Exploring the risk-factor association between depression and incident stroke: a systematic review and meta-analysis

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Abstract: There is growing evidence that depression increases the risk of incident stroke. However, few studies have considered possible residual confounding effects by preexistent cerebrovascular and cardiac diseases. Therefore, we synthesized data from cohort studies to explore whether depressed individuals free of cerebrovascular and cardiac diseases are at higher risk of incident stroke. We searched the electronic databases PubMed and Medline for eligible cohort studies that examined the prospective association between depression and first-ever stroke. A random-effects model was used for quantitative data synthesis. Sensitivity analyses comprised cohort studies that considered a lag period with exclusion of incident strokes in the first years of follow-up to minimize residual confounding by preexistent silent strokes and excluded cardiac disease at baseline. Overall, we identified 28 cohort studies with 681,139 participants and 13,436 (1.97%) incident stroke cases. The pooled risk estimate revealed an increased risk of incident stroke for depression (relative risk 1.40, 95% confidence interval [CI] 1.27–1.53; \( P<0.0001 \)). When we excluded incident strokes that occurred in the first years of follow-up, the prospective association between depression and incident stroke remained significant (relative risk 1.64, 95% CI 1.27–2.11; \( P<0.0001 \)). This positive association also remained after we considered only studies with individuals with cardiac disease at baseline excluded (relative risk 1.43, 95% CI 1.19–1.72; \( P<0.0001 \)). The prospective association of depression and increased risk of first-ever stroke demonstrated in this meta-analysis appears to be driven neither by preexistence of clinically apparent cerebrovascular and cardiovascular diseases nor by silent stroke.

Keywords: stroke, prestroke, depression, risk factor

Introduction

While for many years, supported by the results of two recent high-quality meta-analyses, it has been assumed that depression constitutes a possible risk factor for stroke, the previous hypothetic concept of “vascular depression” is fueling a new debate in the scientific community.¹³ The theory is that the presence of subclinical cerebrovascular diseases may induce neurobiological changes in brain regions, including the dorsal cortex and limbic structures, and therewith predispose, precipitate, or perpetuate depressive syndromes and at the same time occurrence of stroke.³ The findings of a recent large-scale cohort study including more than 10,000 people further substantiate this hypothesis, when no association between depression and incident stroke was evident after considering only cerebrovascular events that occurred beyond the first years of follow-up.⁴ The latter approach may ensure that results are not biased by the fact that people who have had a clinically silent or apparent stroke immediately prior to study initiation have a higher risk of developing depression in the short term (ie, reverse causality).³
To date, a large portion of all conducted cohort studies have not focused on stroke-free participants as the target population. For instance, the meta-analysis by Pan et al identified seven of 28 included studies that enrolled people with a history of stroke, increasing the chance that the findings were biased by reverse causality. Less attention has been given to other vascular comorbidities like cardiac disease in this matter. Apart from the probable causal relationship with cardiac disease causing depression and vice versa, it is a well-known fact that patients with cardiac disease have more than twice the odds of having a stroke than those without cardiac disease. Moreover, these patients generally have a higher burden of vascular risk factors potentiating the manifestation of subclinical cerebrovascular disease that in turn may increase the possibility of residual confounding in longitudinal cohort studies.

Since recently published reviews and meta-analyses, results of other longitudinal cohort studies investigating the prospective association between depression and incident stroke have been published. Modified inclusion criteria derived from conclusions and recommendations of previous research have been implemented to minimize study heterogeneity and biases, and now allow more in-depth conclusions on the potential risk-factor association between depression and stroke. Therefore, we aimed to update and enlarge the earlier knowledge base on the risk of a first-ever stroke in people with depression compared to people without depression by performing a systematic review and meta-analysis of cohort studies. We particularly explored the question of whether people with depression and free of both cerebrovascular and cardiac diseases are at higher risk of incident stroke or whether such association might rather result from reverse causality.

Materials and methods

Literature search and study selection

We performed a systematic literature search using the PubMed and Medline electronic databases. The text words “depression” and “stroke” with the use of the Boolean operator “AND” were used to identify relevant studies that examined the prospective association between depression and incident stroke. We exclusively chose these two text words and their corresponding MeSH (Medical Subject Heading) terms (“depressive disorder” OR “depressive” AND “disorder” OR “depressive disorder” OR “depression” OR “depression AND stroke”) to ensure coverage of all potentially eligible articles stored in the searched electronic databases. In addition, we manually searched the bibliographies of recently published opinion statements, reviews, and meta-analyses that addressed the risk-factor association between depression and stroke to identify further articles of interest. We only considered articles published in the English language for further review and excluded duplicates.

The title and abstract of the identified articles were screened according to the eligibility criteria detailed in the next section by an independent reviewer (KB). Full texts were retrieved and reviewed for definite eligibility. Any ambiguities concerning the eligibility of a full text were discussed with a second independent investigator (JK) to reach a final consensus. In the case of missing information or any obscurities, the lead authors of the identified articles were contacted for clarification. The literature search covered studies from the earliest date available until our last search on August 24, 2014. Our systematic review and meta-analysis complied with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines.

Eligibility criteria

According to our formulated PICOS (participants, interventions, comparators, outcomes, and study design) question, an original epidemiological study was included if 1) participants were nonhospitalized adults with no history of stroke or transient ischemic attack (TIA) at the time of study initiation, 2) the exposure (intervention) variable was defined as depression, whereas assessment of depression had to be prospectively performed at baseline and based upon an objective measure like a neuropsychological mood scale or neuropsychiatric evaluation that complied with the Diagnostic Statistical Manual for Mental Disorders (DSM)-III/IV or the International Classification of Diseases (ICD)-7–10, 3) the comparison group consisted of participants without depression at the time of study initiation, 4) the dichotomous outcome event of interest was a first-ever stroke during the follow-up period, including fatal and nonfatal ischemic stroke, TIA, and intracerebral hemorrhage (hereinafter referred as “all stroke”); and 5) the study design was a community-, population-, or registry-based longitudinal cohort study reporting relative effect estimates, such as hazard ratios (HRs), relative risk (RRs), and odds ratios (ORs).

Data items

The following data items were extracted from the included original full-text articles and entered into a predetermined Excel (Microsoft, Redmond, WA, USA) spreadsheet: 1) article information (name of first author, publication year, journal), 2) study design (population-level, study geographical location, length of follow-up, methods used for assessment of the exposure [eg, applied mood scale, neuropsychiatric...
Assessment of risk of bias in individual studies

Each individual cohort study was reviewed for study quality and potential sources of bias by two independent investigators (KB and JK) using the Newcastle–Ottawa Scale for observational studies recommended by the Cochrane Collaboration. A Briefly, each included study was rated according to three grouping items: selection of cohort, comparability of cohort, and assessment of outcome. High-quality choices were scored with a star, with a maximum of nine stars achievable for all eight subitems (one star for each numbered item within the selection and outcome categories; a maximum of two stars for comparability). Of note, adjustment for the most relevant stroke risk factors (eg, arterial hypertension and age), minimum length of follow-up of 10 years, dropout rate less than 20%, and exposure, as well as outcome ascertained other than self-report (eg, linked primary record sources), were considered high-quality choices.

Synthesis of results and additional analyses

Statistical analyses were performed with the use of the software package Stata (Version 12.1; StataCorp, College Station, TX, USA) and the user-written package Metan for meta-analyses. For simplicity reasons, we reported RR with 95% CI as the common effect estimate for the association between the exposure and the incident cases (hereinafter referred to as overall RR). However, to minimize statistical heterogeneity, the primary and some of the additional analyses were carried out separately for studies reporting HR and OR/RR. The OR and RR reflect the risk at a given time point, and both can be considered interchangeable when the absolute number of cases is small (<10%). Since the event rates were below 10% in the included studies, we merged OR and RR for this purpose. The most fully adjusted effect estimates from the included studies were combined using the DerSimonian and Laird random-effects model. Assuming substantial heterogeneity among a set of large-scale cohort studies gathered from the published literature, this method was chosen a priori, as it cannot be anticipated that variation in observed effect estimates is exclusively a result of sampling error (that would be the assumption for a fixed-effect model).

A test of between-study statistical heterogeneity was performed using the Cochran’s Q statistic (with significant heterogeneity set at \( P < 0.1 \)), and the percentage of variation across studies attributable to heterogeneity was described with the \( I^2 \) statistic. An \( I^2 \) value of 0 indicates no, 25%–50% low, 50%–75% moderate, and >75% high observed heterogeneity. We ascertained potential sources of between-study heterogeneity and the robustness of the obtained results by several sensitivity and subgroup analyses. Statistical significance was set at \( P < 0.05 \). If an included study reported stratified effect estimates, these were pooled using the inverse-variance fixed-effect or DerSimonian and Laird random-effects model (whichever applicable). We assessed publication bias by visual inspection of the funnel plot and the use of Egger’s linear regression and Begg’s rank correlation tests. The Duval and Tweedie trim-and-fill method was applied to identify and adjust for significant small-study effects.

Results

Systematic review

As detailed in the flowchart in Figure 1, a total of 6,090 titles/abstracts were retrieved from the electronic literature and 62 from the bibliographies of published literature. Of these, we independently reviewed 61 full-text articles for eligibility and retained 28 articles for our quantitative data synthesis. Since the previously published meta-analyses covering a literature search until May 2011, nine more published studies investigating the prospective relationship between depression and stroke with 412,699 participants and 6,559 incident stroke cases were identified according to our eligibility criteria and included in our quantitative data synthesis. All included studies were designed as prospective community- or population-based cohort studies, except two that comprised registry-based cohorts. The total number of participants in the included studies was 681,139 with 13,436 (1.97%) incident stroke cases observed during a total follow-up ranging from 3 to 29 years. One study did not report the exact number of incident cases. Diagnosis of depression was largely based on a self-rating depression scale (n=24), with the Center for Epidemiological Studies – Depression (CES-D) being the most commonly used assessment scale (n=11). In only four studies, the baseline depression diagnosis originated from a direct evaluation through a psychologist, psychiatrist, or other physician according to ICD or DSM criteria.

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Most of the studies used medical records and death certificates, to some extent in combination with self- or proxy reporting, for the diagnosis of stroke. In only four studies, the diagnosis of (nonfatal) stroke solely relied on self- or proxy reporting.23,27,30,33 Most studies restricted the end point to ischemic and hemorrhagic stroke (n = 15),4,23,24,26,28–31,36,37,40,41,44,47,48 two studies to ischemic stroke and TIA,22,42 and eleven studies captured all stroke (ischemic, hemorrhagic, and TIA) end points.25,27,32–35,38,39,43,45,46 With regard to stroke survival, three studies recorded only fatal stroke.24,28,32 Fifteen studies excluded participants with cardiac disease at baseline a priori or in additional analyses.25,26,28,31,32,34,36–38,41,42,44,46–48 of which six had not been included in previous meta-analyses. In terms of effect estimates, six studies reported RRs,31,33,35,38,39,47 19 HRs,4,22–28,32,36,37,46–48 and three ORs.29,30,34 A full summary of the included studies, with an assessment of study quality by the Newcastle–Ottawa Scale, is provided in Table 1.

To minimize the possibility of reverse causality, seven studies considered a lag period by excluding the first years of follow-up (first 2 years, n = 2; first 3 years, n = 2; first 5 years, n = 3),4,24,31,32,36,37,41 and one study considered only participants free of cerebrovascular disease who had received a depression diagnosis at least 10 years before baseline.42 Six studies were primarily conducted in participants aged ≥65 years,22,23,32,35,39,41 and seven studies in participants <65 years.4,26,28,30,36,37,44 The age of participants in the remainder of studies overlapped these age strata; however, two studies provided additional analyses for effect estimates stratified by age (<65 and ≥65 years).27,41 Three studies exclusively reported effect estimates stratified by race/sex (white men, white women, black persons) and age (<65 and ≥65 as well as 65–74 and ≥75 years, respectively) strata.23,31,43 From these studies, pooled estimates were used for the current analysis. An additional analysis defining...
Table 1 Characteristics of the included cohort studies

<table>
<thead>
<tr>
<th>Study</th>
<th>Study population* and location</th>
<th>Length of follow-up</th>
<th>Definition of exposure</th>
<th>Definition of outcome</th>
<th>Cases, n</th>
<th>Confounder adjustment</th>
<th>Newcastle–Ottawa Scale</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arbelaez et al²²</td>
<td>5,525 men (42%) and women, ≥65 years; United States</td>
<td>11 years (median), 1989–2000</td>
<td>10-item CES-D ≥8</td>
<td>Fatal/nonfatal ischemic stroke and TIA; medical records and death certificates</td>
<td>611</td>
<td>Age, sex, race, occupation, income, education, marital status, hypertension, diabetes, smoking, history of cardiac disease, lipids, BMI</td>
<td>⭐⭐⭐⭐</td>
</tr>
<tr>
<td>Avendano et al²³</td>
<td>2,604 men (n/a) and women, ≥65 years; United States</td>
<td>12 years, 1982–1994</td>
<td>20-item CES-D ≥21</td>
<td>Fatal/nonfatal ischemic and hemorrhagic stroke; self-reports (nonfatal) and death certificates</td>
<td>270</td>
<td>Age, sex, race, education, income</td>
<td>⭐⭐⭐</td>
</tr>
<tr>
<td>Brunner et al²⁶</td>
<td>10,036 men (67%) and women, 35–55 years; United Kingdom</td>
<td>24 years, 1985–2009</td>
<td>GHQ-30 ≥5</td>
<td>Fatal/nonfatal ischemic and hemorrhagic stroke; hospital records or general practitioner confirmation, death certificates</td>
<td>168</td>
<td>Age, sex, ethnicity</td>
<td>⭐</td>
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<tr>
<td>Everson et al²⁴</td>
<td>6,676 men (46%) and women, 17–94 years; United States</td>
<td>29 years, 1965–1994</td>
<td>18-item HPL Depression Scale ≥5</td>
<td>Fatal ischemic and hemorrhagic stroke; death certificates</td>
<td>169</td>
<td>Age, sex, race, education, alcohol use, smoking, BMI, hypertension, diabetes</td>
<td>⭐⭐⭐</td>
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<tr>
<td>Everson-Rose et al²⁵</td>
<td>6,749 men (47%) and women, 45–84 years; United States</td>
<td>8.5 years (median), 2000–2012</td>
<td>20-item CES-D ≥16</td>
<td>All fatal/nonfatal stroke; clinical diagnosis</td>
<td>195</td>
<td>Age, race, sex, education, study site, systolic blood pressure, alcohol use, smoking, physical activity, BMI, height, use of antihypertensives, diabetes mellitus/fasting blood glucose, lipids</td>
<td>⭐⭐⭐</td>
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<tr>
<td>Gafarov et al²⁶</td>
<td>560 women aged 25–64 years; Russia</td>
<td>16 years, 1995–2010</td>
<td>15-item MOPSY questionnaire</td>
<td>Fatal/nonfatal ischemic and hemorrhagic stroke; medical records and death certificates</td>
<td>35</td>
<td>Age</td>
<td>⭐⭐⭐</td>
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<tr>
<td>Glymour et al²⁷</td>
<td>19,087 men (41%) and women, ≥50 years; United States</td>
<td>8.1 years (mean), 1996–2006</td>
<td>8-item CES-D ≥3</td>
<td>All fatal/nonfatal stroke; self- or proxy-reporting</td>
<td>1,864</td>
<td>Age, race, education, income, wealth, marital status, overweight, obese, alcohol use, smoking, hypertension, diabetes, history of cardiac disease</td>
<td>⭐⭐⭐⭐</td>
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<tr>
<td>Gump et al²⁸</td>
<td>11,216 men, 35–57 years; United States</td>
<td>18.4 years (median), 1981–1999</td>
<td>20-item CES-D ≥16</td>
<td>Fatal ischemic and hemorrhagic stroke; death certificate</td>
<td>167</td>
<td>Age, intervention group, race, education, smoking, blood pressure, alcohol use, cholesterol, history of cardiac disease</td>
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Table 1 (Continued)

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<thead>
<tr>
<th>Study</th>
<th>Study population* and location</th>
<th>Length of follow-up</th>
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<th>Definition of outcome</th>
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<tr>
<td>Hamano et al10</td>
<td>326,229 men (42%) and women, ≥30 years; Sweden</td>
<td>3 years, 2005–2007</td>
<td>Clinical diagnosis by primary care, inpatient or outpatient registries (depressive disorder according to ICD-10)</td>
<td>Fatal/nonfatal ischemic and hemorrhagic stroke; medical records</td>
<td>4,718</td>
<td>Potential confounders (not specified)</td>
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<td>Jackson et al10</td>
<td>10,547 women, 47–52 years; Australia</td>
<td>12 years, 1998–2010</td>
<td>20-item CES-D ≥10</td>
<td>Fatal/nonfatal ischemic and hemorrhagic stroke; self-reports (nonfatal), death certificates</td>
<td>177</td>
<td>Age, education, homeownership, hypertension, diabetes mellitus, heart disease, hysterectomy/oophorectomy, smoking, alcohol use, physical activity, BMI</td>
<td>★☆☆☆☆☆</td>
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<tr>
<td>Jonas et al11</td>
<td>6,095 men (n/a) and women, 25–74 years; United States</td>
<td>16 years (median), 1971–1992</td>
<td>GWB-D ≥12</td>
<td>Fatal/nonfatal ischemic and hemorrhagic stroke; hospital records and death certificates</td>
<td>483</td>
<td>Age, race, sex, education, smoking status, BMI, alcohol use, nonrecreational physical activity, serum cholesterol level, history of diabetes, history of heart disease, and systolic blood pressure</td>
<td>★☆☆☆ ☆☆★</td>
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<tr>
<td>Kamphuis et al12</td>
<td>799 men, 70–90 years; Finland, Italy, Netherlands</td>
<td>7.4 years (mean), 1989–2000</td>
<td>20-item Zung SDS ≥60</td>
<td>All fatal stroke; death certificates</td>
<td>66</td>
<td>Age, country, education, BMI, smoking, alcohol intake, blood pressure, cholesterol, physical activity</td>
<td>★☆☆☆☆☆</td>
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<tr>
<td>Larson et al13</td>
<td>1,703 men (38%) and women, ≥18 years; United States</td>
<td>13 years, 1981–1993 and 1996</td>
<td>DIS (major depression according to DSM-III)</td>
<td>All fatal/nonfatal stroke; self-reports (nonfatal), death certificates</td>
<td>95</td>
<td>Age, sex, education, diabetes, blood pressure, smoking, history of cardiac disease</td>
<td>★☆☆☆☆☆</td>
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<tr>
<td>Li et al14</td>
<td>5,015 men (37%) and women, ≥18 years; Taiwan</td>
<td>9 years, 2001–2009</td>
<td>Clinical diagnosis by psychiatrist (major depressive disorder according to DSM-III)</td>
<td>All fatal/nonfatal stroke; hospital records</td>
<td>156</td>
<td>Age, sex, diabetes mellitus, hypertension, hyperlipidemia, substance comorbidities</td>
<td>★☆☆☆☆☆</td>
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</table>

*Study population and location details include gender, age range, country, and sample size.
<table>
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<th>Sample Size</th>
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<th>Results</th>
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<tr>
<td>Liebetrau et al.</td>
<td>401 men (30%) and women, 85 years; Sweden</td>
<td>3 years, 1986–1990</td>
<td>Clinical diagnosis by psychiatrist (major depression and dysthymia according DSM-III)</td>
<td>All fatal/nonfatal stroke; hospital discharge register, death certificates, self-reports, and key informants</td>
<td>56</td>
<td>Sex, depression at baseline, and blood pressure</td>
<td>*☆☆☆☆☆</td>
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<tr>
<td>Majed et al.</td>
<td>9,601 men, 50–59 years; Ireland, France</td>
<td>10 years (median), from 1991</td>
<td>Fourth quartile of 13-item modified CES-D compared with first quartile</td>
<td>Fatal/nonfatal ischemic and hemorrhagic stroke; hospital or general practitioner records</td>
<td>136</td>
<td>Age, study centers, socioeconomic factors (marital status, education level, employment status), physical activity, smoking, alcohol intake, systolic blood pressure, use of antihypertensives, BMI, total and high-density lipoprotein cholesterol, treatment for diabetes, and use of antidepressants</td>
<td>*☆☆☆☆☆</td>
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<tr>
<td>Nabi et al.</td>
<td>23,282 men (41%) and women, 20–54 years; Finland</td>
<td>7 years, 1998–2005</td>
<td>21-item BDI ≥10</td>
<td>Fatal/nonfatal ischemic and hemorrhagic stroke; hospital discharge register or mortality records</td>
<td>129</td>
<td>Age, sex, education, alcohol use, sedentary lifestyle, smoking, obesity, hypertension, diabetes, incident cardiac disease</td>
<td>*☆☆☆☆☆</td>
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<tr>
<td>Ohira et al.</td>
<td>879 men (35%) and women, 40–78 years; Japan</td>
<td>10.3 years (mean), 1985–1996</td>
<td>Zung SDS ≥35</td>
<td>All fatal/nonfatal stroke; death certificate, medical records, or clinical diagnosis</td>
<td>69</td>
<td>Age, sex, body mass index, systolic blood pressure levels, serum total cholesterol levels, alcohol intake, cigarette smoking, antihypertensive medication, diabetes mellitus</td>
<td>*☆☆☆☆☆</td>
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<tr>
<td>Ostir et al.</td>
<td>2,478 men (31%) and women, ≥65 years; United States</td>
<td>6 years, 1986–1992</td>
<td>20-item CES-D ≥9</td>
<td>All fatal/nonfatal stroke; clinical diagnosis or death certificates</td>
<td>340</td>
<td>Age, sex, race, marital status, education, BMI, smoking, diabetes, hypertension, history of cardiac disease</td>
<td>*☆☆☆☆☆</td>
</tr>
<tr>
<td>Pan et al.</td>
<td>80,574 women, 54–79 years; United States</td>
<td>6 years, 2000–2006</td>
<td>MHI-5 ≤52</td>
<td>Fatal/nonfatal ischemic and hemorrhagic stroke; self-report, medical records, death certificates</td>
<td>1,033</td>
<td>Age, marital status, parental history of myocardial infarction, ethnicity, physical activity level, body mass index, alcohol consumption, smoking status, menopausal status, postmenopausal hormone therapy, current aspirin use, current multivitamin use, Dietary Approaches to Stop Hypertension dietary score, history of hypertension, hypercholesterolemia, diabetes, cancer, heart diseases</td>
<td>*☆☆☆☆☆</td>
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<tbody>
<tr>
<td>Péquignot et al</td>
<td>7,308 men (37%) and women, ≥65 years; France</td>
<td>5.3 years (median), from 1999</td>
<td>20-item CES-D ≥16</td>
<td>Fatal/nonfatal ischemic and hemorrhagic stroke; hospital records, death certificates</td>
<td>141</td>
<td>Age, study center, sex, smoking status, alcohol consumption, high blood pressure, impaired fasting glycemia or diabetes, hypercholesterolemia, living alone, education level, Mini Mental State Examination score</td>
<td>★★☆☆☆☆</td>
</tr>
<tr>
<td>Rahman et al</td>
<td>36,654 men (44%) and women, ≥18 years; Sweden</td>
<td>3.9 years (mean), 2006–2009</td>
<td>Clinical diagnosis by psychiatrist (any depression according to ICD-7)</td>
<td>Fatal/nonfatal ischemic stroke and TIA; death register, hospital discharge register</td>
<td>833</td>
<td>Birth year, sex, smoking status, educational level, hypertension, diabetes, alcohol intake, BMI</td>
<td>★☆☆☆☆★</td>
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<tr>
<td>Salaycik et al</td>
<td>4,120 men (46%) and women, ≥29 years; United States</td>
<td>8 years, from 1990 and 1996</td>
<td>20-item CES-D ≥16</td>
<td>All fatal/nonfatal stroke; medical records</td>
<td>228</td>
<td>Age, sex, blood pressure, diabetes, atrial fibrillation, history of cardiac disease, left ventricular hypertrophy, smoking</td>
<td>★☆☆☆☆★</td>
</tr>
<tr>
<td>Stürmer et al</td>
<td>3,738 men (n/a) and women, 40–65 years; Germany</td>
<td>8.5 years (median), 1992–2003</td>
<td>Personality scale: high category vs medium category</td>
<td>Fatal/nonfatal ischemic and hemorrhagic stroke; medical records (treating doctors), death certificates</td>
<td>62</td>
<td>Age, sex, BMI, smoking status, alcohol consumption, exercise, comorbidity (cancer, hypertension, hyperlipidaemia, diabetes), family history of stroke, education</td>
<td>★☆☆☆☆★</td>
</tr>
<tr>
<td>Surtees et al</td>
<td>20,627 men (43%) and women, 41–80 years; United Kingdom</td>
<td>8.5 years (median), 1996–2006</td>
<td>HLEQ (major depressive disorder according to DSM-IV)</td>
<td>All fatal/nonfatal stroke; clinical diagnosis or death certificate</td>
<td>595</td>
<td>Age, sex, smoking, blood pressure, cholesterol, obesity, history of cardiac disease, diabetes, social class, education, antihypertensive use, family history of stroke, antidepressant use</td>
<td>★☆☆☆★☆</td>
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<tr>
<td>Vogt et al</td>
<td>2,573 men (46%) and women, ≥18 years; United States</td>
<td>15 years, 1970–1985</td>
<td>Depressive Index, upper tertile vs bottom tertile</td>
<td>All fatal/nonfatal stroke; death index and vital records</td>
<td>–</td>
<td>Age, sex, socioeconomic status, length of health plan membership, subjective health status, smoking</td>
<td>★☆☆☆★☆</td>
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</tbody>
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depression in relation to the use of antidepressant medication was performed in six of 27 studies.30,37,40–43

Primary quantitative data synthesis
Of the 28 included studies, 15 reported a significant positive association between the risk of a first-ever stroke and depressive disorder.2,22,25–27,29–33,35,38,40–42 In ten studies, there was a positive association too, though these did not show statistical significance in the primary analyses.23,24,28,34,36,39,43–45,47 Pooling the risk estimates from all of these 28 studies revealed an increased risk of an incident stroke for depression by 40% (overall RR 1.40, 95% CI 1.27–1.53; P<0.0001), with an I² statistic of 49% (P=0.002 by Cochran Q test). This association remained alike when we separately pooled studies that reported ORs/RRs (OR/RR 1.54, 95% CI 1.25–1.89; P<0.0001; n=9) and HRs (HR 1.34, 95% CI 1.21–1.49; P<0.0001; n=19). There was evidence of moderate heterogeneity with an I² statistic of 67% (P=0.002 by Cochran Q test) for studies that reported ORs/RRs, and low heterogeneity with an I² statistic of 36% (P=0.064 by Cochran Q test) for studies that reported HRs. Figure 2 depicts the forest plots for the association between depression and incident stroke.

Sensitivity and subgroup analyses
In order to minimize the possibility of reverse causality, we conducted an additional analysis where we considered studies that excluded incident strokes that occurred in the first year of follow-up separately. Specific data were not available from two studies.32,37 While five of six studies with data available found a significant positive association between depression and a first-ever stroke (effect estimates >1.0, P<0.005),24,31,36,41,42 one study suggested a reverse causal inference for depressive symptoms and stroke after excluding the first 5 years of follow-up (HR 0.94, 95% CI 0.64–1.37; P=0.74).4 Synthesizing these risk estimates revealed a persistent predictive association between depression and incident stroke (overall RR 1.64, 95% CI 1.27–2.11; P<0.0001). The heterogeneity was moderate, with an I² statistic of 60% (P=0.03 by Cochran Q test). This association remained strong after including only studies that reported HRs (HR 1.62, 95% CI 1.17–2.23, P=0.003; I²=66%, P=0.018 by Cochran Q; n=5). Figure 3 illustrates the corresponding forest plot.

We repeated our primary analysis, but excluded studies that enrolled participants with cardiac disease at baseline, and still found a strong independent association between depression and incident stroke (overall RR 1.43, 95% CI 1.19–1.72; P<0.0001), with evidence of moderate
heterogeneity ($I^2=58\%, P=0.003$ by Cochran Q test). Analyzing studies by preferred reported risk estimates did not change the results. The pooled OR/RR was $1.42$ ($95\%$ CI $1.03–1.96$, $P<0.034$; $n=4$), and the pooled HR was $1.45$ ($95\%$ CI $1.13–1.85$, $P<0.003$; $n=11$), both with evidence of moderate heterogeneity ($I^2=67\%$, $P=0.027$ by Cochran Q, and $I^2=58\%$, $P=0.008$ by Cochran Q, respectively). The exclusion of studies that focused on either fatal or nonfatal strokes did not alter the results either (overall RR $1.36$, $95\%$ CI $1.24–1.50$, $P<0.0001$; $I^2=46\%$, $P=0.007$ by Cochran Q). A strong prospective association between depression and incidence of a first-ever stroke remained after focusing on studies that assessed all stroke subtypes (ie, ischemic and hemorrhagic stroke, TIA) as the outcome measure (overall RR $1.50$, $95\%$ CI $1.21–1.86$, $P<0.0001$; $I^2=62\%$, $P=0.004$ by Cochran Q). To minimize misclassification bias, we removed studies that primarily defined the exposure variable by self-reporting, and found a slightly higher risk of incident stroke for clinically diagnosed depression at baseline (overall RR $1.64$, $95\%$ CI $1.21–2.24$, $P=0.002$; $I^2=68\%$, $P=0.015$ by Cochran Q). Similar results to those obtained by the primary analysis were found when restricting the data synthesis to a minimum cohort size of 5,000 participants (overall RR $1.33$, $95\%$ CI $1.23–1.44$, $P<0.0001$; $I^2=35\%$, $P=0.076$ by Cochran Q) and a minimum duration of follow-up of 5 years (overall RR $1.41$, $95\%$ CI $1.27–1.56$, $P<0.0001$; $I^2=42\%$, $P=0.018$ by Cochran Q). Excluding the two registry-based cohort studies from the primary analysis did not change the results (overall RR $1.40$, $95\%$ CI $1.26–1.55$, $P<0.001$; $I^2=48\%$, $P=0.004$ by Cochran Q).

When we stratified the included studies by age, it appeared that the ≥65-year (overall RR $1.45$, $95\%$ CI $1.20–1.75$, $P<0.0001$; $I^2=55\%$, $P=0.029$ by Cochran Q) age-group had a slightly lower risk than the <65-year (overall RR $1.59$, $95\%$ CI $1.35–2.09$, $P<0.0001$; $I^2=52\%$, $P=0.035$ by Cochran Q) age-group. Extending the definition of depression to the additional or exclusive use of antidepressants resulted in an elevation of the prospective risk of stroke (overall RR $1.60$, $95\%$ CI $1.23–2.09$, $P<0.0001$; $I^2=64\%$, $P=0.026$ by Cochran Q).

**Publication bias**

The funnel plot appeared slightly asymmetric, with evidence of small-study effects (ie, tendency that smaller studies show larger treatment effects) demonstrated by both

**Figure 2** Forest plot showing the pooled adjusted risk estimates for first-ever stroke in depressed individuals.

**Note:** Weights are from random effects analysis.

**Abbreviations:** ES, effect size; CI, confidence interval; HR, hazard ratio; OR, odds ratio; RR, relative risk.
The major finding of this meta-analysis comprising almost 700,000 participants, consistent with previous analyses, is that depression increases the risk of first-ever stroke by 40% in the general population. Moreover, our results relating depression to incident stroke are corroborated by recently published similar pooled estimates with an increased overall risk of stroke (34% and 45%, respectively), yet we included nine new studies, adding more than 400,000 participants.\(^1\,^2\) This prospective association appears to be driven neither by the preexistence of clinically apparent cerebro- and cardiovascular disease nor by silent stroke.

While our synthesized results suggest a possible causal association between depression and stroke, recently analyzed data from a large prospective cohort study contradict this contention by showing a nullified incident risk of stroke once a different analytic approach was applied.\(^4\) In this study, the risk of incident stroke was significantly elevated by 60% for depressed individuals, yet no such association was evident with exclusion of incident events that occurred in the first 5 years of follow-up. This analytic “lag period” approach was chosen to avoid the impact of possible reverse causality between preexisting silent cerebrovascular disease and depression, based on the vascular depression concept.\(^9\,^30\) However, this cohort study was limited by the absence of adjustment for vascular risk factors, use of a nonspecific depression screening scale, and an upper age limit of study participants of 55 years.\(^3\) In contrast, several previous cohort studies did not show significant influence of a lag period on the prospective association between the risk of incident stroke.

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**Figure 3** Forest plot showing the pooled adjusted risk estimates for first-ever stroke in depressed individuals according to the analytic approach used in the included study. Lag period indicates that stroke cases that occurred in the first years of follow-up were excluded from the analysis to minimize the possibility of reverse causality.

**Note:** Weights are from random effects analysis.

**Abbreviations:** ES, effect size; CI, confidence interval.
and depression. This is also reflected by the results of our synthesized subanalysis on studies that accounted for residual confounding by excluding the initial follow-up period showing a nearly identical overall (41%) and even pronounced risk once a lag period was defined, as opposed to studies with no lag period (64% and 34%, respectively). Also, the fact that most studies we included in our meta-analysis adjusted properly for well-established vascular risk factors further substantiates the notion that the association between depression and stroke could be independent of coexisting vascular diseases.

In concordance with the original vascular depression concept, which specifically proposed an association between cerebral small-vessel disease and depression in the elderly, a recent meta-analysis confirmed cerebral small-vessel disease being fourfold more common in patients with late- than early onset depression. Another more recent meta-analysis on this subject reported similar though weaker results. However, the development of cerebral small-vessel disease increases with age and is related to an increased risk of stroke. Therefore, viewed in conjunction with the vascular depression concept, it could be anticipated that particularly the elderly suffering from depression have an increased risk of stroke. Still, we found a balanced risk of stroke across groups stratified by age (even slightly lower in absolute numbers for the ≥65-year than the <65-year age-group), and one study included in our meta-analysis even suggested a higher risk of stroke in younger people with depression. Specifically, data originating from the Framingham Heart Study showed that depressive symptoms were significantly associated with a fourfold-increased risk of a first-ever ischemic stroke in individuals younger than 65 years but not in those older than or equal to 65 years. This discrepancy of balanced (or even increased) risk of stroke in younger depressed individuals and the increasing risk of stroke with age in the general population may be partially explained by the fact that depressive individuals less commonly adhere to medication than nondepressed people, and consequently primary stroke prevention may be compromised in depressed individuals. Moreover, unhealthy lifestyle, including smoking, unfavorable food, and lack of physical activity, is also associated with depression and in turn an increased risk of stroke.

Consistent with our finding of a balanced risk of stroke between young and elderly individuals with depression, a previous systematic review of 23 etiological, phenomenological, and outcome studies did not find any identifiable distinguishing clinical features between early onset and late-onset depression. Additionally, it might be speculated that an increased stroke risk in depressed people may arise from pharmacological interferences with platelet aggregation, as recently discussed for antidepressant medication and even suggested by the increased stroke risk (60%) yielded from our subanalysis of six studies that defined depression in relation to the use of antidepressant medication. On the other hand, a recent study did not confirm an increased risk of stroke in individuals with manic or bipolar disorder, a mental illness also known to be associated with unhealthy lifestyle behavior and a pronounced burden of vascular risk factors. This dissonant increase of stroke risk in depressed but not manic or bipolar individuals might suggest that the underlying cerebral pathomechanisms might be specific for depression. Based on the findings of previous studies, these mechanisms might include cerebral inflammation, dysregulation of the hypothalamic–pituitary–adrenal axis, increased platelet reactivity, and autonomic dysfunction.

Our quantitative data synthesis is subject to limitations, mostly associated with clinical and methodological diversity among the included studies, as shown by the moderate heterogeneity in our pooled data set. Clinical heterogeneity is given by the variety of self-reporting mood scales used in the studies included in our analysis, potentially introducing misclassification and information bias. This may particularly apply to participants prescribed antidepressant medication possibly obscuring the actual prevalence of depression at baseline. Another shortcoming relates to the fact that most of the published studies defined the outcome measures as any stroke subtype instead of focusing on a specific subtype (eg, ischemic or hemorrhagic stroke) or etiology. Therefore, stroke subtype- or etiology-specific differences in the relationship between depression and stroke might remain undetected in our synthesized analysis. Particularly given that in young adults, causes of stroke are less frequently attributed to cerebrovascular diseases compared with the elderly, but comprise a broad spectrum of nonatherosclerotic causes (eg, arterial dissection, hypercoagulable state, nonatherosclerotic arteriopathies, or cryptogenic), our finding of balanced stroke risk in different age-groups must be considered with caution. This also relates to hemorrhagic stroke, which has a variety of underlying age-dependent etiologies itself.

Even though we excluded studies that had enrolled participants with a known history of stroke in order to minimize confounding effects arising from potential preexisting cerebrovascular disease, it remains speculative to what extent unknown subclinical cerebrovascular disease might have influenced stroke risk in our analysis. Many studies, however, have consistently shown that particularly severe strokes with...
prominent functional disability predict the development of depression.\textsuperscript{5,67} Conversely, it might be speculated that stroke survivors clinically less severely affected may have a lower, if any, risk of developing a depression. Nevertheless, we reduced the probability of coexisting undetected cerebrovascular disease by performing a sensitivity analysis, removing studies that included patients with a history of cardiac disease, and yielded similar results.

Conclusion

Whether the consistent association of depression and increased risk of first-ever stroke demonstrated in this meta-analysis is indicative of depression-induced neurovascular pathology or is rather attributed to an interplay between two mutually enhancing conditions needs to be further elucidated in long-term studies. These studies should focus on specific stroke subtypes and preferably enroll young individuals free of cerebrovascular and cardiovascular diseases and provide longitudinal cerebral structural and functional assessment.

Disclosure

The authors report no conflicts of interest in this work.

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