Efficacy and safety of ledipasvir/sofosbuvir for hepatitis C among drug users: a systematic review and meta-analysis

Xue Yang, Yang Tang, Di Xu, Guang Zhang, Peng Xu, Houlin Tang* and Lin Pang

Abstract

Background and aims: Limited data is available on the efficacy of direct acting anti-viral drugs on hepatitis C in drug users. The aim of this meta-analysis was to comprehensively analyze the efficacy and safety of LDV/SOF in drug users infected with the hepatitis C virus (HCV).

Methods: The PubMed, Cochrane library, Embase and Web of Science databases were searched for articles published till April 2021 on HCV-positive drug users who were treated with ledipasvir/sofosbuvir (LDV/SOF). The primary endpoint was pooled sustained virological response at 12 weeks (SVR12) with 95% confidence interval (95% CI). Funnel plots and Egger's test were used to assess the publication bias.

Results: A total of 12 studies and 711 subjects treated with LDV/SOF-based regimen for HCV were included, and the pooled SVR12 rate was 89.8% (95% CI 85.9–92.7). The pooled SVR12 rate of genotype 1 drug users was 92.4% (95% CI 88.6–95.0). Subgroup analysis showed that pooled SVR12 rates of patients treated with LDV/SOF and LDV/SOF ± RBV were 89.2% (95% CI 83.4–93.1), 90.4% (95% CI 83.6–94.5) respectively. In addition, the SVR12 rates were 88% (95% CI 70.7–95.7) for 8 weeks, 89.9% (95% CI 81.0–94.9) for 12 weeks and 82.2% (95% CI 24.9–98.5) for 24 weeks of treatment.

Conclusion: LDV/SOF is a safe and relatively effective treatment for hepatitis C in drug users.

Keywords: Hepatitis C, Ledipasvir and sofosbuvir, Drug users, SVR12, Meta-analysis

Introduction

Chronic hepatitis C can lead to liver fibrosis, which eventually progresses to cirrhosis and increases the risk of primary liver cancer [1]. Although the World Health Organization (WHO) has set a goal to eliminate hepatitis C virus (HCV) globally by 2030, its 2017 Global Hepatitis Report shows that 71.1 million people were chronically infected with HCV by 2015 [2, 3]. This goal has been achieved in Iceland, but is challenging to tackle in the United States and sub-Saharan Africa since the HCV epidemic in these regions is related to intravenous drug use [4]. The prevalence of HCV infection among drug users ranges from 30 to 70% [5].

Furthermore, even a considerable burden of HCV infection does not prevent drug use among the long-term abusers [6]. The HCV load is significantly increased in people who inject drugs (PWID), and drug use accounts for 23% of the newly identified infections [2, 7]. Furthermore, prolonged drug use in the absence of suitable interventions can significantly increase the risk of HCV infection [8].

Many PWID with chronic hepatitis C could not be treated in the past due to concerns regarding their alcohol consumption, pre-existing psychiatric disorder, and intravenous drug use [9]. Since 2014, direct-acting antiviral (DAA) drugs have completely revolutionized the treatment of chronic HCV infection, and the...
combination of 2–3 DAA s achieved sustained virological response (SVR) in more than 95% of the treated patients without requiring any interferons [10]. Ledipasvir/sofosbuvir (LDV/SOF) is the first DAA combination approved by the American Association for the Study of Liver Diseases (AASLD) and the American Infectious Disease Society for treating HCV genotype 1 [11]. Moreover, the fixed dose combination of the NS5B polymerase inhibitor SOF and the NS5A inhibitor LDV has marked a new era for patients with chronic HCV with genotype 1a, 1b, and 4 because it is the first drug to be approved by the FDA that does not include peginterferon (PEG) or ribavirin (RBV) [12]. The Asian-Pacific Association for the Study of the Liver (APASL) guidelines advises Ledipasvir (90 mg/day) with sofosbuvir (400 mg/day) for 12 weeks is also associated with high SVR12 rate in HCV genotype 2 infected patients including treatment-experienced and those with cirrhosis [13]. In according to the European Association For The Study Of The Liver(EASL) guidelines, all people who inject drugs (PWIDs) who are infected with HCV, including those receiving OST, those with a history of injecting drug use and those who recently injected drugs, should be treated with the general recommendations [14].

LDV/SOF with or without RBV was the most frequently used medication regimen [15]. Now, none of the guidelines specify a priority treatment plan for HCV treatment for drug users. Whether LDV/SOF has advantages in treating drug users is a question worthy of research.

The poor compliance of chronic drug users is a persistent barrier to treating HCV. In addition, the high cost of DAA drugs and the risk of re-infection further limits the benefits of this treatment regimen among drug users [16]. There are relatively few studies in the effects of antiviral treatment in this population, and fewer on the efficacy of LDV/SOF regimens. Therefore, the aim of this study is to systematically review the studies evaluating the efficacy and safety of LDV/SOF in achieving SVR12 in drug users with HCV.

Methods

Literature search strategy

The study was conducted according to the preferred reporting items for systematic review and meta-analysis (PRISMA) protocols [17]. The Pubmed, Cochrane library, Embase and Web of Science database were searched independently by two investigators (X.Y. and T.Y.) for articles published after April 2021 using the following MeSH terms: “Hepatitis C” (e.g. “HCV”); “Drug User” (e.g. “People who inject drugs”, “IDUS”); “ledipasvir, sofosbuvir drug combination” (e.g. “Harvoni”). The complete literature search strategy for the four databases can be found in Additional file 1. The bibliographies of the selected articles were also searched manually for additional studies. The clinical trial registry for additional trials was also checked. There were no filters for language or publication date.

Study selection

The titles and abstracts were independently screened by two investigators (X.Y. and T.Y.), and the eligible studies were additionally validated by a third investigator (X.D.). The inclusion criteria were as follows: (1) study subjects were hepatitis C patients who use or inject drugs, (2) ledipasvir + sofosbuvir intervention, and (3) clear SVR12 as the outcome. The exclusion criteria and the number of excluded studies were shown in Fig. 1.

Data extraction

The following data was extracted by the two primary investigators using a standard form: (1) study characteristics (writer, region, publication year, study period, study design, setting), (2) drug use characteristics (definition of recent drug use; OST, OAT), (3) treatment characteristics (medications, treatment duration, drug dose, HCV treatment experience), (4) patient characteristics (genotype, sex, age, weight, BMI, race, education, aboriginals, homelessness/unstable housing, employment status, relationship status, involvement in sex work, imprisonment record, annual Income, alcohol overuse, HIV co-infection, HBV co-infection, cirrhosis, prior history of HCC, adherence support), and (5) outcome characteristics (number of SVR12, relapse, reinfection, virological failure, adverse events). The inconsistencies were then confirmed by the third investigator. Since most of the included literature was concerned with LDV/SOF and other drugs, we initially extracted all drug information to obtain complete baseline data. Some relevant information was obtained from ClinicalTrials.com via the NCT numbers.

Quality assessment

The quality of the observational studies was assessed with the Newcastle–Ottawa quality assessment scale (NOS). The studies were scored on the basis of three aspects: selection (4 points in total), comparability (2 points in total) and outcome of study participants (3 points in total). Total score of less than 5 was considered low quality, 6–7 as moderate quality, and greater than 8 as high quality [18]. The Cochrane Collaboration’s tool was used to assess the risk of bias in the randomized controlled trials (RCTs). Seven domains were evaluated: (1) random sequence generation (selection bias), (2) allocation concealment (selection bias), (3) blinding of participants and personnel (performance bias), (4)
assessment (detection bias), (5) incomplete outcome data (attrition bias), (6) selective reporting (reporting bias), and (7) other bias. Each domain was assessed separately with three options: “Low risk”, “Unclear risk” and “High risk” [19]. Two investigators performed the primary assessment, and the third verified and summarized the results.

Outcomes
The primary outcome was the proportion of patients who achieved a sustained virological response (SVR) 12 weeks after discontinuation of treatment (SVR12). The secondary outcomes were as follows: (1) relapse—recurrence of HCV RNA within the post therapy follow-up period [20], (2) reinfection—detection of HCV RNA following end of treatment response or following sustained virologic response [21], (3) virological failure—increase in the HCV RNA level to at least 100 IU/ml from < 15 IU/ml during treatment, by > 1 log10 IU/ml from the lowest levels attained during treatment, or to at least 15 IU/ml after 6 weeks of treatment [22], and (4) adverse events (AEs; any grade. e.g. fatigue, headache, nausea), severe adverse events (SAEs. e.g. death) and discontinuation due to AEs.

Statistical analyses
The outcomes of proportion were pooled using the Wilson score method, and 95% confidence intervals (95% CIs) were used to compare the safety and efficacy of the pooled SVR12 rate with the inverse variance method. Heterogeneity across the included studies was assessed with Cochran Q-statistics and I² statistics. The random effects model was used in case of significant heterogeneity (P<0.10 and I²>50%), otherwise the fixed effects model was adopted. To further evaluate the efficacy and safety of DAA regimes, SVR12 was analyzed in subgroups stratified on the basis of genotypes, sex, presence/absence of intravenous drug use, HCV treatment experience, presence/absence of OST, HIV co-infection, alcohol overuse, homelessness or unstable housing, employment status, and relationship status. Likewise,
the efficacy and safety of LDV/SOF were evaluated in the different treatment regimen and treatment duration sub-

groups. Publication bias was analyzed with funnel plots and Egger’s test. All the statistical tests were two-sided, and

\( P \) value < 0.05 was considered statistically significant. All analyses were conducted using the Meta package in R

(4.0.2).

Results

Search results and study characteristics

We searched a total of 1720 articles from four databases. After removing the duplicate entries, 1349 articles were

scanned further and 55 were selected after excluding those with irrelevant titles and abstracts. After reading

the complete articles, 43 were excluded for various reasons and 12 (11 full-length articles and 1 abstract) [23–34]

were included for the final analysis (Fig. 1). Given the peculiarities of drug users, the sample size in each study was

relatively small. All included articles were published after 2016. The studies were either observational studies or RCTs. In every study population, the males outnumbered the females. The patients were treated with LDV/SOF for 8, 12 and 24 weeks. The characteristics of the studies and drug users were summarized in Tables 1 and 2. The Prisma checklist of this study was shown in the Additional file 4.

Quality of the included studies

Six observational articles were assessed by NOS, and showed moderate quality with an average score of 7. Two studies were of high quality and three of medium quality. Six RCTs were assessed by the Cochrane Collaboration’s tool. None of the trials were conducted in a double-blinded manner, and only two mentioned random assignment, indicating higher risk of bias. The quality assessment of the included studies was summarized in the Additional file 2/Table 1, 2 and 3.

Efficacy of outcomes and SVR12 rate

The SVR12 rate of HCV infection was available for 1069 cases. The pooled estimated SVR12 rate from random-
effects model was 90.6% (95%CI 87.1–93.3, \( I^2 = 59.9\%, P < 0.01 \)) (Fig. 2). No publication bias was found in studies (\( t = 1.6.3, P = 0.13 \)), and the funnel plot and Egger’s test results were shown in the Additional file 3/Figs. 1and2. The SVR12 rate of LDV/SOF regimen was available for 711 cases. The pooled estimated SVR12 rate from the random-effects model was 89.8% (95%CI 85.9–92.7, \( I^2 = 47.6\%, P = 0.03 \)) (Fig. 3). No publication bias was

Table 1 Main characteristics of the studies included in meta-analysis

| Study | Study design | Study period | Publication type | Region | Setting | Regimen | Duration (weeks) | Daily dose of LDV/SOF |
|-------|-------------|--------------|-----------------|--------|---------|---------|------------------|----------------------|
| Akiyama et al. [33] | Randomized trial | NA | Abstract | Sub-Saharan Africa | Multi-center | LDV/SOF | NA | NA |
| Akiyama et al. [34] | Randomized trial | 2013.10–2017.4 | Full-length | New York | Multi-center | LDV/SOF | 4/8/12 | NA |
| Gayam et al. [23] | Retrospective cohort study | 2016.1–2017.12 | Full-length | America | Multi-center | LDV/SOF ±RBV | NA | NA |
| Alimohammadi et al. [24] | Retrospective study | 2015.9–2019.2 | Full-length | Canada | Multi-center | LDV/SOF | NA | NA |
| Coffin et al. [25] | Two-arm randomized trial | 2015–2017 | Full-length | San Francisco | Single-center | LDV/SOF | 8 | 80/400 mg |
| Schütz et al. [26] | Observational study | 2015.9–2016.9 | Full-length | Austria | Multi-center | LDV/SOF | 8 | 90/400 mg |
| Øvrehus et al. [27] | Randomized trial | 2015.4–2016.4 | Full-length | Denmark | Single-center | LDV/SOF/RBV | 4 | 90/400 mg |
| Trabut et al. [28] | Retrospective case–control study | NA | Full-length | French | Multi-center | LDV/SOF ±RBV | 8/12 | NA |
| Grebely et al. [29] | Randomized trial | 2012.10–2016.5 | Full-length | 6 Study locations | Multi-center | LDV/SOF ±RBV | 8/12/24 | 90/400 mg |
| Morris et al. [30] | Observational study | 2016.3–2017.2 | Full-length | Australia | Single-center | LDV/SOF | 8/12/24 | NA |
| Read et al. [31] | Observational cohort study | 2015–2017 | Full-length | Australia | Single-center | LDV/SOF | 8/12/24 | NA |
| Grebely et al. [32] | Randomized trial | 2013.11–2014.4 | Full-length | 88 study locations | Multi-center | LDV/SOF ±RBV | 8/12/24 | 90/400 mg |
| Study                        | Regimen              | SVR12 (n/N) | Definition of recent drug use (measurement method) | OAT/OST | Age (year) | Sex (M/F) | Cirrhosis | HCV genotype | Treatment history | HIV/HBV coinfection |
|-----------------------------|----------------------|-------------|-----------------------------------------------------|---------|------------|-----------|-----------|--------------|-------------------|---------------------|
| Akiyama et al. [33]         | LDV/SOF              | 82/90       | Drug use in the past or during DAA therapy (urine analysis and self-report) | NA      | 51.2 ± 11.0 | NA        | NA        | GT 1,4       | NA                | NA                  |
| Akiyama et al. [34]         | LDV/SOF              | 98/104      | Drug use in the past or during DAA therapy (urine analysis and self-report) | OAT:150 | 97/53      | 41        | 16        | 21           |                   |                     |
| Gayam et al. [23]           | LDV/SOF ± RBV        | 94/101      | Drug use during DAA therapy (urine analysis and self-report) | OAT:58  | 610 ± 9.8   | 93/59     | 53        | GT 1−4       | 29                | 44/NA               |
| Alimohammadi et al. [24]    | LDV/SOF              | 62/68       | Drug use during the past 6 months (urine analysis) | OST:23  | 55          | 56/18     | 15        | NA           | 12/NA             |                     |
| Coffin et al. [25]          | LDV/SOF              | 28/31       | Actively injecting drug use in the past or during DAA therapy (urine analysis) | NA      | 42.43 ± 11.9 | 25/6      | NA        | GT 1         | 0                 | 0/0                 |
| Schütz et al. [26]          | LDV/SOF              | 40/40       | Intravenous drug in the past or during DAA therapy (NR) | OAT:40  | 38.9 ± 8.7  | 29/11     | 0         | GT 1         | 0                 | 0/0                 |
| Ovrehus et al. [27]         | LDV/SOF/RBV          | 12/16       | Drug use in the past or during DAA therapy (self-report) | NA      | 53         | 26/48     | GT 1–3    | 0            | 0                 | NA/NA               |
| Trabut et al. [28]          | LDV/SOF              | 2/3         | Drug use in the past or during DAA therapy (NR) | OST:34  | 46.2 ± 7.3  | 42/8      | 28        | GT 1−4,6     | 12                | 4/1                 |
| Grebely et al. [29]         | LDV/SOF ± RBV        | 49/53       | Drug use 12 months ago (urine analysis) | OST:194 | 48 ± 10.7   | 141/53    | 70        | GT 1–6       | 42                | NA/NA               |
| Morris et al. [30]          | LDV/SOF              | 49/62       | Drug use in the past or during DAA therapy | OST:45  | 45.2 ± 10.5 | 82 / 38   | NA        | GT 1−3       | 10                | NA/NA               |
| Study | Regimen       | SVR12 (n/N) | Definition of recent drug use (measurement method) | OAT/OST | Age (year) | Sex (M/F) | Cirrhosis | HCV genotype | Treatment history | HIV/HBV coinfection |
|-------|---------------|-------------|-----------------------------------------------------|---------|------------|-----------|-----------|--------------|-------------------|---------------------|
| Read et al [31] | LDV/SOF | 30/38 | Injecting drug use in the past or during DAA therapy (urine analysis and self-report) | OST:18 | 45 (25–69) | 48/23 | NA | GT 1–3 | 6 | 8/0 |
|         | SOF/DCV | 23/28 |  |  |  |  |  |  |  |  |  |
|         | OBV/DCV/PTV/r±RBV | 6/6 |  |  |  |  |  |  |  |  |  |
| Grebely et al [32] | LDV/SOF | 45/48 | Drug use 12 months ago (urine analysis) | OST:70 | 47 ±11 | 48/22 | 7 | GT 1 | 8 | NA/NA |
found in studies ($t = 1.78, \ P = 0.11$), and the funnel plot
and Egger’s test results were shown in the Additional
file 3/Figs. 3and4. In eightstudies [23, 25, 26, 28, 29,
31, 32, 34] with a total of 503 patients with HCV geno-
type 1, 469 patients achieved SVR12 after LDV/SOF
treatment. The pooled estimation of SVR12 rate from
random-effects model was 92.4% (95% CI 88.6–95.0,
$I^2 = 31.7\%, \ P = 0.17$) (Fig. 4).

### Subgroup analysis of SVR12 rate

As shown in Table 3, the pooled SVR12 rate was sig-
ificantly higher in the multi-center (93.1%, 95% CI

| Response | SVR12 (N = 1069) | Heterogeneity | $P^b$ value | Studies |
|----------|------------------|---------------|-------------|---------|
|          | Total, n/N       | Rate (95% CI) | $I^2$ (%)   | $P^a$   |
| Overall  | 973/1069         | 90.6 (87.1–93.3) | 59.9 | 0.0040 | 12 |
| By settings |
| Multi-center | 761/814         | 93.1 (91.1–94.7) | 0.0 | < 0.0001 | 0.0040 | 8 |
| Single-center | 212/255        | 82.9 (77.7–87.1) | 0.0 | - | - | 4 |
| By genotypes |
| 1 | 512/549 | 92.4 (88.7–95.0) | 32.3 | 0.1638 | 0.2163 | 8 |
| 2 | 15/15 | 90.0 (61.6–98.0) | 0.0 | - | - | 3 |
| 3 | 36/44 | 80.9 (65.8–90.3) | 0.0 | - | - | 3 |
| 4 | 6/7 | 81.0 (41.8–96.2) | 0.0 | - | - | 2 |
| 6 | 1/1 | 75.0 (10.9–98.7) | - | - | - | 1 |
| By sex |
| Male | 260/294 | 88.6 (79.5–94.0) | 64.8 | 0.1247 | 0.8051 | 5 |
| FEMALE | 123/139 | 87.4 (80.5–92.1) | 0.0 | - | - | 5 |
| By the presence or absence of intravenous drug use |
| Yes | 174/197 | 88.1 (78.1–93.9) | 55.0 | 0.0214 | 0.8787 | 4 |
| No | 165/187 | 89.1 (74.9–95.7) | 69.3 | - | - | 4 |
| By HCV treatment experienced |
| Naive | 205/227 | 90.2 (78.8–95.8) | 72.0 | 0.1741 | 0.6157 | 3 |
| Experienced | 41/47 | 86.9 (73.8–94.0) | 0.0 | - | - | 3 |
| By the presence or absence of OST |
| Yes | 83/97 | 85.0 (76.2–90.9) | 0.0 | 0.8280 | 0.5122 | 3 |
| No | 103/126 | 81.6 (73.8–87.5) | 0.0 | - | - | 3 |
| By the presence or absence of HIV co-infected |
| HIV co-infected | 49/56 | 86.8 (75.3–93.4) | 0.0 | 0.2988 | 0.2620 | 3 |
| Non HIV co-infected | 145/154 | 93.5 (82.3–97.8) | 63.7 | - | - | 2 |
| By the presence or absence of alcohol overuse |
| Overuse | 85/91 | 92.7 (84.6–96.7) | 0.0 | 0.1705 | 0.0492 | 3 |
| No overuse | 71/86 | 82.2 (72.5–90.0) | 0.0 | - | - | 2 |
| By homeless or unstably housed |
| Homeless or unstably housed | 52/61 | 91.1 (64.8–98.3) | 39.7 | 0.0450 | 0.6915 | 2 |
| Stable housing | 45/49 | 86.4 (55.1–97.1) | 70.0 | - | - | 3 |
| By the presence or absence of employed |
| Employed | 14/16 | 83.9 (57.1–95.3) | 0.0 | 0.1438 | 0.5555 | 2 |
| Unemployed | 116/131 | 84.5 (75.9–90.4) | 68.5 | - | - | 3 |
| By living in stable relationship |
| Single | 57/61 | 92.9 (69.4–98.7) | 41.6 | 0.5195 | 0.7838 | 2 |
| In a relationship | 28/29 | 94.8 (80.0–99.0) | 0.0 | - | - | 2 |

*Test of heterogeneity

*Test for subgroup differences
91.1–94.7) compared to the single-center (82.9%, 95% CI 77.7–87.1) cohorts (P < 0.01). In terms of the HCV genotype, the pooled SVR12 rates for GT1, GT2, GT3, GT4 and GT6 were 92.4% (95% CI 88.7–95.0), 90% (95% CI 61.6–98.0), 80.9% (95% CI 65.8–90.3), 81% (95% CI 41.8–96.2) and 75% (95% CI 10.9–98.7) respectively. The pooled rates of SVR12 were similar between males and females [88.6% (95% CI 79.5–94.0) vs 87.4% (95% CI 80.5–92.1)], as well as between the intravenous and non-intravenous users [88.1% (95% CI 78.1–93.9) vs 89.1% (95% CI 74.9–95.7)]. In addition, 86.9% (95% CI 73.8–94.0) of HCV treatment-experienced patients achieved SVR12 compared to 90.2% (95% CI 78.8–95.8) of patients without prior HCV treatment. Pooled SVR12 rates were 85.0% (95% CI 76.2–90.9) for patients with OST and 81.6% (95% CI 73.8–87.5) for those without OST. The pooled rate of SVR12 was lower among patients with HIV co-infection [86.8% (95% CI 75.3–93.4)] compared to the non-HIV subgroup [93.5% (95% CI 82.3–97.8)]. SVR12 was achieved in 92.7% (95% CI 84.6–96.7) of patients with alcohol abuse compared to only 82.2% (95% CI 72.5–90.0) of the patients without alcohol abuse. Furthermore, 91.1% (95% CI 64.8–98.3) of the homeless or unstably housed patients had achieved SVR12, compared to 86.4% (95% CI 55.1–97.1) of those with stable homes. The pooled rates of
SVR12 were 83.9% (95% CI 57.1–95.3) for patients with jobs and 84.5% (95% CI 75.9–90.4) for the unemployed patients. Finally, the SVR12 rates were 94.8% (95% CI 80.0–99.0) and 92.9% (95% CI 69.4–98.7) for patients in stable versus unstable relationships respectively. No significant differences were observed in other subgroup analyses.

Subgroup analysis of LDV/SOF

We also conducted subgroup analysis on the basis of LDV/SOF regimen and treatment duration (Table 4). Among drug users that received LDV/SOF, 89.2% (95% CI 83.4–93.1) achieved SVR. In contrast, 90.4% (95% CI 83.6–94.5) of drug users treated with LDV/SOF ± RBV achieved SVR. Pooled rates of SVR12 were 88% (95% CI 70.7–95.7) at 8 weeks, 89.9% (95% CI 81.0–94.9) at 12 weeks, and 82.2% (95% CI 24.9–98.5) at 24 weeks. There were no significant differences between these subgroups.

Different types of drugs used in drug users

Nine studies included data on the types of drugs used by the patients (Table 5). The number of heroin, cocaine, methamphetamine, buprenorphine, methadone, and cannabis were compared. The heterogeneity analysis showed that the type of drug used was not significantly different among the patients.

Table 5 Types of drugs used by the drug users

| Drugs       | Safety | Rate% (95% CI) | Heterogeneity | Studies |
|-------------|--------|----------------|---------------|---------|
| Heroin      | 129/287| 51.0 (35.9–65.4)| 80.4          | 0.0016  | 4     |
| Cocaine     | 123/424| 25.1 (15.9–37.4)| 83.3          | 0.0005  | 4     |
| Methamphetamine | 58/103 | 58.2 (41.9–72.9)| 56.9          | 0.1279  | 2     |
| Buprenorphine | 78/486 | 15.0 (6.6–30.4) | 89.6< 0.0001  |         | 5     |
| Methadone   | 313/486| 55.7 (29.2–79.3)| 93.8          | <0.0001 | 5     |
| Cannabis    | 30/122 | 24.6 (6.2–61.5) | 92.1          | 0.0004  | 2     |
| Benzodiazepines | 59/272 | 24.0 (14.5–36.9) | 76.9          | 0.0132  | 3     |

Fig. 4 Forest plot of the risk ratio and risk difference of SVR12 in patients with genotype 1 using LDV/SOF in the included literature
cannabis and benzodiazepines users were 129, 123, 58, 78, 313, 30 and 59 respectively. The pooled SVR12 rates were 51% (95% CI 35.9–65.4), 25.1% (95% CI 15.9–37.4), 58.2% (95% CI 41.9–72.9), 15% (95% CI 6.6–30.4), 55.7% (95% CI 29.2–79.3), 24.6% (95% CI 6.2–61.5) and 24.0% (95% CI 14.5–36.9) for the heroin, cocaine, methamphetamine, buprenorphine, methadone, cannabis and benzodiazepines users.

Safety

Most of the included articles did not describe adverse events in detail, and only 8 had data on one or more indicators. As shown in Table 6, the cases with relapse, re-infection, virological failure, AEs, SAEs and discontinuation due to AEs were 4, 4, 11, 333, 17 and 2 respectively, and the pooled rates were 5.1% (95% CI 1.9–12.9), 5.7% (95% CI 2.2–14.2), 4.8% (95% CI 2.5–9.0), 77.8% (95% CI 64.5–87.1), 5.3% (95% CI 1.8–14.5) and 0.5% (95% CI 0.1–1.8). The most common AEs were fatigue (27.3%), nausea (16.1%), headache (13.9%), insomnia (5.1%), diarrhea (4.0%) and pain (2.7%). In patients with SAEs, one death occurred due to opioid drug overdose and one was related to asymptomatic neutropenia. For the two patients who discontinued treatment, detailed medical reasons were not provided.

Discussion

The pooled SVR12 rates of drug users treated with LDV/SOF was 89.8% (95% CI 85.9–92.7) in this meta-analysis, which was lower than that reported by Grebely et al. [35] (97%) for sofosbuvir/velpatasvir and glecaprevir/pibrentasvir(94.6%) [36] among drug users, and slightly higher than that achieved by elbasvir/grazoprevir (89.5%) [37] among drug users. Despite the high SVR12, sofosbuvir/velpatasvir was associated with high rates of AEs (83%), SAEs (7%), discontinuation due to AEs (1%) and virological relapse rates (25%) compared to LDV/SOF (AEs=77.8%, SAEs=5.3%, discontinuation due to AEs=0.5% and virological relapse=5.1%) [35, 38]. Therefore, LDV/SOF is a safe and relatively effective treatment option for treating HCV in drug users.

There are eight known genotypes of HCV, and nearly half (46%) of the global HCV genotypes are GT1, mainly in Europe, North America and Australia, followed by GT3 (30%) primarily distributed in South Asia, particularly the Indian sub-continent [39]. The pooled SVR12 rate was significantly higher in drug users infected with the GT1 (92.4%; 95% CI 88.6–95.0) compared to the total SVR12 rate (90.6%; 95% CI 87.1–93.3). SVR12 was only 80.9% (95% CI 65.8–90.3) in GT3 drug users treated with LDV/SOF. The efficacy of LDV/SOF against GT1 was also better compared to the other genotypes, which may be attributed to the small sample size of the latter. In addition, the SVR rate of paritaprevir, ritonavir, ombitasvir and dasabuvir with/out ribavirin was only 71% as opposed to the 91% achieved by boceprevir and telaprevir in drug users with GT1 virus [40, 41]. Thus, LDV/SOF is a suitable option for drug users with GT1 HCV. In addition to the HCV genotype, the use of LDV/SOF to treat HCV depends on whether the patient has liver cirrhosis [42]. Cirrhosis was mentioned in 8 of the 12 studies, of which two studies recruited subjects without cirrhosis, the exclusion criteria for two studies were decompensated cirrhosis, and the exclusion criterion for one study was that the Fib4 and fibrosis-cirrhosis index exceeded 3.25 and 1.25, respectively. However, there was no data on SVR12 of LDV/SOF in drug users with cirrhosis in all the included studies, and no related studies have been found. The next research needs to pay attention to this aspect.

In the 12 included studies, one study reported SVR12 rates of 95% (21/22) and 94% (45/48) for LDV/SOF + RBV and LDV/SOF respectively. Other studies had shown similar efficacies of LDV/SOF regimen with/out RBV, although inclusion of the latter increased the incidence of AEs and SAEs. Therefore, ribavirin is not recommended for treating HCV in combination with LDV and SOF [43–46]. Furthermore, our results indicate that treatment efficacy did not improve with the duration. While 12 weeks of LDV/SOF treatment had

| Outcomes                        | Safety | Heterogeneity | Studies |
|---------------------------------|--------|---------------|---------|
|                                 | Total, n/N | Rate% (95% CI) | I^2 (%) | P     |
| Virological relapse             | 4/81 | 5.1 (1.9–12.9) | 0.0 | 0.5813 | 2 |
| Virological reinfection         | 4/71 | 5.7 (2.2–14.2) | 0.0 | 0.7930 | 2 |
| Virological failure             | 11/250 | 4.8 (2.5–9.0) | 5.3 | 0.3480 | 3 |
| AEs                             | 333/447 | 77.8 (64.5–87.1) | 85.7 | 0.0001 | 4 |
| SAEs                            | 17/368 | 5.3 (1.8–14.5) | 76.2 | 0.0056 | 4 |
| Discontinuation due to AEs      | 2/453 | 0.5 (0.1–1.8) | 0.0 | 0.8377 | 2 |
greater efficacy compared to the other time points, the least efficacy was observed after 24 weeks. In addition, a meta-analysis showed that 24 weeks LDV/SOF resulted in more SAEs in GT1 chronic hepatitis C patients compared to 8 weeks and 12 weeks treatments [20]. Thus, we recommend a 12-week treatment course of LDV/SOF for drug users with HCV infection.

In the ITT analysis, the SVR12 rate was 95% among people without a history of injecting drug use, 92% among PWID not receiving OAT [47]. A major issue with antiviral therapy for drug users is poor compliance [7]. The pooled SVR12 rates for drug users was lower than that of LDV/SOF-treated children (99%, 95% CI 94–100) [48], adolescents (98%, 95% CI 93–100) [49] and subjects aged 65 years or older (97%, 95% CI 96–98) [50] who were not drug users. It was also lower than the SVR12 rates in HCV patients with HBV (100%) [51], compensated liver disease (94%, 95% CI 84–99) [52] and kidney transplant (100%, 95% CI 94–100) [53] who were treated with LDV/SOF and were not drug users. Eight studies included in this meta-analysis had information regarding compliance of drug users taking LDV/SOF, of which 5 reported a compliance rate > 80% through self-reporting or collection of remaining drug doses. This was consistent with the 99% and 98% compliance rates of HCV-positive drug users through self-reporting and residual pill counting respectively as reported by Cunningham et al. [54]. However, the patients that received treatment twice a day had lower compliance. Two studies [55, 56] showed that false reporting and the loss of pills were associated with an overestimation of cure rates by self-reporting and residual pill counting. Among the remaining studies, one reported poor compliance, one stated that patients with addiction had a compliance rate of 62%, and the third achieved 39.2% and 49.9% compliance rates through non-direct and modified direct observation respectively. Macias et al. [47] showed that the compliance rate of active drug users was 79%, while the SVR12 rate of non-drug users was 95%. Taken together, drug users have relatively lower compliance to anti-viral treatment compared to non-users, which can be attributed to the high rate of loss during follow-up [57]. In this meta-analysis, the compliance among drug users treated with LDV/SOF was fair, as only 4% (29/727) of the cases were lost to follow-up.

There were still some limitations in our study that ought to be considered. First, six of the included studies analyzed effects of LDV/SOF in combination with other drugs. Furthermore, the basic information of drug users treated only with LDV/SOF was not available and the subgroup analysis of LDV/SOF was incomplete. Second, it was prone to include overlapping patients in Ref No 29 and 32. Third, methamphetamine and heroin were the main drugs used by drug addicts, but there was no information regarding whether previous or recent use of different drugs affected SVR12. Fourth, the resistance of drug users to LDV/SOF was also unknown. Fifth, the sample size was small due to the few clinical studies published so far.

Conclusion
This meta-analysis is the first to provide a comprehensive analysis of LDV/SOF treatment for drug users, with a total of 711 participants from 12 individual studies. Although there were still some limitations, the data showed that LDV/SOF is a safe and relatively effective for drug users with hepatitis C. Larger and better designed studies are needed to evaluate the effects of LDV-SOF and the treatment resistance in drug users. Future studies need to focus on providing effective measures to improve medication compliance among drug users and reduce loss during follow-up.

Abbreviations
LDV/SOF: Ledipasvir/sofosbuvir; CI: Confidence interval; RBV: Ribavirin; NA: Not applicable; OBV: Ombitasvir; PTV: Paritaprevir; r: Ritonavir; DSV: Dasabuvir; VEL: Velpatasvir; DCV: Daclatasvir; SMV: Simeprevir; VOX: Voxilaprevir; NR: Not reported; PEG: Pegylated interferon; AE: Any adverse events; CAE: Common adverse events; SAEs: Serious adverse events.

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Authors’ contributions
The articles were collected and analyzed by XY, YT and DX, and XY wrote the manuscript. PX and GZ revised the manuscript and raised questions to further improve the manuscript. HL T and LP supervised the entire study. All authors read and approved the final manuscript.

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Availability of data and materials
All data generated or analyzed in this study are included in this article and its supplementary information files.
Declarations

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Authors declare no conflict.

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