Statin use and the risk of *Clostridium difficile* infection: a systematic review with meta-analysis

Raseen Tariq, Dhruvika Mukhija, Arjun Gupta, Darrell S Pardi, Sahil Khanna

1Division of Gastroenterology and Hepatology, Mayo Clinic, Rochester, MN, 2Department of Internal Medicine, Rochester General Hospital, Rochester, NY, 3Department of Internal Medicine, Cleveland Clinic, Cleveland, OH, 4Department of Internal Medicine, University of Texas Southwestern Medical Center, Dallas, TX, 5Division of Gastroenterology and Hepatology, University of California San Diego, La Jolla, CA, USA

**Purpose:** Statins have pleiotropic effects beyond cholesterol lowering by immune modulation. The association of statins with primary *Clostridium difficile* infection (CDI) is unclear as studies have reported conflicting findings. We performed a systematic review and meta-analysis to evaluate the association between statin use and CDI.

**Patients and methods:** We searched MEDLINE, Embase, and Web of Science from January 1978 to December 2016 for studies assessing the association between statin use and CDI. The Newcastle–Ottawa Scale was used to assess the methodologic quality of included studies. Weighted summary estimates were calculated using generalized inverse variance with random-effects model.

**Results:** Eight studies (6 case–control and 2 cohort) were included in the meta-analysis, which comprised 156,722 patients exposed to statins and 356,185 controls, with 34,849 total cases of CDI available in 7 studies. The rate of CDI in patients with statin use was 4.3%, compared with 7.8% in patients without statin use. An overall meta-analysis of 8 studies using the random-effects model demonstrated that statins may be associated with a decreased risk of CDI (maximally adjusted odds ratio [OR], 0.80; 95% CI, 0.66–0.97; *P*=0.02). There was significant heterogeneity among the studies, with an *I*² of 79%. No publication bias was seen. Meta-analysis of studies that adjusted for confounders revealed no protective effect of statins (adjusted OR, 0.84; 95% CI, 0.70–1.01; *P*=0.06, *F*=75%). However, a meta-analysis of only full-text studies using the random-effects model demonstrated a decreased risk of CDI with the use of statins (OR 0.77; 95% CI, 0.61–0.99; *P*=0.04, *F*=85%).

**Conclusion:** Meta-analyses of existing studies suggest that patients prescribed a statin may be at decreased risk for CDI. The results must be interpreted with caution given the significant heterogeneity and lack of benefit on analysis of studies that adjusted for confounders.

**Keywords:** *Clostridium difficile* infection, incidence, meta-analysis, statins

**Introduction**

*Clostridium difficile* infection (CDI) is the most common cause of hospital-acquired diarrhea and is increasingly being recognized in the community. Despite extensive preventive efforts, such as antibiotic stewardship, and emerging treatment strategies, an increasing incidence and worsening outcomes of CDI have been demonstrated. Novel risk factors for CDI, such as the use of proton pump inhibitors, have been identified that place persons previously considered to be at low risk, now at risk for CDI. Some risk factors for CDI such as an aging, immunocompromised population are not modifiable. In this setting, innovative methods to reduce the incidence of CDI are required.
HMG-CoA reductase inhibitors (ie, statins) are among the most common medications prescribed in the USA. From 2003 to 2012, the percentage of American adults aged 40 years and older taking a statin increased from 18% to 26%. The American College of Cardiology and the American Heart Association recommend statin therapy for all patients with cardiovascular disease and increased cholesterol levels and for patients aged 40–75 years who have diabetes mellitus or an estimated 10-year risk of cardiovascular disease of 7.5% or higher. Although the approved indications to use statins are largely cardiovascular, they have been shown to improve outcomes in infections such as pneumonia, pulmonary hypertension, new-onset inflammatory bowel disease, venous thromboembolism, autoimmune conditions such as systemic lupus erythematosus, and certain cancers such as hepatocellular carcinoma and gastric cancer. Statin use has been found to prevent infections in patients with cirrhosis and to be associated with decreased risks of severe sepsis and decompensation and all-cause mortality in compensated liver disease secondary to hepatitis C.

By inhibiting the production of isoprenoid intermediates, which are required for the activation of intracellular messengers, statins have pleiotropic effects on inflammatory and immunomodulatory pathways. Conceivably, statins may modify the risk of CDI. This remains a pertinent question given the many number of patients taking statins and therefore are at risk for CDI. A retrospective study indicated that statin users may, in fact, be at higher risk for CDI given the ability of statins to affect the interaction of the C. difficile organism and its toxins with colonic epithelium. In contrast, a large case–control study demonstrated a 22% lower risk of CDI in statin users versus nonusers. Given the conflicting results, we performed a systematic review and meta-analysis to study the association between the use of statins and the risk of CDI.

**Patients and methods**

All procedures used in this meta-analysis were reported according to the Preferred Reporting Items for Systematic Reviews and meta-analyses (PRISMA) guidelines.

**Selection criteria**

The studies considered in this meta-analysis were case–control studies, cohort studies, or clinical trials that included a study population of patients who did and did not receive statin therapy and that evaluated the occurrence of CDI, with no restrictions on study setting (inpatient or outpatient). We excluded studies that did not evaluate CDI as an outcome. Studies were also excluded from meta-analyses if there were insufficient data to determine an estimate of an odds ratio and 95% CI. We included published full-text articles and studies in abstract form.

**Data sources and search strategy**

We conducted a comprehensive search of Ovid MEDLINE In-Process & Other Non-Indexed Citations, Ovid MEDLINE, Ovid Embase, Ovid Cochrane Central Register of Controlled Trials, Ovid Cochrane Database of Systematic Reviews, Web of Science, and Scopus from January 1978 through December 2016. The search strategy was designed and conducted by study investigators (SK and RT) and the Mayo Clinic library staff, independently. The search was limited to studies published in English. Controlled vocabulary supplemented with keywords was used to search for studies of statin use and CDI. Main keywords used in the search were the following: *Clostridium difficile*, *C. difficile*, *Clostridium difficile* infection, *CDI*, *Clostridium difficile*–associated diarrhea or CDAD, or pseudomembranous colitis AND hmg coa OR hydroxymethylglutaryl OR hmg OR coa OR coenzyme OR atorvastatin OR cerivastatin OR compactin OR fluindostatin OR lovastatin OR mevinolin OR pitavastatin OR pravastatin OR rosuvastatin OR simvastatin OR statin AND outcomes, infection. The detailed search strategy is shown in Table S1.

Two authors (SK and RT) independently reviewed the titles and abstracts of the identified studies, and those that did not answer the research question of interest were excluded. The full texts of the remaining articles were reviewed to determine inclusion criteria fulfillment. The reference lists of articles with information on the topic were also reviewed for additional pertinent studies. A flow diagram of the included studies is shown in Figure 1.

The ROBINS-I risk of bias was used by 2 investigators (SK and RT) to assess the methodologic quality of case–control and cohort studies. In this scale, observational studies were scored across 3 categories using the following parameters: selection (3 questions), classification of exposure (3 questions), classification of missing data (5 questions), and bias in the selection of reported result (4 questions). For each question, 1 point was given if the study met the criterion (Table 1). Studies with a cumulative score of 10 or more were considered to be of moderate to high quality. Any discrepancies were addressed by joint re-evaluation of the original article.
763 Total citations
574 Rejected after title review
189 Potentially relevant studies
172 Rejected after abstract review
17 Articles eligible for inclusion and detailed assessment of selection criteria
9 Rejected
7 Insufficient data
2 Not assessing risk of CDI
8 Included studies

Figure 1 Flow diagram of study selection process.
Abbreviation: CDI, Clostridium difficile infection.

Table 1 ROBINS-I tool for bias assessment

<table>
<thead>
<tr>
<th>Author, year</th>
<th>Selection of participants</th>
<th>Classification of exposure</th>
<th>Classification of missing data</th>
<th>Bias in measurement of outcomes</th>
<th>Bias in selection of reported result</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Case–control studies</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Motzkus-Feagans et al, 201119</td>
<td>3</td>
<td>2</td>
<td>4</td>
<td>3</td>
<td>n/a</td>
<td>12</td>
</tr>
<tr>
<td>Naggie et al, 201124</td>
<td>3</td>
<td>2</td>
<td>4</td>
<td>3</td>
<td>n/a</td>
<td>12</td>
</tr>
<tr>
<td>Nseir et al, 201325</td>
<td>2</td>
<td>2</td>
<td>3</td>
<td>n/a</td>
<td>10</td>
<td></td>
</tr>
<tr>
<td>Elashery et al, 201427</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>2</td>
<td>n/a</td>
<td>8</td>
</tr>
<tr>
<td>Kumarappa et al, 201226</td>
<td>2</td>
<td>1</td>
<td>3</td>
<td>1</td>
<td>n/a</td>
<td>7</td>
</tr>
<tr>
<td>Ewelukwa et al, 201429</td>
<td>1</td>
<td>1</td>
<td>3</td>
<td>2</td>
<td>n/a</td>
<td>7</td>
</tr>
<tr>
<td>Cohort studies</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>McGuire et al, 200918</td>
<td>1</td>
<td>1</td>
<td>3</td>
<td>2</td>
<td>n/a</td>
<td>7</td>
</tr>
<tr>
<td>Tartof et al, 201528</td>
<td>2</td>
<td>2</td>
<td>3</td>
<td>2</td>
<td>n/a</td>
<td>10</td>
</tr>
</tbody>
</table>

Notes: Scoring criteria (maximum score 15):
- Bias in selection of participants into the study
  - Was selection of participants into the study (or into the analysis) based on participant characteristics observed after the start of intervention?
  - Do start of follow-up and start of intervention coincide for most participants?
  - Were adjustment techniques used that are likely to correct for the presence of selection biases?
- Bias in classification of intervention/exposure
  - Were intervention/exposure groups clearly defined?
  - Was the information used to define intervention/exposure group recorded at the start of the intervention/exposure?
  - Could classification of intervention/exposure status have been affected by knowledge of the outcome or risk of the outcome?
- Bias due to missing data
  - Were outcome data available for all, or nearly all, participants?
  - Were participants excluded due to missing data on intervention status?
  - Were participants excluded due to missing data on any variables that was required for analysis?
  - Are the proportion of participants and reasons for missing data similar across interventions?
  - Is there evidence that results were robust due to the presence of missing data?
- Bias in measurement of outcomes
  - Could the outcome measure have been influenced by knowledge of the intervention/exposure received?
  - Were outcome assessors aware of the intervention/exposure received by study participants?
  - Were the methods of outcome assessment comparable across intervention/exposure groups?
  - Were any systematic errors in measurement of the outcome related to intervention/exposure received?
- Bias in measurement of outcomes
  - Is the reported effect estimate likely to be selected, on the basis of the results, from multiple outcome “measurements” within the outcome domain?
  - Is the reported effect estimate likely to be selected, on the basis of the results, from multiple “analyses” of the intervention–outcome relationship?
  - Is the reported effect estimate likely to be selected, on the basis of the results, from different “subgroups”?  

Abbreviation: n/a, not available.
Data abstraction
Data were independently abstracted to a predetermined collection form by 2 investigators (SK and RT). Data were collected for each study, including study setting and design, year of publication, location, primary outcome (CDI) reported, and number of patients in each group (exposed vs not exposed and CDI vs no CDI). Conflicts in data abstraction were resolved by consensus, referring to the original article.

Outcomes assessed
Our primary analysis focused on assessing the risk of CDI and its association with statin use in studies that adjusted for potential confounders and in full-text studies.

Statistical analyses
We used the random-effects model described by DerSimonian and Laird\(^1\) to calculate weighted summary estimates using generalized inverse variance. Adjusted odds ratios (ORs), when available, or ORs (calculated for each study) were used in the analysis. Summary estimates are presented as ORs with 95% confidence intervals (CIs). We assessed heterogeneity within groups with the \(I^2\) statistic, which estimates the proportion of total variation across studies that is due to heterogeneity in study patients, design, or interventions rather than chance; \(I^2\) values greater than 50% suggest substantial heterogeneity.\(^2\) The presence of publication bias was assessed by the visual inspection of funnel plots.\(^2\) All \(P\)-values were 2-tailed. For all tests (except for heterogeneity), a \(P\)-value <0.05 was considered statistically significant. Calculations were performed and graphs were constructed using RevMan (Review Manager, version 5.3; Cochrane Inc).

A priori-defined analyses including studies that controlled for confounders, moderate- to high-quality studies, and full-text studies only and studies that recruited inpatients only were performed.

Results
Search results
The described search strategy revealed 763 potentially relevant studies; titles were screened and relevant articles were identified (Figure 1). In all, 17 articles were reviewed, of which 9 were excluded for various reasons (Figure 1). A total of 8 studies were included in this meta-analysis, of which only 3 were abstracts; all 8 observational studies evaluated the risk of CDI with statins.\(^1,18,19,24-29\) Two other studies that described the risk of recurrent CDI with statins were separately analyzed.\(^30,31\)

Quality of included studies
The median ROBINS-I score for case–control studies was 9.3 (range, 7–12) and for cohort studies was 8.5 (range, 7–10) out of 15 points. Four of the 8 included studies were considered to be of moderate to high quality, with a cumulative score of 10 or more. Table 1 shows the methodologic quality of all included studies.

Characteristics of included studies
The included studies comprised a total of 156,722 patients exposed to statins and 356,185 controls, with 34,849 total cases of CDI (available in 7 studies, 1 study did not report the total number of patients included). The characteristics of the 8 included studies are shown in Table 2. Seven studies were performed in USA and 1 in Israel. The earliest study recruitment period began in 2002, and the latest ended in 2015. All observational studies assessed medication exposure through review of medical records.

Statin use and CDI risk
The rate of CDI in patients taking statins was 4.3% (6,828/156,722), compared with 7.8% (28,021/356,185) in patients not taking statins. An overall meta-analysis of all 8 studies using the random-effects model demonstrated that statins were associated with a 20% decreased risk of CDI (maximally adjusted OR, 0.80; 95% CI, 0.66–0.97; \(P=0.02\)) (Figure 2A). There was a significant heterogeneity among the studies, with an \(F\) of 79%. No publication bias was seen (Figure 2B).

Primary analyses
Of the 8 included studies, 4 studies had been adjusted for potential confounders (Table 2). Analysis of studies that adjusted for confounders revealed no protective effect of statins (adjusted OR, 0.84; 95% CI, 0.70–1.01; \(P=0.06\)) (Figure 3). Meta-analysis of only full-text studies using the random-effects model demonstrated a decreased risk of CDI with use of statins (OR 0.77; 95% CI, 0.61–0.99; \(P=0.04\), \(F=85\%\)) (Figure 4).

Subgroup analyses
Given the significant heterogeneity in meta-analysis of all the included studies, we performed subgroup analyses to better understand the heterogeneity. However, no single source of heterogeneity was identified; the \(F\) remained increased in the subgroup analyses.
Subgroup analysis of moderate- to high-quality studies

Four of the 8 included studies were considered to be of moderate to high quality based on ROBINS-I scoring. Subgroup analysis of only these studies also revealed a significantly decreased risk of CDI with the use of statins (OR 0.73; 95% CI, 0.6–0.89; \( P = 0.04 \), \( I^2 = 81\% \)) (Figure 5).

Subgroup analysis of studies with inpatients only

Five of the 8 studies included in our meta-analysis included inpatients,\textsuperscript{19,24,25,27,28} 2 included both inpatients and outpatients,\textsuperscript{26,29} and 1 study included only intensive care unit (ICU) patients.\textsuperscript{18} The subgroup analysis of studies that included only inpatients also revealed a decreased risk of CDI with use of statins (OR 0.81; 95% CI, 0.68–0.95; \( P = 0.01 \), \( I^2 = 76\% \)) (Figure 6).

Statin and CDI outcomes

During our search, we found 2 other studies that evaluated the risk of recurrent CDI with statins.\textsuperscript{30,31} Meta-analysis of these 2 studies revealed that statins had no effect on the recurrence of CDI (unadjusted OR, 0.92; 95% CI, 0.74–1.15; \( P = 0.47 \))
Two additional studies evaluated the association of statins with risk of 30-day CDI mortality.\textsuperscript{32,33} Meta-analysis of these 2 studies also revealed a nonsignificant result (unadjusted OR, 0.91; 95% CI, 0.34–2.39; \( P = 0.84 \)) (Figure 7B).

**Discussion**

To our knowledge, this study is the first meta-analysis to explore an association between statins and incident CDI. From our analysis of studies that were conducted over more
than a decade (2002–2012), we conclude that statins may be associated with a decreased risk of CDI, with a 20% risk reduction. However, no protective effect was seen by pooling only studies that had been adjusted for potential confounders. Five of the 8 included studies showed that statins decreased the risk of incident CDI, but 1 study showed an increased risk with the use of these agents.18 However, that study was limited to ICU patients with sepsis and included very few patients with CDI.

The development of CDI is secondary to antibiotic exposure, which leads to altered intestinal microbiota.34 This is followed by infection by a toxigenic strain of C. difficile via feco-oral transmission, which leads to diarrhea secondary to inflammation from toxin exposure. Factors that facilitate infection or increase the virulence of C. difficile are being increasingly recognized. These include host-related factors such as immunosuppression, advanced age, hospitalization, severe illness, gastrointestinal surgery, obesity, and acid suppression (eg, proton pump inhibitors),4,35–38 and organism-related factors such as expression of certain surface adhesins and flagellar genes and toxin C. difficile transferase.39,40 However, data are limited on organism- and host-specific factors that afford protection against CDI. Our study helps address a major gap in knowledge by proposing statins as having a potentially protective role in CDI. Statins are known to interfere with molecules involved in endothelial adhesion and transendothelial migration of polymorphonuclear cells to sites of inflammation.41 Statins also reduce the activation of sites of inflammation.41 Statins also reduce the activation of
Infection and Drug Resistance downloaded from https://www.dovepress.com/ by 54.70.40.11 on 24-Jan-2020
For personal use only.
Powered by TCPDF (www.tcpdf.org)

the monocyte/macrophage system and reduce the cytotoxicity of T cells.\(^4\) It is also possible that the protective effects of statins relate to \(C.\ difficle\) toxins targeting Rho-GTPase proteins in the host cytosol,\(^4\) which is also a major target for the action of statins.\(^4\) Another potential mechanism for the protection from CDI offered by statins could be the alteration of microbiota by statins, although the specific effects of statins on the intestinal microbiome are unknown. Statins are also known to be bactericidal in vitro and in vivo,\(^4\) but these effects have been seen at concentrations much higher than those achieved in the blood during treatment with statins; therefore, the bactericidal effect of statins most likely does not protect against infections directly.

We performed a subgroup analysis by separating studies that included inpatients only; this analysis also demonstrated a decreased risk of CDI with statin use. The results also remained statistically significant when the analysis was limited to moderate- to high-quality studies. Analysis of studies that had been adjusted for potential confounders revealed no protective effect of statin. It is possible that the protective effect seen in the overall analysis might be a healthy effect rather than a true association between statins and decreased CDI risk.

With \(C.\ difficle\) being declared an emergent threat by the Centers for Disease Control and Prevention, it has become increasingly important to continue to explore novel and innovative ways to combat it, especially if the intervention can have other potential health-related benefits concurrently, as with statins. Widespread use of statins may lead to a significant decrease in health care–related costs due to their indirect effects even outside their primary indications, such as CDI inhibition. Some studies recommend the use of statins for primary prevention of cardiovascular events, even without hyperlipidemia.\(^4\)

The strengths of our study include a comprehensive literature review with a large patient population and stability of our results in subgroup analysis. Our study also has several limitations. The individual studies included in our meta-analysis varied in several ways, including study design, patient population, study setting (inpatient vs outpatient vs ICU), NAP1 (North American pulsed-field gel electrophoresis type 1) status, use of International Classification of Diseases –Ninth Classification codes versus medical records, lack of information on obvious confounders such as exposure to antibiotics (class, duration, and dose) in all studies, and the duration, dose, and type of statins used. These different aspects led to substantial heterogeneity. Only 4 studies had been controlled for different confounding factors. We were unable to perform analyses in which all confounding factors could be accounted or controlled for, including continuous versus intermittent use of statin and duration and dose of statins and exposure to antibiotic use. Although no publication bias was seen on visual inspection of funnel plot, the results of this test should be interpreted with caution since fewer than 10 studies were included.

**Conclusion**

In conclusion, our study highlights that statins may have a role in the prevention of CDI. However, this protective
effect was not seen after adjusting for potential confounders. Nevertheless, providers should continue to encourage and reinforce the use of statins when indicated in patients, not only for their direct cardiovascular benefits, but also for indirect benefits such as prevention of CDI, especially in the setting of systemic antibiotic exposure. Elective withholding of statin medications in hospitalized patients with multiple risk factors for CDI should be discouraged. However, the results of this meta-analysis must be interpreted with caution given the significant heterogeneity among the included studies and discrepancies seen on subgroup analyses. Further prospective studies would be required to evaluate the protective effect of statins on reducing the risk of CDI, including dose and duration of these medications.

Acknowledgments
The abstract of this paper was presented as a poster at the American College of Gastroenterology Conference in Orlando, FL, on October 15th, 2017. The abstract was published in the conference abstract supplement of the American Journal of Gastroenterology (Am J Gastroenterol 112:SS6; doi:10.1038/ajg.2017.296).

Author contributions
RT: study design, acquisition of data, analysis and interpretation of data, drafting of the manuscript, critical revision of the manuscript for important intellectual content. DM: study design, acquisition of data, drafting of the manuscript. AG: study design, acquisition of data, drafting of the manuscript. SS: study design, analysis and interpretation of data, critical revision of the manuscript for important intellectual content. DSP: study design, interpretation of data; critical revision of the manuscript. SK: study concept and design, acquisition of data, analysis and interpretation of data, drafting of the manuscript, critical revision of the manuscript for important intellectual content, statistical analysis, administrative support, study supervision. All authors have approved the final version of the manuscript.

Disclosure
The authors report no conflicts of interest in this work.

References


Supplementary material

<table>
<thead>
<tr>
<th>No</th>
<th>Searches</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Clostridium difficile/</td>
</tr>
<tr>
<td>2</td>
<td>exp Enterocolitis, Pseudomembranous/</td>
</tr>
<tr>
<td>3</td>
<td>exp Clostridium Infections/</td>
</tr>
<tr>
<td>4</td>
<td>(((clostridium or clostroidal) adj3 (enterocolitis or enteritis or colitis or disease* or infection* or diarrhea or diarrhoea)) or “antibiotic associated colitis” or “bacillus difficilis” or “C difficile” or CDAD or clostridioses or clostridiosis or “clostridium difficile” or “clostridium difficultis” or “pseudomembranous colitis” or “pseudomembranous enteritis” or “pseudomembranous enterocolitis”).mp.</td>
</tr>
<tr>
<td>5</td>
<td>1 or 2 or 3 or 4</td>
</tr>
<tr>
<td>6</td>
<td>exp Hydroxymethylglutaryl-CoA Reductase Inhibitors/ or statins.mp.</td>
</tr>
<tr>
<td>7</td>
<td>(“hmg coa” or ((hydroxymethylglutaryl or hmg) adj (coa or coenzyme)) or atorvastatin or cerivastatin or compactin or fluidostatin or lovastatin or mevinolin* or pitavastatin or pravastatin or rosuvastatin or simvastatin).mp. [mp=title, abstract, heading word, drug trade name, original title, device manufacturer, drug manufacturer, device trade name, keyword, floating subheading].</td>
</tr>
<tr>
<td>8</td>
<td>exp hydroxymethylglutaryl coenzyme A reductase inhibitor/</td>
</tr>
<tr>
<td>9</td>
<td>or/6-8</td>
</tr>
<tr>
<td>10</td>
<td>6 or 7 or 8 or 9</td>
</tr>
<tr>
<td>11</td>
<td>exp survival/</td>
</tr>
<tr>
<td>12</td>
<td>exp death/</td>
</tr>
<tr>
<td>13</td>
<td>exp mortality/</td>
</tr>
<tr>
<td>14</td>
<td>mortality.fs.</td>
</tr>
<tr>
<td>15</td>
<td>exp survival analysis/</td>
</tr>
<tr>
<td>16</td>
<td>exp evidence based medicine/</td>
</tr>
<tr>
<td>17</td>
<td>exp meta analysis/</td>
</tr>
<tr>
<td>18</td>
<td>exp Meta-Analysis as Topic/</td>
</tr>
<tr>
<td>19</td>
<td>exp “systematic review”/</td>
</tr>
<tr>
<td>20</td>
<td>exp Guideline/ or exp Practice Guideline/</td>
</tr>
<tr>
<td>21</td>
<td>exp controlled study/</td>
</tr>
<tr>
<td>22</td>
<td>exp Randomized Controlled Trial/</td>
</tr>
<tr>
<td>23</td>
<td>exp triple blind procedure/</td>
</tr>
<tr>
<td>24</td>
<td>exp Double-Blind Method/</td>
</tr>
<tr>
<td>25</td>
<td>exp Single-Blind Method/</td>
</tr>
<tr>
<td>26</td>
<td>exp latin square design/</td>
</tr>
<tr>
<td>27</td>
<td>exp comparative study/</td>
</tr>
<tr>
<td>28</td>
<td>exp intervention studies/</td>
</tr>
<tr>
<td>29</td>
<td>exp Cross-Sectional Studies/</td>
</tr>
<tr>
<td>30</td>
<td>exp Cross-Over Studies/</td>
</tr>
<tr>
<td>31</td>
<td>exp Cohort Studies/</td>
</tr>
<tr>
<td>32</td>
<td>exp longitudinal study/</td>
</tr>
<tr>
<td>33</td>
<td>exp retrospective study/</td>
</tr>
<tr>
<td>34</td>
<td>exp prospective study/</td>
</tr>
<tr>
<td>35</td>
<td>exp clinical trial/</td>
</tr>
<tr>
<td>36</td>
<td>clinical study/</td>
</tr>
<tr>
<td>37</td>
<td>exp case–control studies/</td>
</tr>
<tr>
<td>38</td>
<td>exp confidence interval/</td>
</tr>
<tr>
<td>39</td>
<td>exp multivariate analysis/</td>
</tr>
<tr>
<td>40</td>
<td>((event adj based) or (meta adj analys*) or (systematic adj review*) or guideline* or (control adj3 study) or (control adj3 trial) or (randomized adj3 study) or (randomized adj3 trial) or (randomised adj3 study) or (randomised adj3 trial) or “pragmatic clinical trial” or (double adj blind*) or (double adj mask*) or (singl adj blind*) or (singl adj mask*) or (tripl adj blind*) or (tripl adj mask*) or (trebl adj blind*) or (trebl adj mask*) or “latan square” or placebo* or nocebo* or multivariate or “comparative study” or “comparative survey” or “comparative analysis” or (intervention adj2 study) or (intervention adj2 trial) or “cross-sectional study” or “cross-sectional analysis” or “cross-sectional survey” or “cross-sectional design” or “prevalence study” or “prevalence analysis” or “prevalence survey” or “disease frequency study” or “disease frequency analysis” or “disease frequency survey” or crossover or “cross-over” or cohort* or longitudinal study or longitudinal analysis or longitudinal study or longitudinal evaluation or longitudinal analysis or (retrospective or “ex post facto”) adj3 (study or survey or analysis or design)) or retrospective* or “prospective study” or “prospective survey” or “prospective analysis” or prospective* or “concurrent study” or “concurrent survey” or “concurrent analysis” or “clinical study” or “clinical trial” or “case control study” or “case base study” or “case referent study” or “case referent study” or “case referent study” or “case compeer study” or “case comparison study” or “matched case control” or “multicenter study” or “multi-center study” or “odds ratio” or “confidence interval” or “change analysis” or ((study or trial or random* or control*) and compar*)).mp,pt.</td>
</tr>
</tbody>
</table>

(Continued)
<table>
<thead>
<tr>
<th>No</th>
<th>Searches</th>
</tr>
</thead>
<tbody>
<tr>
<td>42</td>
<td>(clinical trial, all or clinical trial, phase i or clinical trial, phase ii or clinical trial, phase iii or clinical trial, phase iv or clinical trial or controlled clinical trial or multicenter study or observational study or randomized controlled trial or pragmatic clinical trial or comparative study or controlled clinical trial or guideline or practice guideline or meta analysis or multicenter study or observational study or randomized controlled trial or pragmatic clinical trial or systematic reviews) [Limit not valid in Embase, CCTR, CDSR; records were retained]</td>
</tr>
<tr>
<td>43</td>
<td>(editorial or erratum or letter or note or addresses or autobiography or bibliography or biography or comment or dictionary or directory or interactive tutorial or interview or lectures or legal cases or legislation or news or newspaper article or overall or patient education handout or periodical index or portraits or published erratum or video-audio media or webcasts) [Limit not valid in Embase, Ovid MEDLINE(R), Ovid MEDLINE(R) In-Process, CCTR, CDSR; records were retained]</td>
</tr>
<tr>
<td>44</td>
<td>limit to English language [Limit not valid in CDSR; records were retained]</td>
</tr>
</tbody>
</table>