Prevalence of potential drug-drug interactions and associated factors among outpatients and inpatients in Ethiopian hospitals: a systematic review and meta-analysis of observational studies

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Abstract

Background: Drug-drug interaction is an emerging threat to public health. Currently, there is an increase in comorbid disease, polypharmacy, and hospitalization in Ethiopia. Thus, the possibility of drug-drug interaction occurrence is high in hospitals. This study aims to summarize the prevalence of potential drug-drug interactions and associated factors in Ethiopian hospitals.

Methods: A literature search was performed by accessing legitimate databases in PubMed/MEDLINE, Google Scholar, and Research Gate for English-language publications. To fetch further related topics advanced search was also applied in Science Direct and HINARI databases. The search was conducted on August 3 to 25, 2019. All published articles available online until the day of data collection were considered. Outcome measures were analyzed with Open Meta Analyst and CMA version statistical software. Der Simonian and Laird’s random effect model, I² statistics, and Logit event rate were also performed.

Results: A total of 14 studies remained eligible for inclusion in systematic review and meta-analysis. From the included studies, around 8717 potential drug-drug interactions were found in 3259 peoples out of 5761 patients. The prevalence of patients with potential drug-drug interactions in Ethiopian hospitals was found to be 72.2% (95% confidence interval: 59.1, 85.3%). Based on severity, the prevalence of major, moderate, and minor potential drug-drug interaction was 25.1, 52.8, 16.9%, respectively, also 1.27% for contraindications. The factors associated with potential drug-drug interactions were related to patient characteristics such as polypharmacy, age, comorbid disease, and hospital stay.

Conclusions: There is a high prevalence of potential drug-drug interactions in Ethiopian hospitals. Polypharmacy, age, comorbid disease, and hospital stay were the risk factors associated with potential drug-drug interactions.

Keywords: Drug-drug interactions, Hospitals, Ethiopia

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Background

Drug-drug interactions (DDIs) are types of adverse drug events (ADEs) that can occur when the effect of a drug is altered by another drug that is taken. Commonly it ends up with a qualitative and/or quantitative change in drug action [1]. They may change the diagnostic, preventive, and therapeutic activity of any drug and results in treatment failure, the toxicity of medications, and alternation of drug efficacy [2].

It can be categorized based on the severity and mechanisms by which drugs interact with each other [3, 4]. Based on their severity, DDIs can be mild, moderate, or severe. Major DDIs may be life-threatening or may cause prolonged or permanent damage. Moderate DDIs may require medical intervention or change in therapy. Whereas minor DDIs do not usually require a change in therapy. Regardless of the DDI severity, the patient should be monitored for possible manifestations of the interaction [3]. DDIs can also be classified as pharmaceutical, pharmacokinetic, and pharmacodynamic nature of drugs; the influence of disease on drug metabolism; prescriber issues such as multiple drug prescription by multiple prescribers, inadequate knowledge of prescribers’ on DDIs or poor recognition of the relevance of DDIs by prescribers are among the risk factors significantly associated with the occurrence of potential DDIs [5–10].

DDIs are common in cardiovascular, Human Immunodeficiency Virus-infected, psychiatric patients, and renal and hepatic insufficiency (CKD, cirrhosis) patients. Because this type of patient requires multiple types of drugs, their kidney and liver may decrease the excretion and metabolize the ability of medications. Therefore, the occurrence of DDIs in this type of patient may be significant [5–7, 11, 12].

DDIs are more frequent in hospitalized patients, patients who stay in the hospital for a longer time, and/or receive more drugs per day [13–16]. Hospitalized patients are more likely to be affected by DDIs because of severe and multiple illnesses, comorbid conditions, chronic therapeutic regimens, poly-pharmacy, and frequent modification in therapy [17]. Among hospitalized patients, elderly patients are at higher risk of potential DDIs, and the occurrence of potential DDIs ranges from 3 to 69%, depending on the specific area and population. The increased prevalence was found to be related to the presence of multiple chronic illnesses, the use of multiple medications, and altered pharmacokinetics in elderly patients [8].

Physicians and pharmacists alert fatigue is a common reason for the occurrence of drug-drug interactions for patients receiving interacting drugs. Even though computerized DDI alert systems could decrease the occurrence of DDIs, numerous alerts produced by such system lead physician and pharmacist alert fatigue. This alert fatigue results in a considerable override of DDI alerts. A study done in Japan showed physicians override DDI alerts at a high rate in computerized drug interaction alert system [18].

DDIs may have undesirable or harmful effects in addition to their desirable effects [4]. Clinically significant DDIs may cause potential harm to patients, harmful outcomes, and resulting in an estimated cost of more than $1 billion per year to governmental health care system expenditure [19].

DDI is being an evolving public health problem currently [20]. In Ethiopia, now a day, polypharmacy is increasing due to a rise in the occurrence of comorbid conditions in the hospital health care system [21, 22], where large number of patients are hospitalized. So, there is a high possibility of DDIs. Furthermore, due to economic problems, the probability of monitoring patients with comorbid diseases using sophisticated instruments is not feasible; causing the patient to DDIs.

As a result, potential DDIs causing serious risk to patient health. Therefore, this study attempted to review and quantitatively estimate the prevalence of potential DDIs and associated risk factors in hospitals, both among inpatients and outpatients in Ethiopia.

Methods

Study protocol

The review protocol was created based on Preferred Reporting Items for Systematic Review and Meta-analysis (PRISMA). The checklist was strictly followed while reporting this systematic review and meta-analysis (Additional file 1: Table 1) [23]. The review protocol is registered on PROSPERO with reference ID number: CDR 42020149416. The published methodology is also available at https://www.crd.york.ac.uk/prospero/display_record.php?ID=CDR42020149416.

Screening and eligibility of studies

WA designed the study. Two authors WA and GA screened the title and abstracts of the studies based on the inclusion and exclusion criteria. They also collected the full texts, evaluated the eligibility of the studies for final inclusion, assessed the quality of the study, and analyzed the data. AI commented on the review and meta-analysis.
Inclusion and exclusion criteria

**Inclusion criteria**

- √ Observational studies addressing the prevalence of potential DDIs and conducted in Ethiopia (prospective, retrospective and descriptive cross-sectional studies)
- √ All male and female patients in any age (pediatrics, adults, and geriatric) and admitted to hospital wards or visited outpatients
- √ All published articles without time limit
- √ Patients who had any disease and admitted to hospital wards or visited outpatients
- √ Studies which were published by English language and provided sufficient data for the review

**Exclusion criteria**

- √ Articles with missing or insufficient outcomes
- √ Studies that were conducted in primary health care settings
- √ Articles not published in peer reviewed journal.

**Table 1** Quality assessment of included studies in the review

| Studies                              | Total scores | Quality   |
|--------------------------------------|--------------|-----------|
| Gunasekaran et al., 2016 [25]        | 9            | Moderate  |
| Behailu Terefe Tesfaye et al., 2017 [6] | 12           | High      |
| Diksis et al, 2019 [5]               | 12           | High      |
| Chelkeba L et al., 2013 [26]         | 12           | High      |
| B.Akshaya Srikanth et al., 2014 [27] | 12           | High      |
| Admassie, et al, 2013 [28]           | 10           | High      |
| Henok Getachew et al, 2016 [29]      | 12           | High      |
| Teka et al, 2016 [30]                | 12           | High      |
| Zeru Gebretsadik et al., 2017 [31]   | 11           | High      |
| Haftay Berhane Mezgebe, 2015 [7]     | 11           | High      |
| Teklay et al., 2014 [32]             | 11           | High      |
| Yesuf TA, et al, 2017 [33]           | 10           | High      |
| Tesfaye and Nedi, 2017 [34]          | 11           | High      |
| Kibrom et al, 2018 [35]              | 11           | High      |

Fig. 1 PRISMA flow diagram showing the selection process
Search strategy and data sources
We had searched literatures from a legitimate database such as HINARI, Science direct, PubMed/MEDLINE, Google Scholar, and Research Gate for English-language publications. The literature search was performed to retrieve relevant findings closely related to the prevalence of potential DDIs and associated factors with DDIs among outpatients and inpatients in Ethiopian hospitals.

The search was conducted with the aid of carefully selected search-words without specification in time. “Prevalence”, “occurrence”, “potential DDIs”, “associated factors” and “Ethiopia” were the search words used in this review and meta-analysis. AND/OR words were used for the identification of the articles. The search was conducted from August 3–25, 2019 and all published articles available online until the day of data collection were considered.

Data extraction
A standardized data extraction form was prepared in Microsoft Excel by the investigators. Important information which was related to study characteristics such as:

Table 2 General characteristics of studies included for systematic review and Meta-analysis

| Region   | Study area          | Author and publication year | Study design | Pathology         | Target population                          | Study setting         | Interaction database       |
|----------|---------------------|------------------------------|--------------|-------------------|---------------------------------------------|------------------------|---------------------------|
| Oromia   | Middle East Ethiopia, Adama | Gunasekaran et al., 2016 [25] | Retrospective CS | All                | All hospitalized patients                    | All wards             | Medscape online            |
|          | southeast of AA, Bishoftu | Behailu Terefe Tesfaye et al., 2017 [6] | CS | HIV/AIDS          | All HIV infected patients                    | ART Clinic            | Medscape online & Drug.com |
| South West Ethiopia, Jimma | Diksis et al., 2019 [5] | Prospective CS | Cardiac disorder | Cardiac adult patients | Medical ward | Micromedex 3.0 DRUG-REAX* |
|          | Chelkeba L et al., 2013 [26] | CS | Cardiac disorder | Patients on CV medication in OPD | Cardiac clinic | Micromedex 2* |
| Amhara   | North West Ethiopia, Gondar | B.Akshaya Srikanth et al., 2014 [27] | Prospective CS | All                | All hospitalized patients                    | Medical ward          | www.drugs.com              |
|          | Admasie et al., 2013 [28] | Retrospective CS | All | All hospitalized patients | Inpatients and Out patients | Micromedex 2* |
|          | Henok Getachew et al., 2016 [29] | Retrospective CS | All | All hospitalized pediatric patients | Pediatric ward | Micromedex 2 |
| Tigray   | Northern Ethiopia | Teka et al., 2016 [30] | CS | All | All hospitalized patients | Medical ward | Micromedex 2.0 |
|          | Zeru Gebretsadik et al., 2017 [31] | Retrospective CS | All | All patients who come for medical service | Outpatient pharmacy | Micromedex 2.0 |
|          | Hafay Berhane Mezgebe, 2015 [7] | Retrospective CS | Psychiatric illness | Patients with psychiatric illness | Psychiatric unit | Micromedex 2.0 Drug-Reax* |
|          | Teklay et al., 2014 [32] | Prospective CS | DVT | Patients on warfarin therapy | Medical ward | Micromedex 2* online |
|          | Yesuf TA, et al., 2017 [33] | CS | All | All hospitalized patients | Medical ward | Micromedex 2* |
| AA       | TASH | Tesfaye and Nedi, 2017 [34] | CS | All | All hospitalized patients | Medical ward | Medscape online |
| SPHMMC   | Kibrom et al., 2018 [35] | Retrospective CS | All | Adult patients | Medical ward | Micromedex 3.0 DRUG-REAX* |

Abbreviations: HIV Human Immune Deficiency Virus, AIDS Acquire Immune Deficiency Syndrome, ART Antiretroviral Therapy, CV Cardio Vascular, OPD Outpatient Department, CS Crosssectional Study, TASH Tikur Anbessa Specialized Hospital, SPHMMC Saint Paulos Millennium Medical College
| Region   | Author                                      | Pathology     | Target population                     | Study setting | No. of patients | No. of patients with DDIs (%) | Prevalence of patients with DDIs (%) | No. of potential DDIs | Major | Moderate | Minor | Unknown & Contraindication |
|----------|---------------------------------------------|---------------|----------------------------------------|---------------|-----------------|-----------------------------|--------------------------------------|----------------------|--------|----------|-------|--------------------------|
| Oromia   | Gunasekaran et al., 2016 [25]              | All           | All hospitalized patients              | All wards     | 300             | 267 (89.00%)               | 62 (23.2%)                          | 95 (35.58%)          | 110 (41.2%) |                     |        |                         |
|          | Behailu Terefe Tesfaye et al., 2017 [6]    | HIV/AIDS      | All HIV infected patients              | ART Clinic    | 350             | 350 (100.00%)             | 2 (0.08%)                           | 176 (72.69%)          | 662 (27.2%) |                     |        |                         |
|          | Diksis et al., 2019 [5]                    | Cardiac disorder | Cardiac adult patients             | Medical ward  | 200             | 195 (97.50%)               | 316 (32.7%)                         | 441 (45.6%)           | 210 (21.7%) |                     |        |                         |
|          | Chelleba L et al., 2013 [26]               | Cardiac disorder | Patients on CV medication in OPD       | Cardiac clinic | 322             | 297 (92.24%)               | 88 (29.6%)                          | 200 (67.34%)          | 9 (3.03%)     |                     |        |                         |
| Amhara   | B.Akshaya Srikanth et al., 2014 [27]       | All           | All hospitalized patients              | Medical ward  | 100             | 78 (78.00%)               | 53 (12.8%)                          | 253 (61.26%)          | 107 (25.9%) |                     |        |                         |
|          | Admassie, et al., 2013 [28]                | All           | All hospitalized patients              | Inpatients and Outpatient | 2180          | 711 (32.61%)               | 127 (59.5%)                         | 1020 (77.04%)         | 177 (13.4%) | Contraindication = 11 (0.83%) |        |                         |
|          | Henok Getachew et al., 2016 [29]          | All           | All hospitalized pediatric patients   | Pediatric ward | 384             | 176 (45.83%)               | 40 (10.2%)                          | 201 (51.15%)          | 152 (38.7%) |                     |        |                         |
| Tigray   | Teka et al., 2016 [30]                     | All           | All hospitalized elder patients       | Medical ward  | 140             | 87 (62.14%)               | 46 (51.6%)                          | 36 (43.9%)            | 0 (0.00%)     | Contraindication = 5 (6.1%) unknown = 22 (6.65%) |        |                         |
|          | Zeru Gebretsadik et al., 2017 [31]        | All           | All patients who come for medical service | Outpatient pharmacy | 596             | 275 (46.14%)               | 34 (1103%)                         | 210 (63.44%)          | 87 (263%)     |                     |        |                         |
|          | Haftay Berhane Megebe, 2015 [7]           | Psychiatric illness | Patients with psychiatric illness | Psychiatric unit | 216             | 176 (81.48)               | 198 (43.8%)                         | 232 (51.33%)          | 22 (48.7%)    | Contraindication = 13 (2.88%) |        |                         |
|          | Teklay et al., 2014 [32]                  | DVT           | Patients on warfarin therapy          | Medical ward  | 133             | 132 (99.25)               | 11,827.6%                          | 310 (72.43%)          | 0 (0.00%)     |                     |        |                         |
|          | Yesuf TA, et al., 2017 [33]               | All           | All hospitalized patients              | Medical ward  | 204             | 135 (53.43)               | 150 (80.6%)                         | 36 (19.35%)           | 0 (0.00%)     | Contraindication = 80 (43%) |        |                         |
| Addis Ababa | Tesfaye and Nedi, 2017 [34]              | All           | All hospitalized patients              | Medical ward  | 252             | 197 (78.17)               | 94 (13.1%)                          | 385 (53.55%)          | 240 (33.4%)   |                     |        |                         |
|          | Kibrom et al., 2018 [35]                  | All           | Adult patients                         | Medical ward  | 384             | 209 (54.43)               | 105 (35.7%)                         | 157 (53.4%)           | 32 (10.9%)     | Contraindication = 2 (0.68%) |        |                         |

Abbreviations: HIV Human Immune Deficiency Virus, AIDS Acquire Immune Deficiency Syndrome, ART Antiretroviral Therapy, CV Cardio Vascular, OPD Outpatient Department
sample size described, potential or actual DDIs assessed, measures in place to ensure that results are valid and limitations of the study listed. Each criterion is related to a quality assessment criterion with scores 0 or 1 and the total quality scores ranged from 0 to 12 (scores 0 to 6 = poor quality, 7 to 9 scores = moderate quality, 10 to 12 points = high quality) (Table 1).

Outcome measurements
The outcome measure in this review and meta-analysis is the prevalence of potential DDIs. It primarily aimed to assess the pooled estimates of potential DDIs in the hospitals of Ethiopia. This study has also two secondary outcome measures: Associated risk factors for potential DDIs and number of potential DDIs (major, moderate, and minor) in Ethiopian hospitals.

Data processing and statistical analysis
Analysis of the pooled estimate of outcome measures i.e. Prevalence of potential DDIs, as well as subgroup analysis, were done by Open Meta Analyst advanced software. CMA version-3 software was used for publication bias assessment. The presence of publication bias was evaluated by using Egger’s regression tests and presented with funnel plots of standard error. Furthermore, the precision was presented with the Logit event rate. A statistical test with a \( P \) value of less than 0.05 (one-tailed) was considered significant [36].

Heterogeneity assessment
Heterogeneity may be defined as any type of variability between studies in a systematic review and meta-analysis. When there is variability in participants, interventions, and outcomes studied, we call it clinical heterogeneity. In this review and meta-analysis, Der Simonian and Laird’s random-effects model were used by considering clinical heterogeneity among studies. Variability in study design and risk of bias may be described as methodological heterogeneity [37].

Variation in intervention effects being evaluated in different studies is defined as statistical heterogeneity. This type of heterogeneity is usually a result of clinical or methodological heterogeneity or both among studies. Statistical heterogeneity is assessed by using Cochran’s Q- statistics, chi-squared and I\(^2\) tests. In this review and meta-analysis, clinical heterogeneity of studies was assessed using I\(^2\) statistics. Based on the result of the statistical test, I\(^2\) statistics value of less than 25% were considered as low heterogeneity and I\(^2\) statistics value from 50 to 75% and I\(^2\) statistics value greater than 75% were considered as medium and high heterogeneity respectively [38].

Results
Article search results
A total of 69 articles were identified through the search strategy. After duplication was removed, 49 articles have remained for screening. From these, 30 articles were excluded by their titles and abstracts. The remaining 19 articles were then evaluated as per predetermined eligibility criteria for inclusion. Five articles were also excluded with justification (Additional file 2: Table 2). Finally, a total of 14 full-text articles that passed the eligibility criteria and quality assessment were included for final review and analysis (Fig. 1).

General characteristics of the included studies
A total of 14 studies were included for systematic review and meta-analysis and important information that were related to study characteristics were presented in Table 2. All studies employed were observational cross-sectional study designs i.e. six retrospectives cross-sectional study (CS); three prospective CS and five CS design. The year of publication of included studies ranges from 2013 to 2019. The study included a wide range of population
characteristics (pediatric, adult, and geriatric patients). Regarding geographic distribution, 14 studies were obtained from three regions and one city administration (Addis Ababa). The studies included all types of disease which had been treated in a medical ward and outpatient setting.

Nine articles analyzed patients with all type of pathologies without focusing on any specific disease, two articles analyzed patients with the cardiac disorder, one article studied HIV patients and one article analyzed patients with psychiatric disorders.

Nine articles studied DDIs in inpatient ward (seven articles in a medical ward; one article in a pediatric ward; one article in all wards); four articles studied DDIs in the outpatient setting (ART Clinic, Cardiac Clinic, Psychiatric unit, and Outpatient pharmacy) and one article studied inpatients and outpatient setting.

Among the fourteen studies analyzed, six different databases were used to detect potential interactions. About half of the studies used Micromedex® 2.0 database systems (seven articles; 50.0%), two articles (14.2%) used Medscape online, two articles (14.2%) used Micromedex® 3.0 database systems. The other three articles used Medscape online and drug.com, Drug.com and Micromedex online (Table 2).

Quality of included studies
The quality of the included studies ranges from moderate to high quality (Additional file 3: Table 3).

Study outcome measures
Prevalence of potential DDIs
The prevalence and number of potential DDIs for each study are presented in Table 3. From 14 studies, the pooled prevalence of patients with potential DDIs in Ethiopian Hospitals was found to be 72.2% with 95% CI between 59.1 and 85.3). Figure 2 showed heterogeneity across 14 studies was high (I² = 99.78%, p < 0.001).

Based on the severity of DDIs, the pooled prevalence of potential DDIs was 25.1, 52.8, 16.9, and 1.27% for major, moderate, minor potential DDIs and contraindications respectively. Figures 3, 4, and 5 showed heterogeneity across 14 studies was high.

| Studies                        | Estimate (95% C.I.) | Moderate potential DDIs/Total no. of potential DDIs |
|-------------------------------|--------------------|------------------------------------------------------|
| Gunasekaran et al 2016        | 0.356 (0.290, 0.413) | 95/267                                               |
| Tesfaye et al 2017            | 0.727 (0.709, 0.745) | 1767/2431                                            |
| Dale et al 2019               | 0.456 (0.426, 0.487) | 441/796                                              |
| Chekeba et al 2013            | 0.673 (0.620, 0.727) | 200/297                                              |
| Srikant et al 2014            | 0.613 (0.566, 0.660) | 263/413                                              |
| Admassese et al 2013          | 0.766 (0.741, 0.791) | 1020/1335                                             |
| Getachew et al 2016           | 0.511 (0.482, 0.541) | 201/393                                              |
| Tekle et al 2016              | 0.414 (0.310, 0.517) | 36/67                                                |
| Zeno et al 2017               | 0.395 (0.344, 0.446) | 230/353                                              |
| Berhanu Megesha 2015          | 0.499 (0.453, 0.544) | 232/465                                              |
| Tekey et al 2014              | 0.724 (0.692, 0.767) | 310/428                                              |
| Yasef TA et al 2017           | 0.135 (0.094, 0.176) | 36/66                                                |
| Tesfaye and Necl 2017         | 0.535 (0.499, 0.572) | 365/719                                              |
| Kirbom et al 2018             | 0.375 (0.320, 0.430) | 111/296                                              |
| Overall (P²=99.78 %, P<0.001) | 0.528 (0.431, 0.624) | 5297/8717                                             |
Based on the mechanisms of DDIs involved, seven studies documented well but the remaining seven studies didn’t document well the mechanisms of DDIs (Table 4).

**Factors associated with potential DDIs**
The factors associated with potential DDIs were related to patient characteristics (Table 5).

**Common interacting drug-combinations**
The most common contraindications, major, and moderate DDIs are presented in Table 6.

**Test of heterogeneity, subgroup analysis, and publication bias**

**Test of heterogeneity**
In this review and meta-analysis, there is clinical and statistical heterogeneity. The tests of heterogeneity showed significant heterogeneity ($I^2 = 99.78\%$, $p < 0.001$). To differentiate heterogeneity, sensitivity analysis, subgroup analysis, and Meta-regression was done.

**Sensitivity analyses**
There was no significant change in the degree of heterogeneity even if an attempt was done to exclude the expected outliers as well as one or more of the studies from the analysis. Therefore, fourteen studies were included for the meta-analysis.

**Subgroup analyses**
Subgroup analysis also conducted based on Region and Study setting. Subgroup analysis based on a region revealed that the highest prevalence of potential DDIs was observed at Oromia Region, 94.9% (95% CI: 90.3 to 99.5) followed by Tigray Region with a prevalence of 68.6% (95% CI: 42.6 to 94.5) (Fig. 6).

Subgroup analysis based on study setting revealed that the highest prevalence of potential DDIs was observed at outpatient: 80.0% (95% CI: 58.9 to 101.1 followed by inpatient: 73.2% (95% CI: 60.8 to 85.7 and inpatient and outpatient setting: 32.6% (95% CI: 30.6 to 34.6).

Univariate meta-regression for prevalence of potential DDIs revealed that sampling distribution is a source of heterogeneity (regression coefficient = 7.36; $p$-value = 0.0067) (Fig. 7).

**Publication bias**
Funnel plots of standard error with logit effect size i.e. event rate supplemented by statistical tests confirmed that there is no evidence of publication bias on studies reporting the prevalence of potential DDIs.

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**Table 4** Studies of the prevalence of DDIs according to the mechanisms involved in Ethiopian Hospitals

| Authors                     | Mechanism of DDIs | Pharmacokinetic | Pharmacodynamics | Unknown |
|-----------------------------|-------------------|-----------------|------------------|--------|
| Gunasekaran et al., 2016    | 164 (61.42%)      | 101 (37.83%)    | 2 (0.75%)        |
| Behailu Terefe Tesfaye et al., 2017 | 1059 (43.56%)   | 1335 (54.92%)   | 37 (1.52%)       |
| Diksis et al., 2019         | 245 (25.34%)      | 574 (59.36%)    | 148 (15.3%)      |
| Henok Getachew et al., 2016 | 197 (50.13%)      | 181 (46.06%)    | 15 (3.82%)       |
| Yesuf TA et al., 2017       | 142 (53.38%)      | 124 (46.62%)    | 0 (0.09%)        |
| Tesfaye and Nedi, 2017      | 358 (49.79%)      | 321 (44.65%)    | 40 (5.56%)       |
| Kibrom et al., 2018         | 142 (47.97%)      | 87 (29.39%)     | 67 (22.6%)       |

Footnote: Seven studies did not report the mechanisms of drug-drug interaction
and associated factors in Ethiopian Hospitals because there is no higher concentration of studies on one side of the mean than the other at the bottom of the plot (Fig. 8).

**Discussion**

This systematic review and meta-analysis aimed to review and summarize the prevalence of potential DDIs and associated factors through reviewing and quantitatively summarizing the pieces of evidence available in Ethiopia. It was conducted and attempted to analyze 14 original studies addressing the topic. From all included studies, 5761 patients were included for pooled estimation of the primary outcome. A total of 8717 potential DDIs was found in 3259 patients. This means 2.67 DDIs per patient was suffering at least one DDI (calculated by dividing the number of potential DDIs/number of patients who suffer at least one potential DDI). On the other word, 1.5 DDIs were occurred per 100 patients (calculated by dividing the number of potential DDIs by the number of patients).

The overall prevalence of patients with potential DDIs in Ethiopia was found to be 72.2% (95%CI: 59.1, 85.3%). Based on the severity of DDIs, the pooled prevalence of potential DDIs was 25.1, 52.8, 16.9, and 1.27% for major, moderate, minor potential DDIs and contraindications respectively. These potential DDIs are more likely to produce negative outcomes. The analysis showed a high prevalence of DDIs which indicates the countries drug-drug interactions problem in the Ethiopians Hospitals. So, prescribers should prescribe interacting drugs in a monitored way.

The review showed that all DDIs studies in Ethiopia assessed potential DDIs, while no study was performed on actual DDIs. This may be due to

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**Table 5** Associated factors for potential DDIs

| Factors                              | Description                                                                 |
|--------------------------------------|-----------------------------------------------------------------------------|
| No of prescribed drugs (Polypharmacy) | Patients taking three or more than three concomitant drugs are at higher risk of the occurrence of potential DDIs [27, 28] |
|                                      | There is an association of the occurrence of one or more potential DDIs with the number of medications prescribed per patient who took more than four medications [35] |
|                                      | Polypharmacy (five or more medications) is an important factor which leads to potential DDIs [5, 29–31, 33, 34] |
| Co-morbid disease                    | Co-morbid condition independently increased the potential DDIs almost 2-folds [33] |
| Age                                  | Older age was found to be predisposing factors for the occurrence of DDI [5, 28, 30, 31] |
|                                      | Potential DDIs were occurring more frequently in the age group of 2–6 years than any other age group of the pediatric population [29] |
| Hospital stay                        | The chance of taking multiple drugs increases with longer stays (greater than or equal to seven) in the hospital, which in turn increases the risk for potential DDIs [5] |
| International Normalized ratio (INR value) | Increase in international normalized ratio value was found to be strongly associated with DDI and hence the risk of bleeding [32] |

Footnote: Ten studies did not report the mechanisms of drug-drug interaction.
Fig. 6 Subgroup analysis of the prevalence of potential DDIs based on region

| Studies                  | Estimate (95% C.I.) | Prevalence of potential DDIs/Total patients |
|--------------------------|---------------------|---------------------------------------------|
| Gunesekaran et al        | 0.890 (0.855, 0.925) | 267/300                                     |
| Tesfaye et al            | 0.999 (0.995, 1.003) | 300/300                                     |
| Chisa et al              | 0.975 (0.967, 0.997) | 149/200                                     |
| Chekeba et al            | 0.922 (0.933, 0.952) | 207/322                                     |
| Subgroup Oromia (p^2=95.31 %, P=0.000) | 0.549 (0.503, 0.595)  | 1169/1172                                   |
| Srikanth et al           | 0.780 (0.699, 0.861) | 78/100                                      |
| Admasse et al            | 0.326 (0.306, 0.346) | 711/2100                                    |
| Getachew et al           | 0.450 (0.408, 0.508) | 176/394                                     |
| Subgroup Amhara (p^2=88.45 %, P=0.000) | 0.510 (0.305, 0.731)  | 905/2644                                    |
| Teka et al               | 0.621 (0.541, 0.702) | 87/140                                      |
| Zime G/Findik et al      | 0.461 (0.423, 0.501) | 275/596                                     |
| Berhane Megatu           | 0.815 (0.762, 0.867) | 178/216                                     |
| Teklay et al             | 0.962 (0.970, 1.007) | 113/133                                     |
| Yewul TA et al           | 0.598 (0.538, 0.668) | 109/216                                     |
| Subgroup Tigray (p^2=99.49 %, P=0.000) | 0.686 (0.624, 0.745)  | 778/1289                                    |
| Tesfaye and Nedi         | 0.702 (0.723, 0.733) | 197/202                                     |
| Kibrom et al             | 0.546 (0.494, 0.594) | 209/394                                     |
| Subgroup Addis Ababa (p^2=97.65 %, P=0.000) | 0.663 (0.430, 0.894)  | 404/634                                     |
| Overall (p^2=99.78 %, P=0.000) | 0.722 (0.591, 0.853)  | 3259/3741                                   |

Fig. 7 Univariate meta-regression model using sample size for the prevalence of potential DDIs
identifying actual DDIs is much more complicated than potential DDIs.

The analysis showed that the occurrence of potential DDIs in the inpatient and outpatient settings reported by studies (inpatient: 73.2% (95% CI: 60.8 to 85.7%; outpatient: 80.0% (95% CI: 58.9 to 101.1%); inpatient and outpatient setting: 32.6% (95% CI: 30.6 to 34.6%). The prevalence of potential DDIs in this review is higher than another review in a developed nation in which 33% of the general population developed potential DDIs [39]. The high incidence of DDIs may be associated with a high number of drugs per prescription that was observed in individual studies. Otherwise, our review included only patients treated in the inpatient department, outpatient department, HIV clinic, and heart and cardiac clinics.

The prevalence of potential drug-drug interactions in the outpatient setting is higher than in the inpatient setting. The possible explanations for this finding. First, ART Clinic, Cardiac Clinic, Psychiatric unit, and Outpatient pharmacy were considered as outpatient settings. Moreover, the number of drugs and pathologies treated was different. This result helps hospitals to plan activities to prevent the occurrence of potential DDIs. So, hospitals can able to identify and follow up potential risk health care areas i.e. outpatient, inpatient, and other areas and help patients easily.

Similarly, this review showed all (100%) HIV infected patients treated in the outpatient setting [6]97.5% of adult patients with heart diseases treated in inpatient ward [5] and 92.23% cardiac disorder patients treated in the outpatient setting [26] were susceptible to DDIs. A high number of prescribed drugs, prescribing drugs with many potential DDIs, pharmacodynamics nature of drugs used in cardiology, and the influence of heart disease on drug metabolism may cause the high occurrence of potential DDIs in this group of patients. One finding in a developed country showed that 80% of hospitalized patients with heart diseases were susceptible to DDIs [40].

In this review and meta-analysis, age, polypharmacy, comorbid disease, and hospital stay were significantly associated with the occurrence of potential DDIs in the hospitals. Similarly, the finding from a review in a developed country highlighted these risk factors. Many studies had emphasized that the high occurrence of potential DDIs in old age is due to physiological changes related to age, comorbid diseases, and a high rate of medication use [41]. In addition to older age, potential DDIs were occurring more frequently in the age group of 2–6 years than any other age group of the pediatric population [29]. This is due to wide-ranging of patient ages and body-weights, limited physiologic reserve, medications dosing errors and ineptitude to properly communicate with healthcare workers [8].

Different studies were also supported as polypharmacy and comorbid disease increases the likelihood of the occurrence of potential DDIs [15, 33, 42, 43]. In the review, taking five or more medications was an important factor for identifying potential DDIs.
that leads to potential DDIs [5, 29–31, 33, 34]. This may be due to the probability of taking interacting drugs is increased. Likewise, the prevalence of potential DDIs from this review would likely have been higher.

Comorbid disease increases the occurrence of potential DDIs. Because the reason might be, the drugs prescribed for the comorbid disease are often used in combination that leads to the possibility of the occurrence of potential DDIs. Furthermore, increased hospital stay leads to the occurrence of potential DDIs. Since, hospitalized patients are more likely exposed to multiple illnesses, comorbid conditions, chronic therapeutic regimens, poly-pharmacy, and frequent modification during their stay of therapy [17].

The first limitation of this review and meta-analysis was the drug-drug interactions found were the only potential and doesn’t address the actual DDIs due to a lack of studies. Some of the studies included in the review and meta-analysis had small sample sizes. These might have led to bias. The other limitation of this review was Egger’s test funnel plots revealed as there is no publication bias but this estimation may not be accurate as small studies are included for the review and there are studies that had small size. The fourth limitation of this study was clinical heterogeneity among included studies, so it should be considered with caution. The classification of severity may be defined differently between studies, so this may be another limitation of this study.

Conclusion
The prevalence of patients with potential DDIs in Ethiopian Hospitals was found to be high i.e. 72.2% (95% CI: 59.1, 85.3%). As of these, the most prevalent DDIs were moderate severity, 52.8%. In this review polypharmacy, age, comorbid disease, and hospital stay were the risk factors associated with potential DDIs. This review and meta-analysis had considerable clinical heterogeneity among included studies, so it should be considered with caution.

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WA designed the study, WA and GA collected scientific studies, assessed the quality of the study, extracted and analyzed the data. AI commented on the review. WA also prepared the manuscript for publication. All authors have read and approved the manuscript.

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