	0.12	Moderate	Infrequent values in compositional categorical data (few studies focused on whole population)
²¹ Russell et al (2020)	New Zealand	Australasia	High	Community	Only children	Not reported	3838	Self-report	7	374	0.10	Moderate	Infrequent values in compositional categorical data (only one study focused on children population)
²² Barnett et al (2012)	UK	Europe	High	Primary care	Whole population	Not reported	1751841	Medical records and administrative database	40	406427	0.23	Low	Infrequent values in compositional categorical data (few studies focused on whole population)

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias	Rationale for exclusion
²³ St Sauveret al(2015)	USA	North America	High	Primary care	Whole population	Not reported	106061	Medical records and administrative database	20	34592	0.33	Moderate	Infrequent values in compositional categorical data (few studies focused on whole population)
²⁴ Vetrano et al (2016)	Denmark, Finland, Iceland, Italy, the Netherlands, Norway, United Kingdom, Czech Republic, France, Sweden and Germany, Canada	Multiple continents	High	Care homes	Only older people	82.2	6903	Medical records and administrative database	13	5098	0.74	Moderate	Infrequent values in compositional categorical data (only one study focused on care home)

MM: Multimorbidity. No of participants: The total number of participants in the denominator for estimating prevalence in a study (which could be a subset in some included studies)

Table S4: Characteristics of 193 included studies

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias
²⁵ Aarts et al (2012)	The Netherlands	Europe	High	Primary care	All adults	55.4	1184	Medical records and administrative database	23	420	0.35	Moderate
²⁶ Aarts et al (2011a)	The Netherlands	Europe	High	Community	Middle aged and older	70	15188	Self-report	Not reported	7729	0.51	Moderate
²⁷ Aarts et al (2011b)	The Netherlands	Europe	High	Primary care	All adults	55.4	1763	Medical records and administrative database	23	985	0.56	Moderate
²⁸ Abizanda et al (2014)	Spain	Europe	High	Primary care	Only older people	78.6	842	Medical records and administrative database	14	580	0.69	Moderate
²⁹ Agborsangaya et al (2012)	Canada	North America	High	Community	All adults	46.6	4003	Self-report	16	919	0.23	Moderate
³⁰ Agborsangaya et al (2013)	Canada	North America	High	Community	All adults	47.8	4803	Self-report	16	1729	0.36	Moderate
³¹ Agborsangaya et al (2014)	Canada	North America	High	Community	All adults	47.7	4752	Self-report	16	1597	0.34	Moderate
³² Ahrenfeldt et al (2019)	Europe	Europe	High	Community	Middle aged and older	66.25	244258	Self-report	10	90652	0.37	Moderate
³³ Alimohammadian et al (2017)	Iran	Asia	Low or middle	Community	Middle aged and older	Not reported	49946	Self-report	8	10035	0.20	Moderate
³⁴ Angst et al (2002)	Switzerland	Europe	High	Primary care	All adults	Not reported	591	Medical records and administrative database	10	201	0.34	High

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³⁵ Appa et al (2014)	USA	North America	High	Community	All adults	60.2	1997	Self-report	16	1417	0.71	Moderate
³⁶ Adams et al (2017)	USA	North America	High	Community	All adults	Not reported	400000	Self-report	12	191600	0.48	Moderate
³⁷ Ahmadi et al (2016)	Iran	Asia	Low or middle	Community	Middle aged and older	52.1	49946	Self-report	8	10035	0.20	Moderate
³⁸ Amaral et al (2018)	Brazil	South America	Low or middle	Community	Only older people	Not reported	264	Self-report	8	175	0.66	Moderate
³⁹ An et al (2016)	South Korea	Asia	High	Community	Middle aged and older	54.8	10118	Self-report	8	3228	0.32	Moderate
⁴⁰ Araujo et al (2018)	Brazil	South America	Low or middle	Community	All adults	Not reported	4001	Self-report	12	1160	0.29	Moderate
⁴¹ Arnold-Reed et al (2018)	Australia	Australasia	High	Primary care	All adults	38.2	4285	Medical records and administrative database	43	2269	0.53	Moderate
⁴² Arokiasamy et al (2015)	6 low middle income countries (China, Ghana, India, Mexico, Russia, South Africa)	Multiple continents	Low or middle	Community	All adults	Not reported	42236	Self-report	8	9250	0.22	Moderate
⁴³ Sinnige et al (2015)	The Netherlands	Europe	High	Primary care	Middle aged and older	66.9	120480	Medical records and administrative database	29	74733	0.62	Moderate
⁴⁴ Zemedikun et al (2018)	UK	Europe	High	Community	Middle aged and older	Median age 58	502643	Medical records and administrative database	36	95710	0.19	Moderate
⁴⁵ Wensing et al (2001)	The Netherlands	Europe	High	Primary care	All adults	Not reported	3867	Self-report	25	626	0.16	Moderate
⁴⁶ Mounce et al (2018)	UK	Europe	High	Community	Middle aged and older	Not reported	4564	Self-report	15	1553	0.34	Moderate
⁴⁷ Taylor et al (2010)	Australia	Australasia	High	Community	All adults	Not reported	3206	Self-report	7	547	0.17	Low

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias
⁴⁸ Vancampfort et al (2019)	Six low and middle income countries (China, Ghana, India, Mexico, Russia, and South Africa)	Multiple continents	Low or middle	Community	Middle aged and older	62.4	34129	Self-report	11	15529	0.46	Moderate
⁴⁹ Vancampfort et al (2018)	Six low and middle income countries (China, Ghana, India, Mexico, Russia, and South Africa)	Multiple continents	Low or middle	Community	Only older people	72.6	14585	Self-report	11	8780	0.60	Moderate
⁵⁰ Aubert et al (2016)	Switzerland	Europe	High	Primary care	Middle aged and older	63.5	1002	Medical records and administrative database	17	676	0.67	Moderate
⁵¹ Autenrieth et al (2013)	Germany	Europe	High	Community	Only older people	75.7	1007	Self-report	13	658	0.65	Moderate
⁵² Bahler et al (2015)	Switzerland	Europe	High	Community	Only older people	74.9	229493	Medical records and administrative database	22	175752	0.77	Moderate
⁵³ Vancampfort et al (2017)	44 low and middle income countries	Multiple continents	Low or middle	Community	All adults	38.3	194431	Self-report	11	27518	0.14	Moderate
⁵⁴ Banjare et al (2014)	India	Asia	Low or middle	Community	Only older people	Not reported	310	Self-report	20	176	0.57	Moderate
⁵⁵ Barra et al (2015)	USA	North America	High	Community	All adults	45.36	43079	Self-report	Not reported	22412	0.52	Moderate
⁵⁶ Bernard et al (2016)	Australia	Australasia	High	Hospitals	Only older people	81.8	306	Medical records and administrative database	19	125	0.41	High
⁵⁷ Biswas et al (2019)	Bangladesh	Asia	Low or middle	Community	All adults	Not reported	8763	Self-report	3	1078	0.12	Moderate

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias
⁵⁸ Blakemore et al (2016)	UK	Europe	High	Primary care	Only older people	75	4377	Self-report	24	2631	0.60	Moderate
⁵⁹ Blyth et al (2008)	Australia	Australasia	High	Community	Only older people	76.9	1685	Self-report	18	920	0.55	Moderate
⁶⁰ Bowling et al (2019)	USA	North America	High	Community	Middle aged and older	56.7	4217	Self-report	12	3053	0.72	Moderate
⁶¹ Britt et al (2008)	Australia	Australasia	High	Primary care	All adults	Not reported	9156	Medical records and administrative database	18	3398	0.37	Moderate
⁶² Broeiro-Goncalves et al (2019)	Portugal	Europe	High	Hospitals	All adults	59.8	800376	Medical records and administrative database	22	335357	0.42	Moderate
⁶³ Bruce et al (2010)	Canada	North America	High	Community	All adults	37.8	453	Self-report	4	163	0.36	High
⁶⁴ Burgers et al (2010)	France, Germany, Canada, Australia, Netherlands, New Zealand, UK, USA	Multiple continents	High	Community	All adults	Not reported	8973	Self-report	7	4037	0.45	Moderate
⁶⁵ Burke et al (2017)	US, Europe, Asia	Multiple continents	High	Community	Only older people	Not reported	4668	Self-report	9	2165	0.46	Moderate
⁶⁶ Buurman et al (2016)	The Netherlands	Europe	High	Hospitals	Only older people	78.2	639	Medical records and administrative database	35	440	0.69	Moderate
⁶⁷ Calderon-Larranaga et al (2017)	Sweden	Europe	High	Primary care	Only older people	74.6	3363	Self-report	60	2980	0.89	Moderate
⁶⁸ Camargo-Casas et al (2018)	Colombia	South America	Low or middle	Community	Only older people	71.1	2000	Self-report	12	808	0.40	Moderate

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias
⁶⁹ Canevelli et al (2019)	Italy	Europe	High	Primary care	Only older people	75.1	185	Medical records and administrative database	18	162	0.88	High
⁷⁰ Chamberlain et al (2020)	USA	North America	High	Community	All adults	Not reported	198941	Self-report	21	78527	0.39	Low
⁷¹ Chen et al (2018)	China	Asia	Low or middle	Community	Only older people	Not reported	30774	Medical records and administrative database	33	25101	0.82	Low
⁷² Chen et al (2018)	China	Asia	Low or middle	Community	Middle aged and older	Not reported	3737	Self-report	16	1722	0.46	Moderate
⁷³ Cheung et al (2013)	Hong Kong (SAR of China)	Asia	High	Community	Middle aged and older	71.3	1145	Self-report	18	654	0.57	Moderate
⁷⁴ Chu et al (2018)	Hong Kong (SAR of China)	Asia	High	Primary care	Middle aged and older	Not reported	382	Medical records and administrative database	40	206	0.54	Moderate
 ⁷⁵ Chudasama et al (2019) 	UK	Europe	High	Community	Middle aged and older	Median age:58	491939	Medical records and administrative database	36	96622	0.20	Moderate
⁷⁶ Cimarras-Otal et al (2014)	Spain	Europe	High	Community	All adults	Not reported	22190	Self-report	20	7830	0.35	Moderate
⁷⁷ Chin et al (2016)	Hong Kong (SAR of China)	Asia	High	Primary care	All adults	Median age: 48	9259	Self-report	8	2350	0.25	Moderate
⁷⁸ Agrawal et al (2016)	India, China, Russia, Mexico, South Africa, Ghana	Multiple continents	Low or middle	Community	All adults	57.8	40166	Self-report	9	9238	0.23	Moderate
⁷⁹ Gu et al (2018)	China	Asia	Low or middle	Community	Only older people	Not reported	411	Self-report	13	232	0.56	Moderate
⁸⁰ Gunn et al (2012)	Australia	Australasia	High	Primary care	All adults	50.89	6864	Self-report	12	2154	0.31	Moderate

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias
⁸¹ Han et al (2013)	USA	North America	High	Primary care	Only older people	76	159	Medical records and administrative database	18	117	0.74	High
⁸² Hanlon et al (2018)	UK	Europe	High	Community	All adults	Not reported	493737	Medical records and administrative database	42	161576	0.33	Low
⁸³ Jantsch et al (2018)	Brazil	South America	Low or middle	Community	All adults	42	3092	Self-report	11	912	0.29	Moderate
⁸⁴ John et al (2003)	USA	North America	High	Community	Only older people	71.3	992	Self-report	11	732	0.74	High
⁸⁵ Johnson-Lawrence et al (2017)	USA	North America	High	Community	All adults	49.9	115097	Self-report	9	27278	0.24	Moderate
⁸⁶ Johnston et al (2019)	UK	Europe	High	Community	All adults	48	7184	Self-report	Not reported	388	0.05	Moderate
⁸⁷ Jones et al (2016)	USA	North America	High	Community	Only older people	Not reported	6964	Self-report	10	4951	0.71	Moderate
⁸⁸ Jovic et al (2016)	Serbia	Europe	Low or middle	Community	All adults	49.4	13103	Self-report	13	3522	0.27	Moderate
⁸⁹ Juul-Larsen et al (2020)	Denmark	Europe	High	Hospitals	Only older people	Median age: 78	369	Self-report	34	311	0.84	Moderate
⁹⁰ Hudon et al (2008)	Canada	North America	High	Community	All adults	Not reported	16782	Self-report	25	5343	0.32	Low
⁹¹ Hussain et al (2015)	Indonesia	Asia	Low or middle	Community	Middle aged and older	Not reported	9438	Self-report	12	3369	0.36	Moderate
⁹² Ie et al (2017)	USA	North America	High	Hospitals	Only older people	Not reported	1084	Medical records and administrative database	24	1036	0.96	High
⁹³ Ishizaki et al (2019)	Japan	Asia	High	Community	Only older people	76.9	2525	Self-report	9	1121	0.44	Moderate

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⁹⁴ Danon-Hersch et al (2012)	Switzerland	Europe	High	Community	Only older people	Not reported	1283	Self-report	12	448	0.35	Moderate
⁹⁵ de Heer et al (2013)	USA	North America	High	Community	All adults	47.72	1002	Self-report	19	378	0.38	Moderate
⁹⁶ Demirchyan et al (2013)	Armenia	Asia	Low or middle	Community	All adults	58.8	721	Self-report	Not reported	564	0.78	High
⁹⁷ Fabbri et al (2015)	Italy	Europe	High	Community	Only older people	73.6	1018	Self-report	15	458	0.45	Moderate
⁹⁸ Fillenbaum et al (2000)	USA	North America	High	Community	Only older people	73.44	4034	Self-report	5	1181	0.29	Moderate
⁹⁹ Kaneko et al (2019)	Japan	Asia	High	Community	Only older people	Not reported	253	Self-report	Not reported	135	0.53	Moderate
¹⁰⁰ Kang et al (2017)	South Korea	Asia	High	Primary care	All adults	32.2	590	Medical records and administrative database	14	153	0.26	Moderate
¹⁰¹ Gandhi et al (2020)	USA	North America	High	Community	All adults	Not reported	9499	Self-report	8	3379	0.36	Moderate
¹⁰² Costa et al (2018)	Brazil	South America	Low or middle	Community	Only older people	Not reported	1451	Self-report	29	1343	0.93	Moderate
¹⁰³ Rizzuto et al (2017)	Sweden	Europe	High	Community	Only older people	Not reported	1099	Self-report	36	774	0.70	Moderate
¹⁰⁴ Dhalwani et al (2017)	UK	Europe	High	Community	Middle aged and older	Not reported	5476	Self-report	18	1156	0.21	Moderate
¹⁰⁵ Elixhauser et al (1998)	USA	North America	High	Hospitals	All adults	57.1	1779167	Medical records and administrative database	30	619150	0.35	Low
¹⁰⁶ Fabbri et al (2015)	USA	North America	High	Hospitals	Only older people	72.3	695	Self-report	15	440	0.63	Moderate

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¹⁰⁷ Fortin et al (2014)	Canada	North America	High	Community	Middle aged and older	57.8	1196	Self-report	14	599	0.50	Moderate
¹⁰⁸ Fuchs et al (1998)	Israel	Asia	High	Community	Only older people	Not reported	1820	Self-report	14	1174	0.65	Moderate
¹⁰⁹ Galenkamp et al (2011)	The Netherlands	Europe	High	Community	Middle aged and older	69.2	2046	Self-report	7	876	0.43	High
(2016) 110 Galenkamp et al	Germany, UK, Italy, The Netherlands, Spain and Sweden	Europe	High	Community	Only older people	74.2	2792	Self-report	8	1358	0.49	Moderate
111 Gamma et al (2001)	Switzerland	Europe	High	Community	All adults	Not reported	407	Self-report	14	53	0.13	High
¹¹² Ge et al (2018)	Singapore	Asia	High	Community	All adults	51.4	1940	Self-report	17	715	0.37	Moderate
¹¹³ Ge et al (2019)	Singapore	Asia	High	Community	All adults	51.3	1932	Self-report	17	564	0.29	Moderate
¹¹⁴ Gould et al (2016)	USA	North America	High	Community	Only older people	74.82	4184	Self-report	7	2932	0.70	Moderate
¹¹⁵ Habib et al (2014)	Lebanon	Asia	Low or middle	Community	All adults	46.6	2501	Self-report	Not reported	665	0.27	Moderate
¹¹⁶ Harrison et al (2017)	Australia	Australasia	High	Primary care	All adults	Not reported	8707	Medical records and administrative database	28	2838	0.33	Moderate
¹¹⁷ Hayek et al (2017)	Israel	Asia	High	Community	All adults	47.2	4325	Self-report	10	1579	0.37	Moderate
Henninger et al (2012)	USA	North America	High	Community	Only older people	76	3212	Self-report	9	1753	0.55	Moderate
¹¹⁹ Hernandez et al (2019)	Ireland	Europe	High	Community	Middle aged and older	Not reported	6101	Self-report	31	4468	0.73	Moderate

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias
¹²⁰ Ho et al (2014)	Singapore	Asia	High	Community	Middle aged and older	66.15	1844	Self-report	12	830	0.45	Moderate
¹²¹ Khan et al (2019)	Bangladesh	Asia	Low or middle	Community	All adults	58.6	12338	Self-report	6	1031	0.08	Low
¹²² Kiliari et al (2013)	Cyprus	Europe	High	Community	All adults	53	465	Self-report	Not reported	132	0.28	Moderate
¹²³ King et al (2018)	USA	North America	High	Community	All adults	Not reported	5541	Self-report	11	3342	0.60	Moderate
¹²⁴ Kingston et al (2018)	UK	Europe	High	Community	All adults	Not reported	9723900	Self-report	12	5250906	0.54	High
¹²⁵ Koyanagi et al (2018)	China, Ghana, India, Mexico, Russia, and South Africa	Multiple continents	Low or middle	Community	Middle aged and older	62.1	32715	Self-report	10	16324	0.50	Moderate
¹²⁶ Kriegsman et al (2004)	The Netherlands	Europe	High	Community	Middle aged and older	69.2	2489	Self-report	7	519	0.21	Moderate
¹²⁷ Kristensen et al(2019)	Germany	Europe	High	Community	Middle aged and older	63.47	19605	Self-report	13	12600	0.64	Moderate
¹²⁸ Kristensen et al (2019)	Germany	Europe	High	Community	Middle aged and older	64.37	7604	Self-report	13	5140	0.68	Moderate
¹²⁹ Kunna et al (2017)	China, Ghana	Multiple continents	Low or middle	Community	Middle aged and older	Not reported	15864	Self-report	7	4731	0.30	Low
¹³⁰ Kuwornu et al (2014)	Canada	North America	High	Community	All adults	51.05	3284	Self-report	15	1143	0.35	Moderate
¹³¹ Lai et al (2019)	Hong Kong (SAR of China)	Asia	High	Community	All adults	Not reported	69636	Self-report	14	3898	0.06	Moderate
¹³² Lai et al (2018)	Hong Kong (SAR of China)	Asia	High	Community	All adults	Not reported	300	Self-report	11	48	0.16	Moderate
¹³³ Laires et al (2019)	Portugal	Europe	High	Community	All adults	Not reported	15196	Self-report	13	6671	0.44	Moderate

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias
¹³⁴ Lang et al (2015)	USA	North America	High	Community	Middle aged and older	53.4	3058	Self-report	6	948	0.31	Moderate
¹³⁵ Le Cossec et al (2016)	France	Europe	High	Community	Middle aged and older	70	15325	Self-report	4	3528	0.23	Moderate
¹³⁶ Lee et al (2007)	USA	North America	High	Hospitals	Middle aged and older	Not reported	741847	Medical records and administrative database	11	302792	0.41	Low
¹³⁷ Lee et al (2018)	Taiwan	Asia	High	Community	Only older people	Not reported	20898	Medical records and administrative database	Not reported	4234	0.20	High
¹³⁸ Li et al (2016)	UK	Europe	High	Primary care	All adults	Not reported	27806	Self-report	12	10332	0.37	Moderate
¹³⁹ Li et al (2019)	USA	North America	High	Community	Middle aged and older	67.4	14996	Self-report	8	9805	0.65	Moderate
¹⁴⁰ Lujic et al (2017)	Australia	Australasia	High	Community	Middle aged and older	70.2	90352	Self-report	8	33792	0.37	Moderate
¹⁴¹ Lupianez-Villanueva et al (2018)	14 European countries	Europe	High	Community	All adults	Not reported	14000	Self-report	13	3416	0.24	Moderate
¹⁴² Zhou et al (2018)	Bangladesh, India and China	Asia	Low or middle	Community	All adults	Not reported	18696	Self-report	9	3512	0.19	Moderate
¹⁴³ Zhang et al (2019)	China	Asia	Low or middle	Community	Only older people	70.5	11707	Self-report	11	5104	0.44	Moderate
¹⁴⁴ Wong et al (2010)	Canada	North America	High	Community	Only older people	Not reported	740	Self-report	7	489	0.66	Moderate
¹⁴⁵ Weimann et al (2016)	South Africa	Africa	Low or middle	Community	All adults	34	18526	Self-report	4	506	0.027	Moderate
¹⁴⁶ Wang et al (2017)	Australia	Australasia	High	Community	All adults	44	8820	Self-report	8	2539	0.29	Moderate
¹⁴⁷ Wang et al (2019)	South Africa	Africa	Low or middle	Community	Only older people	Not reported	2627	Self-report	5	439	0.17	Moderate

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias
¹⁴⁸ Wade et al (2019)	New Zealand	Australasia	High	Community	All adults	59.05	7654	Self-report	12	2786	0.36	Moderate
¹⁴⁹ Maciejewski et al (2019)	USA	North America	High	Community	Only older people	77.1	20124230	Medical records and administrative database	19	1442544 6	0.72	Moderate
¹⁵⁰ Marengoni et al (2016)	Sweden	Europe	High	Community	Only older people	74.4	3155	Medical records and administrative database	14	1654	0.52	Moderate
¹⁵¹ Marengoni et al (2009)	Sweden	Europe	High	Community	Only older people	Not reported	1099	Self-report	22	575	0.52	Moderate
¹⁵² Marques et al (2018)	13 European countries	Europe	High	Community	All adults	50.2	32931	Self-report	6	7113	0.22	Moderate
¹⁵³ Mavaddat et al (2014)	UK	Europe	High	Primary care	Middle aged and older	58.7	11439	Self-report	6	1006	0.09	Moderate
¹⁵⁴ McDaid et al (2013)	Ireland	Europe	High	Community	Middle aged and older	Not reported	6018	Self-report	8	733	0.12	High
¹⁵⁵ Melis et al (2014)	Sweden	Europe	High	Hospitals	Only older people	83.75	390	Medical records and administrative database	39	213	0.55	Moderate
¹⁵⁶ Min et al (2007)	USA	North America	High	Community	Only older people	81	372	Self-report	9	230	0.62	High
¹⁵⁷ Momtaz et al (2010)	Malaysia	Asia	High	Community	Only older people	69.26	385	Self-report	16	165	0.43	Moderate
¹⁵⁸ Mondor et al (2018)	Canada	North America	High	Community	All adults	Not reported	27195	Medical records and administrative database	17	11390	0.42	Moderate
¹⁵⁹ Muggah et al (2012)	Canada	North America	High	Community	All adults	Not reported	28450000	Medical records and administrative database	9	4523550	0.16	Moderate
¹⁶⁰ Nagel et al (2008)	Germany	Europe	High	Community	Middle aged and older	56.5	13781	Self-report	15	9275	0.67	Moderate

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias
¹⁶¹ Niedzwiedz et al(2019)	USA	North America	High	Community	Middle aged and older	67.2	2272	Self-report	8	1491	0.66	Moderate
¹⁶² Nunes et al (2016)	Brazil	South America	Low or middle	Community	All adults	45.75	2927	Self-report	11	852	0.29	Moderate
¹⁶³ Nunes et al (2017)	Brazil	South America	Low or middle	Community	All adults	43.7	60202	Self-report	22	13365	0.22	Moderate
¹⁶⁴ Nunes et al (2015)	Brazil	South America	Low or middle	Community	Only older people	Not reported	1593	Self-report	17	1295	0.81	Moderate
¹⁶⁵ Olaya et al (2017)	Spain	Europe	High	Community	Only older people	71.75	2113	Self-report	7	1088	0.51	Moderate
¹⁶⁶ Olivares et al (2017)	Argentina	South America	High	Community	All adults	43	1044	Self-report	Not reported	346	0.33	Moderate
¹⁶⁷ Park et al (2018)	South Korea	Asia	High	Community	Middle aged and older	62.7	5996	Self-report	25	1607	0.27	Moderate
¹⁶⁸ Patel et al (2006)	Mexico	South America	Low or middle	Community	Middle aged and older	73	7852	Self-report	5	1833	0.23	Moderate
¹⁶⁹ Pati et al (2016)	India	Asia	Low or middle	Community	All adults	44.96	103	Self-report	18	24	0.23	Moderate
¹⁷⁰ Pati et al (2019)	India	Asia	Low or middle	Primary care	All adults	44	1649	Self-report	21	567	0.34	Moderate
¹⁷¹ Payne et al (2013)	UK	Europe	High	Primary care	All adults	49	180815	Medical records and administrative database	40	54945	0.30	Moderate
¹⁷² Perez et al (2020)	Sweden	Europe	High	Community	Only older people	72.8	2596	Self-report	60	2213	0.85	Moderate
¹⁷³ Petersen et al (2019)	South Africa	Africa	Low or middle	Primary care	All adults	Not reported	2549	Self-report	Not reported	893	0.35	Moderate
174 Pfortmueller et al (2013)	Switzerland	Europe	High	Hospitals	All adults	Median age: 28	3170	Medical records and administrative database	18	1183	0.37	High

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias
¹⁷⁵ Pressley et al (1999)	USA	North America	High	Hospitals	Only older people	Not reported	5934	Medical records and administrative database	Not reported	3534	0.60	Moderate
¹⁷⁶ Prior et al (2016)	Denmark	Europe	High	Community	All adults	Not reported	118410	Self-report	39	33937	0.29	Moderate
¹⁷⁷ Ribeiro et al (2018)	Brazil	South America	High	Community	Only older people	70	820	Self-report	8	270	0.33	Moderate
¹⁷⁸ Ruel et al (2014)	Australia	Australasia	High	Community	All adults	50	1854	Self-report	8	585	0.32	Moderate
¹⁷⁹ Ruel et al (2014)	China	Asia	Lor or middle	Community	All adults	49	1020	Self-report	11	346	0.34	Moderate
¹⁸⁰ Ryan et al (2018)	Ireland	Europe	High	Community	Middle aged and older	Not reported	4823	Self-report	16	2588	0.54	Moderate
¹⁸¹ Schmidt et al (2016)	Austria, Belgium, Denmark, France, Germany, Italy, the Netherlands, Spain, Sweden, and Switzerland	Europe	High	Community	Only older people	Not reported	56609	Self-report	11	13794	0.24	Moderate
¹⁸² Schottker et al (2016)	Germany	Europe	High	Primary care	Middle aged and older	Median age:70	2547	Medical records and administrative database	14	251	0.10	Moderate
¹⁸³ Seo et al (2017)	South Korea	Asia	High	Community	Middle aged and older	Not reported	156747	Self-report	15	42006	0.27	Moderate
¹⁸⁴ She et al (2019)	China	Asia	Low or middle	Hospitals	Only older people	68.9	1497	Self-report	22	1255	0.84	Moderate
¹⁸⁵ Singh et al (2019)	India	Asia	Low or middle	Community	All adults	41	16287	Self-report	5	1531	0.09	Moderate
¹⁸⁶ Stepanova et al (2015)	USA	North America	High	Community	All adults	34.7	26225	Self-report	13	9992	0.38	High
¹⁸⁷ Stickley et al (2020)	USA	North America	High	Community	All adults	44.9	15311	Self-report	9	3996	0.26	High

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias
¹⁸⁸ Streit et al (2014)	Switzerland	Europe	High	Primary care	Middle aged and older	63.5	1002	Medical records and administrative database	17	676	0.67	Moderate
¹⁸⁹ Stubbs et al (2018)	China, Ghana, India, Mexico, Russia, South Africa	Multiple continents	Low or middle	Community	Middle aged and older	62.4	34129	Self-report	13	19317	0.57	Moderate
¹⁹⁰ Su et al (2016)	China	Asia	Low or middle	Community	Only older people	Not reported	2058	Self-report	10	1012	0.49	Moderate
¹⁹¹ Sundstrup et al (2017)	USA	North America	High	Community	All adults	43.5	10427	Self-report	8	2489	0.24	High
¹⁹² Takahashi et al (2016)	USA	North America	High	Hospitals	All adults	57	6402	Medical records and administrative database	Not reported	3140	0.49	High
¹⁹³ Tinetti et al (2011)	USA	North America	High	Community	Only older people	72.6	5298	Self-report	5	1200	0.23	High
¹⁹⁴ Troelstra et al (2020)	The Netherlands	Europe	High	Community	All adults	Not reported	604	Self-report	26	321	0.53	High
¹⁹⁵ van Zon et al (2020)	USA	North America	High	Community	Middle aged and older	53.8	10719	Self-report	8	2390	0.22	Moderate
¹⁹⁶ Vancampfort et al (2017)	China, Ghana, India, Mexico, Russia, and South Africa	Multiple continents	Low or middle	Community	All adults	Median age: 62	32585	Self-report	11	14524	0.45	Moderate
¹⁹⁷ Vassilaki et al (2015)	USA	North America	High	Primary care	Only older people	78.5	2176	Medical records and administrative database	17	1884	0.87	Moderate
¹⁹⁸ Vassilaki et al (2016)	USA	North America	High	Primary care	Only older people	79	1449	Medical records and administrative database	17	1237	0.85	Moderate
¹⁹⁹ Villarreal et al (2015)	Panama	South America	High	Primary care	Only older people	78.2	304	Self-report	7	227	0.75	Moderate
²⁰⁰ Violan et al (2019)	Spain	Europe	High	Primary care	Only older people	75.4	916619	Medical records and	60	853085	0.93	Moderate

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias
								administrative database				
²⁰¹ von Strauss et al (2000)	Sweden	Europe	High	Community	Only older people	Not reported	502	Self-report	15	155	0.31	Moderate
²⁰² Vos et al (2013)	The Netherlands	Europe	High	Community	Only older people	71.9	315	Self-report	21	202	0.64	Moderate
²⁰³ Vu et al (2019)	Vietnam	Asia	Low or middle	Hospitals	Only older people	71.9	405	Medical records and administrative database	Not reported	146	0.36	High
²⁰⁴ Wang et al (2018)	USA	North America	High	Community	All adults	47	3086	Self-report	20	1109	0.36	Moderate
²⁰⁵ Wang et al (2017)	China	Asia	Low or middle	Community	Only older people	69.24	2705	Self-report	17	1230	0.45	Moderate
²⁰⁶ Wijers et al (2019)	Spain	Europe	High	Community	Middle aged and older	74.2	707	Self-report	21	491	0.69	Moderate
²⁰⁷ Williams et al (2016)	USA	North America	High	Community	All adults	Not reported	23789	Self-report	9	9213	0.39	Moderate
²⁰⁸ Woldesemayat et al (2018)	Ethiopia	Africa	Low or middle	Primary care	All adults	Not reported	411	Self-report	18	73	0.18	Moderate
²⁰⁹ Yao et al (2020)	China	Asia	Low or middle	Community	Middle aged and older	57.7	10084	Self-report	15	3243	0.32	Moderate
²¹⁰ Yorke et al (2017)	USA	North America	High	Community	Middle aged and older	66.6	5877	Self-report	7	3391	0.58	Moderate
²¹¹ You et al (2019)	China	Asia	Low or middle	Community	Only older people	72	5296	Self-report	27	2201	0.42	Moderate
²¹² Zhang et al (2020)	China	Asia	Low or middle	Community	Only older people	74.14	4348	Self-report	15	2338	0.54	Moderate
²¹³ Khanam et al (2011)	Bangladesh	Asia	Low or middle	Community	Only older people	69.5	452	Medical records and administrative database	9	243	0.54	Moderate

Author	Country	Continent	Country income	Setting	Study population	Mean age	No of participants	Source	No of conditions measured	No of MM cases	Proportion with MM	Risk of bias
²¹⁴ Cornell et al (2009)	USA	North America	High	Primary care	All adults	62.4	1645314	Medical records and administrative database	45	1327382	0.81	Moderate
²¹⁵ Cassell et al (2018)	UK	Europe	High	Primary care	All adults	Not reported	403985	Medical records and administrative database	36	109884	0.27	Moderate
²¹⁶ Wong et al (2019)	Hong Kong (SAR of China)	Asia	High	Community	All adults	45.67	1014	Self-report	5	124	0.12	Moderate
²¹⁷ Puth et al (2017)	Germany	Europe	High	Community	All adults	Not reported	19294	Self-report	17	7640	0.40	Moderate

MM: Multimorbidity. No of participants is the total number of participants in the denominator for estimating prevalence in a study (which could be a subset in some included studies)

Table S5: Associations between predictors

	Mean age (lm) Unadjusted coefficient estimates	No of conditions (nb) Unadjusted incident rate ratio			
Mean age		1.0 (1.0-1.0)			
Source					
Self-report	59.7 (57.1-62.3) (intercept)	Ref			
Database	7.0 (1.5-12.5)*	1.8 (1.5-2.2)***			
Continent					
Europe	66.8 (62.8-70.9) (intercept)	Ref			
North America	-7.0 (-12.8 to -1.1)*	0.6 (0.5-0.8)***			
Australasia	-8.0 (-17.5-1.6)	0.8 (0.6-1.1)			
Asia	-8.4 (-14.6 to -2.2)**	0.6 (0.5-0.8)***			
South America	-8.5 (-18.0-1.1)	0.6 (0.4-0.9)**			
Africa	-32.8 (-57.8 to -8.0)**	0.4 (0.2-0.8)*			
Multiple continents	-7.6 (-18.3-3.2)	0.5 (0.3-0.7)***			
Setting					
Community	59.8 (57.2-62.5) (intercept)	Ref			
Primary care	3.5 (-2.5-9.6)	1.7 (1.4-2.1)***			
Hospitals	10.2 (1.5-19.0)*	1.8 (1.3-2.4)***			
Study population					
All adults	48.3 (46.6-50.0) (intercept)	Ref			
Middle-aged and older	15.4 (12.7-18.0)***	0.9 (0.7-1.1)			
Only older people	26.2 (23.7-28.7)***	1.2 (0.9-1.4)			

*<0.05 **<0.01 ***<0.001

Ref: Reference category. lm: Linear regression. nb: Negative binomial regression

Table S6: Risk of bias assessment of included studies

Author	Selection bias	Study design	Confounders	Blinding	Data collection method	Withdrawals and dropouts	Publication bias	Conflict of interest	Overall rating	Reporting of MM measure and definition
^{25.} Aarts et al (2012)	Moderate	Moderate	Moderate	High	Low	Low	Moderate	Low	Moderate	Yes
^{26.} Aarts et al (2011)	Low	High	Moderate	High	Moderate	Moderate	Moderate	Low	Moderate	No
^{27.} Aarts et al (2011)	Moderate	Moderate	Moderate	High	Low	Low	Moderate	Low	Moderate	Yes
^{28.} Abizanda et al (2014)	Moderate	Moderate	Moderate	High	Moderate	Moderate	Moderate	Low	Moderate	Yes
^{29.} Agborsangaya et al (2012)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{30.} Agborsangaya et al (2013)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{31.} Agborsangaya et al (2014)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{32.} Ahrenfeldt et al (2019)	Low	Moderate	Moderate	High	Low	High	Moderate	Low	Moderate	No
^{33.} Alimohammadian et al (2017)	Low	Moderate	Moderate	High	Low	High	Moderate	Low	Moderate	Yes
^{34.} Angst et al (2002)	Moderate	Moderate	Moderate	High	Low	High	High	Unclear	High	No
^{35.} Appa et al (2014)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{36.} Adams et al (2017)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{37.} Ahmadi et al (2016)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{38.} Amaral et al (2018)	High	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes

Author	Selection bias	Study design	Confounders	Blinding	Data collection method	Withdrawals and dropouts	Publication bias	Conflict of interest	Overall rating	Reporting of MM measure and definition
^{39.} An et al (2016)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{40.} Araujo et al (2018)	Low	Moderate	Low	High	Moderate	High	Moderate	Low	Moderate	Yes
^{41.} Arnold-Reed et al (2018)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{42.} Arokiasamy et al (2015)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{43.} Sinnige et al (2015)	Low	Moderate	Moderate	High	Low	Low	Moderate	Low	Moderate	Yes
^{44.} Zemedikun et al (2018)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{45.} Wensing et al (2001)	Moderate	Moderate	Moderate	High	Moderate	Moderate	Moderate	Unclear	Moderate	Yes
^{46.} Mounce et al (2018)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{47.} Taylor et al (2010)	Low	Moderate	Moderate	High	Low	Low	Moderate	Low	Low	Yes
^{48.} Vancampfort et al (2019)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{49.} Vancampfort et al (2018)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{50.} Aubert et al (2016)	Moderate	Moderate	Moderate	High	Moderate	Low	Moderate	Low	Moderate	Yes
^{51.} Autenrieth et al (2013)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{52.} Bahler et al (2015)	Low	Moderate	Moderate	High	Moderate	Low	Moderate	Low	Moderate	Yes
^{53.} Vancampfort et al (2017)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes

Author	Selection bias	Study design	Confounders	Blinding	Data collection method	Withdrawals and dropouts	Publication bias	Conflict of interest	Overall rating	Reporting of MM measure and definition
^{54.} Banjare et al (2014)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{55.} Barra et al (2015)	Moderate	Moderate	High	High	Moderate	High	Moderate	Unclear	Moderate	No
^{56.} Bernard et al (2016)	High	Moderate	High	High	Moderate	Low	Moderate	Low	High	No
^{57.} Biswas et al (2019)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	No
^{58.} Blakemore et al (2016)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	No
^{59.} Blyth et al (2008)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{60.} Bowling et al (2019)	Low	Moderate	Moderate	High	Moderate	Low	Moderate	Low	Moderate	Yes
^{61.} Britt et al (2008)	Low	Moderate	High	High	Moderate	Moderate	Moderate	Low	Moderate	Yes
^{62.} Broeiro-Goncalves (2019)	Low	Moderate	High	High	Moderate	Moderate	Moderate	Low	Moderate	Yes
^{63.} Bruce et al (2010)	High	Moderate	Moderate	High	Low	High	Moderate	Unclear	High	No
^{64.} Burgers et al (2010)	Low	Moderate	High	High	Moderate	Low	Moderate	Low	Moderate	Yes
^{65.} Burke et al (2017)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{66.} Buurman et al (2016)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{67.} Calderon-Larranaga et al (2017)	Moderate	Moderate	Moderate	High	Low	Low	Moderate	Low	Moderate	Yes
^{68.} Camargo-Casas et al (2018)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes

Author	Selection bias	Study design	Confounders	Blinding	Data collection method	Withdrawals and dropouts	Publication bias	Conflict of interest	Overall rating	Reporting of MM measure and definition
^{69.} Canevelli et al (2019)	High	High	High	High	Moderate	High	Moderate	Low	High	Yes
^{70.} Chamberlain et al (2020)	Low	Moderate	Moderate	High	Low	Low	Moderate	Low	Low	Yes
^{71.} Chen et al (2018)	Low	Moderate	High	High	Low	Low	Moderate	Low	Low	Yes
^{72.} Chen et al (2018)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{73.} Cheung et al (2013)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{74.} Chu et al (2018)	Low	Moderate	Moderate	High	Moderate	Moderate	Moderate	Low	Moderate	Yes
^{75.} Chudasama et al (2019)	Moderate	Moderate	Low	High	Moderate	Low	Moderate	Low	Moderate	Yes
^{76.} Cimarras-Otal et al (2014)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{77.} Chin et al (2016)	Moderate	Moderate	Moderate	High	Moderate	Low	Moderate	Low	Moderate	Yes
^{78.} Agrawal et al (2016)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{79.} Gu et al (2018)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{80.} Gunn et al (2012)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{81.} Han et al (2013)	High	High	Moderate	High	Moderate	High	Moderate	Unclear	High	No
^{82.} Hanlon et al (2018)	Low	Moderate	Moderate	High	Low	Low	Moderate	Low	Low	Yes
^{83.} Jantsch et al (2018)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes

Author	Selection bias	Study design	Confounders	Blinding	Data collection method	Withdrawals and dropouts	Publication bias	Conflict of interest	Overall rating	Reporting of MM measure and definition
^{84.} John et al (2003)	Moderate	High	Moderate	High	Low	High	Moderate	Low	High	No
^{85.} Johnson-Lawrence et al (2017)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{86.} Johnston et al (2019)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	No
^{87.} Jones et al (2016)	Low	Moderate	Moderate	High	Low	Low	Moderate	Unclear	Moderate	Yes
^{88.} Jovic et al (2016)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{89.} Juul-Larsen et al (2020)	High	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{90.} Hudon et al (2008)	Low	Moderate	Moderate	High	Low	Low	Moderate	Low	Low	Yes
^{91.} Hussain et al (2015)	Moderate	Moderate	High	High	Moderate	Low	Moderate	Low	Moderate	Yes
^{92.} Ie et al (2017)	High	High	Moderate	High	Moderate	Low	Moderate	Low	High	Yes
^{93.} Ishizaki et al (2019)	Moderate	Moderate	Low	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{94.} Danon-Hersch et al (2012)	Moderate	High	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{95.} de Heer et al (2013)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{96.} Demirchyan et al (2013)	High	Moderate	Low	High	Moderate	High	Moderate	Low	High	No
^{97.} Fabbri et al (2015)	Moderate	Moderate	High	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{98.} Fillenbaum et al (2000)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes

Author	Selection bias	Study design	Confounders	Blinding	Data collection method	Withdrawals and dropouts	Publication bias	Conflict of interest	Overall rating	Reporting of MM measure and definition
^{99.} Kaneko et al (2019)	Moderate	Moderate	High	High	Moderate	Low	Moderate	Low	Moderate	No
^{100.} Kang et al (2017)	High	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{101.} Gandhi et al (2020)	Moderate	Moderate	Moderate	High	High	High	Moderate	Low	Moderate	Yes
^{102.} Costa et al (2018)	High	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{103.} Rizzuto et al (2017)	High	Moderate	High	High	Moderate	Moderate	Moderate	Low	Moderate	Yes
^{104.} Dhalwani et al (2017)	Moderate	Moderate	Low	High	Moderate	High	Moderate	Low	Moderate	Yes
^{105.} Elixhauser et al (1998)	Low	Moderate	High	High	Low	Low	Moderate	Unclear	Low	Yes
^{106.} Fabbri et al (2015)	High	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{107.} Fortin et al (2014)	Low	Moderate	Low	High	Moderate	Moderate	Moderate	Low	Moderate	Yes
^{108.} Fuchs et al (1998)	Moderate	Moderate	Moderate	High	Moderate	Low	Moderate	Unclear	Moderate	No
^{109.} Galenkamp et al (2011)	Low	Moderate	High	High	Moderate	High	Moderate	Unclear	High	No
^{110.} Galenkamp et al (2016)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{111.} Gamma et al (2001)	High	Moderate	High	High	Moderate	High	Moderate	Unclear	High	No
^{112.} Ge et al (2018)	Moderate	Moderate	High	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{113.} Ge et al (2019)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes

Author	Selection bias	Study design	Confounders	Blinding	Data collection method	Withdrawals and dropouts	Publication bias	Conflict of interest	Overall rating	Reporting of MM measure and definition
^{114.} Gould et al (2016)	Moderate	Moderate	Moderate	High	High	High	Moderate	Unclear	Moderate	Yes
^{115.} Habib et al (2014)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	No
^{116.} Harrison et al (2017)	Low	Moderate	High	High	Moderate	Low	Moderate	Unclear	Moderate	No
^{117.} Hayek et al (2017)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{118.} Henninger et al (2012)	Moderate	Moderate	High	High	Moderate	High	Moderate	Unclear	Moderate	No
^{119.} Hernandez et al (2019)	Moderate	Moderate	Moderate	High	High	High	Moderate	Unclear	Moderate	Yes
^{120.} Ho et al (2014)	Moderate	Moderate	High	High	Low	Low	Moderate	Low	Moderate	Yes
^{121.} Khan et al (2019)	Low	Moderate	Low	High	Low	High	Moderate	Low	Low	Yes
^{122.} Kiliari et al (2013)	High	Moderate	High	High	Moderate	Moderate	Moderate	Low	Moderate	No
^{123.} King et al (2018)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{124.} Kingston et al (2018)	Low	Moderate	High	High	Moderate	High	Moderate	Unclear	High	Yes
^{125.} Koyanagi et al (2018)	Low	Moderate	Moderate	High	Moderate	Low	High	Low	Moderate	Yes
^{126.} Kriegsman et al (2004)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{127.} Kristensen et al (2019)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{128.} Kristensen et al (2019)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes

Author	Selection bias	Study design	Confounders	Blinding	Data collection method	Withdrawals and dropouts	Publication bias	Conflict of interest	Overall rating	Reporting of MM measure and definition
^{129.} Kunna et al (2017)	Low	Moderate	Low	High	Moderate	Low	High	Low	Low	Yes
^{130.} Kuwornu et al (2014)	Moderate	Moderate	High	High	Moderate	Low	Moderate	Low	Moderate	Yes
^{131.} Lai et al (2019)	Low	Moderate	High	High	Moderate	Low	Moderate	Low	Moderate	Yes
^{132.} Lai et al (2018)	High	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{133.} Laires et al (2019)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{134.} Lang et al (2015)	Moderate	Moderate	Moderate	High	Moderate	Moderate	Moderate	Low	Moderate	Yes
^{135.} Le Cossec et al (2016)	Low	Moderate	Moderate	High	Moderate	Moderate	Moderate	Low	Moderate	Yes
^{136.} Lee et al (2007)	Low	Moderate	High	High	Low	Low	Moderate	Low	Low	Yes
^{137.} Lee et al (2018)	Low	Moderate	High	High	High	Low	Moderate	Unclear	High	No
^{138.} Li et al (2016)	Low	Low	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{139.} Li et al (2019)	Low	Moderate	Low	High	Moderate	Moderate	Moderate	Low	Moderate	No
^{140.} Lujic et al (2017)	Low	Moderate	Moderate	High	Low	High	Moderate	Low	Moderate	Yes
^{141.} LupianezUnclearVillanueva et al (2018)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{142.} Zhou et al (2018)	Moderate	Moderate	Moderate	High	Moderate	Low	High	Low	Moderate	Yes
^{143.} Zhang et al (2019)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes

Author	Selection bias	Study design	Confounders	Blinding	Data collection method	Withdrawals and dropouts	Publication bias	Conflict of interest	Overall rating	Reporting of MM measure and definition
^{144.} Wong et al (2010)	High	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{145.} Weimann et al (2016)	Low	Moderate	Low	High	Moderate	High	Moderate	Low	Moderate	Yes
^{146.} Wang et al (2017)	Low	Moderate	Low	High	Moderate	High	Moderate	Low	Moderate	Yes
^{147.} Wang et al (2019)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	No
^{148.} Wade et al (2019)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{149.} Maciejewski et al (2019)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{150.} Marengoni et al (2016)	Moderate	Moderate	High	High	Moderate	High	Moderate	Low	Moderate	Yes
^{151.} Marengoni et al (2009)	Moderate	Moderate	High	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{152.} Marques et al (2018)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{153.} Mavaddat et al (2014)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{154.} McDaid et al (2013)	Low	Moderate	High	High	Moderate	High	Moderate	Low	High	Yes
^{155.} Melis et al (2014)	High	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{156.} Min et al (2007)	High	Moderate	High	High	Moderate	High	Moderate	Unclear	High	Yes
^{157.} Momtaz et al (2010)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{158.} Mondor et al (2018)	Low	Moderate	Low	High	Moderate	High	Moderate	Low	Moderate	Yes

Moderate Moderate rate Moderate rate Moderate	High Low High	High High High	Moderate Moderate	Low High	Moderate	Low	Moderate	No
rate Moderate	High	c	Moderate	High	Madarata			
	C	High			woderate	Low	Moderate	Yes
rate Moderate		1	Moderate	Low	Moderate	Unclear	Moderate	Yes
	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
Moderate	Moderate	High	Low	High	Moderate	Low	Moderate	Yes
rate Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
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Author	Selection bias	Study design	Confounders	Blinding	Data collection method	Withdrawals and dropouts	Publication bias	Conflict of interest	Overall rating	Reporting of MM measure and definition
^{174.} Pfortmueller et al (2013)	Moderate	Moderate	High	High	High	High	Moderate	Unclear	High	No
^{175.} Pressley et al (1999)	Low	Moderate	High	High	Moderate	Low	Moderate	Unclear	Moderate	No
^{176.} Prior et al (2016)	Low	Moderate	Moderate	High	Moderate	Low	Moderate	Unclear	Moderate	Yes
^{177.} Ribeiro et al (2018)	High	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{178.} Ruel et al (2014)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{179.} Ruel et al (2014)	Moderate	Moderate	Moderate	High	Low	High	Moderate	Low	Moderate	Yes
^{180.} Ryan et al (2018)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{181.} Schmidt et al (2016)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{182.} Schottker et al (2016)	Moderate	Moderate	Moderate	High	Moderate	Moderate	Moderate	Low	Moderate	Yes
^{183.} Seo et al (2017)	Low	Moderate	Moderate	High	Low	High	Moderate	Low	Moderate	No
^{184.} She et al (2019)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{185.} Singh et al (2019)	Low	Moderate	Moderate	High	Low	Low	Moderate	Unclear	Moderate	Yes
^{186.} Stepanova et al (2015)	Low	High	High	High	High	High	High	Unclear	High	Yes
^{187.} Stickley et al (2020)	Low	Moderate	High	High	Moderate	High	Moderate	Low	High	Yes
^{188.} Streit et al (2014)	Moderate	Moderate	Moderate	High	High	High	Moderate	Unclear	Moderate	Yes

Author	Selection bias	Study design	Confounders	Blinding	Data collection method	Withdrawals and dropouts	Publication bias	Conflict of interest	Overall rating	Reporting of MM measure and definition
^{189.} Stubbs et al (2018)	Low	Moderate	High	High	Moderate	Low	Moderate	Low	Moderate	Yes
^{190.} Su et al (2016)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{191.} Sundstrup et al (2017)	Low	Moderate	High	High	Moderate	High	Moderate	Unclear	High	Yes
^{192.} Takahashi et al (2016)	Moderate	Moderate	High	High	High	Low	Moderate	Low	High	No
^{193.} Tinetti et al (2011)	Low	Moderate	High	High	High	High	Moderate	Unclear	High	No
^{194.} Troelstra et al (2020)	High	Moderate	High	High	Moderate	Low	Moderate	Unclear	High	Yes
^{195.} van Zon et al (2020)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{196.} Vancampfort et al (2017)	Low	Moderate	Moderate	High	Moderate	Low	Moderate	Low	Moderate	Yes
^{197.} Vassilaki et al (2015)	Low	Moderate	Moderate	High	Moderate	Low	Moderate	Unclear	Moderate	Yes
^{198.} Vassilaki et al (2016)	Low	Moderate	Moderate	High	Moderate	Low	Moderate	Unclear	Moderate	Yes
^{199.} Villarreal et al (2015)	High	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	No
^{200.} Violan et al (2019)	Low	Moderate	Moderate	High	High	Low	Moderate	Low	Moderate	Yes
^{201.} von Strauss et al (2000)	Moderate	Moderate	Moderate	High	Moderate	Low	Moderate	Low	Moderate	No
^{202.} Vos et al (2013)	Moderate	Moderate	High	High	Moderate	High	Moderate	Low	Moderate	No
^{203.} Vu et al (2019)	High	Moderate	High	High	Moderate	High	Moderate	Low	High	No

Author	Selection bias	Study design	Confounders	Blinding	Data collection method	Withdrawals and dropouts	Publication bias	Conflict of interest	Overall rating	Reporting of MM measure and definition
^{204.} Wang et al (2018)	Moderate	Moderate	Low	High	Moderate	Low	Moderate	Unclear	Moderate	Yes
^{205.} Wang et al (2017)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{206.} Wijers et al (2019)	Low	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	No
^{207.} Williams et al (2016)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	No
^{208.} Woldesemayat et al (2018)	High	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{209.} Yao et al (2020)	Moderate	Moderate	Low	High	Moderate	High	Moderate	Low	Moderate	Yes
^{210.} Yorke et al (2017)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{211.} You et al (2019)	Moderate	Moderate	Moderate	High	Moderate	High	Moderate	Low	Moderate	Yes
^{212.} Zhang et al (2020)	Moderate	Moderate	Low	High	Moderate	Low	Moderate	Low	Moderate	Yes
^{213.} Khanam et al (2011)	Moderate	Moderate	Moderate	High	Moderate	Low	Moderate	Unclear	Moderate	Yes
^{214.} Cornell et al (2009)	Low	Moderate	High	High	Moderate	Low	Moderate	Unclear	Moderate	Yes
^{215.} Cassell et al (2018)	Low	Moderate	High	High	Moderate	Moderate	Moderate	Low	Moderate	No
^{216.} Wong et al (2019)	High	Moderate	Moderate	High	Moderate	High	Moderate	Unclear	Moderate	Yes
^{217.} Puth et al (2017)	Moderate	Moderate	Moderate	High	Moderate	Moderate	Moderate	Low	Moderate	Yes

Table S7: Output of adjusted meta-analytic model based on 217 studies

	Pooled prevalence of multimorbidity of each subgroup (%, 95% CI)	Meta-regression Unadjusted Odds Ratio (95% CI)	Meta-regression Adjusted Odds Ratio (95% CI) R ² 42.4%	FMI
Group of mean age		R ² 27.0%		
<59	30.4 (27.0-33.9)	Ref	Ref	Ref
59-73	43.5 (38.0-49.1)	1.8 (1.3-2.3)***	2.0 (1.6-2.6)***	0.3
≥74	67.8 (61.3-73.7)	6.4 (4.6-8.9)***	4.7 (3.4-6.5)***	0.2
No of conditions		R ² 6.9%		
<9	29.9 (24.9-35.4)	Ref	Ref	Ref
9-19	43.5 (39.1-47.9)	1.8 (1.3-2.5)***	1.7 (1.3-2.2)***	0.1
20-43	46.7 (38.4-55.2)	2.1 (1.4-3.1)***	2.2 (1.5-3.3)***	0.2
≥44	54.5 (32.6-74.8)	2.8 (1.5-5.4)**	2.8 (1.6-4.8)***	0.1
Setting		R ² 3.7%		
Community	37.8 (34.4-41.4)	Ref	Ref	Ref
Primary care	51.2 (41.6-60.7)	1.7 (1.2-2.5)**	1.8 (1.2-2.6)**	0.1
Hospital	47.1 (31.9-63.0)	1.5 (0.9-2.4)	0.8 (0.5-1.3)	0.1
Care home	73.9 (72.8-74.9)	4.6 (0.6-36.6)	1.5 (0.3-8.4)	0.04
Source		R ² 2.8%		
Self-report	38.3 (34.4-42.2)	Ref	Ref	Ref
Database	48.9 (42.2-55.6)	1.5 (1.1-2.1)**	0.8 (0.6-1.1)	0.1
Continent		R ² 7.4%		
North America	48.9 (42.1-55.7)	Ref	Ref	Ref
Europe	44.0 (37.7-50.4)	0.8 (0.6-1.2)	0.5 (0.4-0.7)***	0.1
Australasia	28.2 (20.3-37.6)	0.4 (0.2-0.8)**	0.4 (0.2-0.6)***	0.08
Asia	34.3 (28.6-40.5)	0.5 (0.4-0.8)**	0.5 (0.3-0.7)***	0.1
South America	47.5 (31.2-64.4)	0.9 (0.5-1.8)	0.8 (0.5-1.3)	0.1
Africa	13.8 (4.5-35.2)	0.2 (0.06-0.5)***	0.2 (0.1-0.5)***	0.1
Multiple continents	41.4 (31.0-52.6)	0.7 (0.4-1.4)	0.7 (0.4-1.2)	0.1

*<0.05 **<0.01 ***<0.001

Ref: Reference category. FMI: Fraction of missing information.

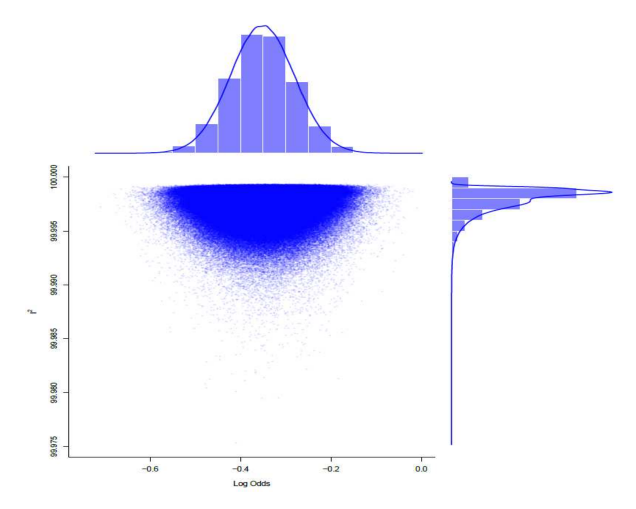
Supplemental material

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Table S8: Definition of variables

Variable name	Definition				
Study setting					
Community	Studies that used population surveys, insurance claims databases, or research databases				
Primary care	Studies that were carried out in primary care settings				
Hospital	Studies that were carried out in hospital settings				
Data source					
Self-report	Studies that collected data using self-report or interviews				
Medical records and administrative	Studies that collected data using electronic medical records, medical chart reviews, insurance claims				
databases	databases, pharmacy databases, or research databases				
Study population					
All adults	Studies with a sample of population aged 18 and older (n=45), aged 20 and older (n=8), aged 21 and				
	older (n=3), aged 25 and older (n=2), or others (n=27) (e.g. aged 16 and older, or aged 17 and older)				
Middle-aged and older	Studies with a sample of population aged 50 and older (n=25), aged 40 and older (n=5), aged 40 and				
	older (n=10), or others (n=6) (e.g. aged 57 and older, or aged 45 and older)				
Only older people	Studies with a sample of population aged 65 and older (n=22), aged 60 and older (n=25), aged 70 and				
	older (n=5) or others (n=11) (e.g. aged 68 and older, aged 77 and older, aged 78 and older, or aged 80				
	and older)				

Figure S1: Graphical display of study effect sizes and heterogeneity



No obvious subgroup effects were identified

Figure S2: Process of examining and identifying outlying studies in meta-analysis

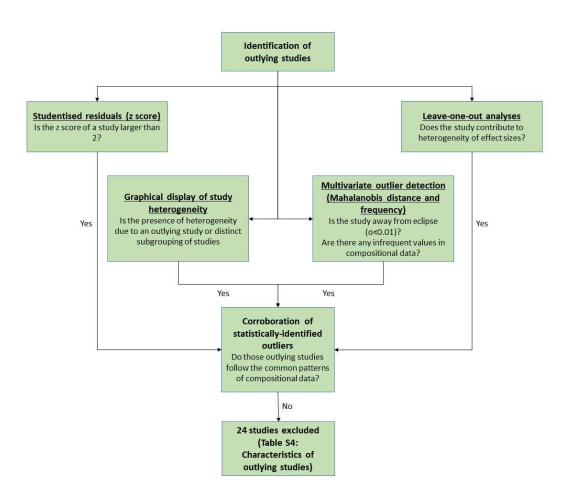
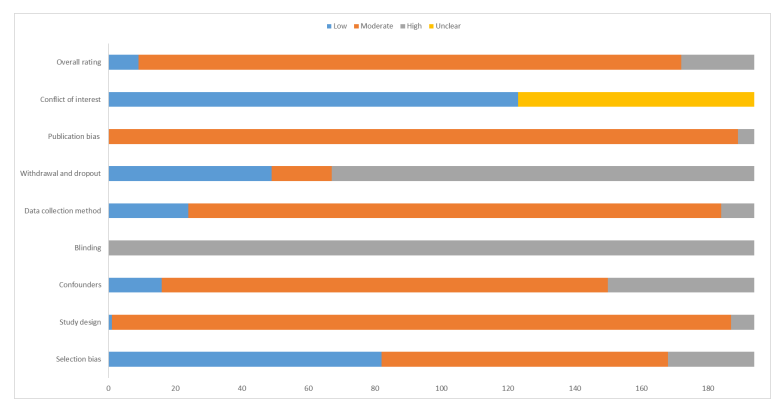


Figure S3: Summary of risk of bias assessment



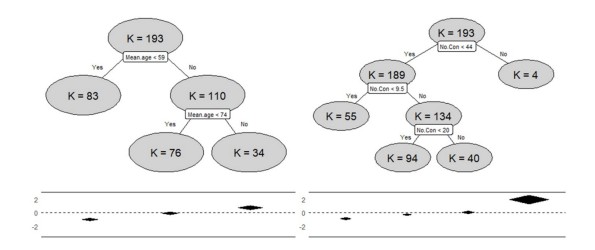


Figure S4: Meta-regression trees for predicting the pooled estimated prevalence of multimorbidity (based on 'mean age' and 'number of conditions' predictors. unit: log(odds))

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