Comparative Efficacy and Safety of 11 Drugs as Therapies for Adults With Neuropathic Pain After Spinal Cord Injury: A Bayesian Network Analysis Based on 20 Randomized Controlled Trials

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Objective: To provide an updated analysis of the efficacy and safety of drugs for the management of neuropathic pain (NP) after spinal cord injury (SCI) based on Bayesian network analysis.

Methods: A Bayesian network meta-analysis of literature searches within PubMed, Cochrane Library, Embase, and Web of Science databases from their inception to February 21 2021 was conducted without language restrictions. Paired and network meta-analyses of random effects were used to estimate the total standardized mean deviations (SMDs) and odds ratios (ORs).

Results: A total of 1,133 citations were identified and 20 RCTs (including 1,198 patients) involving 11 drugs and placebos for post-SCI NP selected. The 5 outcomes from all 11 drugs and placebos had no inconsistencies after Bayesian network analysis. BTX-A gave the most effective pain relief for the 4 weeks, following a primary outcome. No significant differences were found among drugs with regard to adverse events of the primary outcome. Gabapentin, BTX-A, and pregabalin were found to be the most helpful in relieving secondary outcomes of mental or sleep-related symptoms with differences in SMDs, ranging from −0.63 to −0.86. Tramadol triggered more serious adverse events than any of the other drugs with differences in ORs ranging from 0.09 to 0.11.

Conclusion: BTX-A, gabapentin, pregabalin, amitriptyline, ketamine, lamotrigine, and duloxetine were all effective for NP management following SCI. Lamotrigine and gabapentin caused fewer side effects and had better efficacy in relieving mental or sleep-related symptoms caused by SCI-related NP. Tramadol, levetiracetam, carbamazepine, and cannabinoids could not be recommended due to inferior safety or efficacy.

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Keywords: neuropathic pain, spinal cord injury, network meta-analysis, efficacy, safety
INTRODUCTION

Spinal cord injury (SCI) is a rare event with devastating consequences and an estimated average global incidence of 23 cases per million (1) and a prevalence of between 236 and 1,298 per million (2). Patients present with a range of functional impairments, including sensory, motor, and autonomic dysfunctions, depending on the location and severity of the injury (3). Debilitating chronic neuropathic pain (NP) tends to affect 40% of patients following SCI (4–6) and represents a highly disabling clinical condition (7). As a result, treatment of post-SCI NP is vital to mitigate the impact on body function and overall quality of life.

Anticonvulsants, antidepressants, analgesics, and cannabinoids have all been used to treat NP (8–11), but refractory pain following SCI is common (12). Despite the modest short-term benefits of drug therapy, the balance between long-term benefits and damage has been overlooked. Clinical decision-makers and pharmaceutical enterprises have interests in the suitability of drugs for pain relief, making a comparison of drug efficacy and safety valuable. Most studies to date have been limited by manpower and material resources and have thus compared individual drugs with placebo or, at most, compared two drugs. Little valid data can be derived from direct comparisons, stimulating a longstanding debate about drug efficacy and safety (13).

Network meta-analysis extends pairwise meta-analyses to enable the pooling of data from many clinical trials, comparing at least two treatments. Inferences regarding the relative efficacy of individual drugs with placebo or, at most, compared two drugs. Little valid data can be derived from direct comparisons, stimulating a longstanding debate about drug efficacy and safety (13).

Network meta-analysis extends pairwise meta-analyses to enable the pooling of data from many clinical trials, comparing at least two treatments. Using Bayes’ theorem of existing datasets provides a framework for comprehensive evaluation of drug efficacy and safety (16). Twenty RCTs were included in the present study, and Bayesian network analysis was performed to compare the efficacy and safety of different drugs for the treatment of NP in adults following SCI. The objective was to perform an evidence-based analysis for the benefit of clinical practitioners.

METHODS

Data Sources and Study Strategy

A literature search of PubMed, Cochrane Library, Embase, and Web of Science from inception to February 21, 2021 was conducted without language restrictions. The subject words “spinal cord injury,” “neuropathic pain,” “treatment,” and “randomized controlled trials” were used. Subject words were combined with related free words (from Mesh or Emtree). The search strategies used in PubMed and Embase are shown in Supplementary Data 1, 2.

Study inclusion criteria were as follows: (1) single or double-blinded [non-blinded assessors of subjective measurement scale outcomes in RCTs generate substantially biased effect sizes (17)] randomized controlled trials (RCTs) that compared drugs with a placebo or a second drug; (2) patients were adults (≥ 18 years old) with SCI (rated from A to D according to the American Spinal Injury Association (ASIA) impairment scale; (3) patients had been diagnosed with NP according to the criteria of the International Association for the Study of Pain guidelines: (a) the onset of pain within 1 year following SCI; (b) no primary relation of pain to movement, inflammation or other local tissue damage; (c) use of adjectives ‘hot-burning,” “tingling,” “pricking,” “pins and needles,” “sharp,” “shooting,” “squeezing,” “cold,” “electric” or “shock like” to describe the quality of pain; (d) persistent pain for at least 1 month; (e) assessment of pain intensity using numeric rating scale (NRS ≥ 4) or visual analog scale (VAS ≥ 40; 0–100 mm) where 0 = no pain and 10 or 100 = unbearable pain, respectively. Exclusion criteria were as follows: studies including non-drug therapy and non-RCTs; abstracts; studies with incomplete data, redundancy, or insufficient raw data; case reports; reviews; and meta-analyses.

Two researchers conducted independent screenings of titles and abstracts of the following retrieval. Thereafter, both researchers read the full texts, extracted data, and engaged in the discussion to arrive at a joint decision on validity for inclusion. A formal file extraction form was developed. The following data were extracted: author, subjects, sample size, average age, sex, intervention and control measures, follow-up time, intervention evaluation tools, and study design.

Selection Criteria and Study Design

Primary outcomes were efficacy (a standardized pain score at 4 weeks) and safety (adverse events). Secondary outcomes included efficacy (a standardized pain score at 8 weeks), standardized mental or sleep-related assessment scores, and incidence of serious adverse events as supplemental safety outcomes (Supplementary Tables 1–5). Pain, mental, and sleep-related assessment differences were defined as the score of intensity changes (rating scales shown in Table 1) from the baseline to the end point. Patients’ assessments for overall treatment were included among the secondary outcomes with lower scores on the specified rating scale, signifying better overall assessments. Some studies used more than one rating scale, necessitating standardization of continuous data. Results were recorded as close as possible to the 4-week follow-up time point for all analyses. If data were unavailable at the 4-week time point, data from between 3 and 17 weeks were used (time points close to 4 weeks were prioritized for primary outcomes and over 8 weeks for secondary outcomes).

Study Quality Control

Cochrane Handbook quality evaluation criteria were used by two researchers for independent evaluations of the publications. Criteria were as follows: random sequence generation, allocation hiding, participant and researcher blinding, result in evaluator blinding, outcome indicator integrity, selective reporting, and other sources of bias. Each item was scored as low, unclear, or high risk of bias.

Statistical Analysis

Bayesian framework network meta-analysis was performed using OpenBUGS [version 3.2.3; (38)] and R [version 3.6.2; (39)] for repetitive proof. Standardized mean differences (SMDs) for continuous outcomes and summary odds ratios (ORs) for
| Ref. no. | Author & year of publication | Country | Drugs intervention (I) & comparison (C) | Of SCI NP participants enrolled (I/C) | Sex (male/female) | Grade of ASIA | Age (Year) [Mean (SD) or (Median [IQR / Range]) (I/C)] | NP duration (Month) [Mean (SD) or (Median [IQR/Range]) (I/C)] | Follow-up time (Weeks) | Pain evaluation tools | Overall assessment of risk of bias | Evaluation tools of mental or sleep-related symptom relief |
|---------|-------------------------------|---------|----------------------------------------|----------------------------------------|------------------|--------------|-------------------------------------------------------|-----------------------------------------------------------|-------------------------|------------------------|--------------------------|----------------------------------|
| (18)    | Nct 2012                      | UK      | cannabinoids & placebo                 | 116 (56 / 60)                          | 91 / 25          | NA           | 48.7 (12.97) / 47.6 (12.69)                           | NA                                                                            | 7                       | NRS                    | high                     | NRS                              |
| (19)    | Cardenas 2013                 | USA     | pregabalin & placebo                   | 219 (111 / 108)                        | 176 / 43         | NA           | 46.1 (12.7) / 45.6 (13.8)                             | ≥3                                                                          | 16                      | NRS                    | high                     | MOS-SS                           |
| (20)    | Agarwal 2017                  | India   | amitriptyline & lamotrigine             | 147 (74 / 73)                          | 136 / 11         | A-D          | 29.6 (18–40)                                          | NA                                                                            | 3                       | SFMPQ2                 | high                     | NA                               |
| (21)    | Amr 2010                      | Egypt   | ketamine & gabapentin                  | 40 (20 / 20)                           | 33 / 7           | A-D          | 48.6 (10.1) / 48.7 (9.7)                             | 8 (6–17) / 9 (7–18)                                                     | 4                       | VAS (100)               | moderate                 | NA                               |
| (22)    | Amr 2011                      | Egypt   | ketamine & gabapentin                  | 40 (20 / 20)                           | 33 / 7           | A-D          | 48.6 (10.1) / 48.7 (9.7)                             | 8 (6–17) / 9 (7–18)                                                     | 8                       | VAS (100)               | moderate                 | NA                               |
| (23)    | Andresen 2016                 | Denmark | cannabinoids & placebo                 | 73 (36 / 37)                           | 54 / 19          | A-D          | 58.6 (11.3) / 54.1 (11.7)                            | ≥3                                                                         | 12                      | NRS                    | moderate                 | NA                               |
| (24)    | Salinas 2012                  | Colombia| carbamazepine & placebo                | 48 (24 / 22)                           | 42 / 4           | A-D          | 45.6 (18–70)                                          | NA                                                                        | 24                      | VAS (100)               | high                     | SF-36 Scale                      |
| (25)    | Siddall 2006                  | Australia| pregabalin & placebo                   | 137 (70 / 67)                          | 114 / 23         | A-D          | 50.3 (23–78) / 49.8 (21–80)                         | 24.5 (88.8) / 42.5 (88.8)                                          | 12                      | NRS                    | high                     | MOS-sleep scale                |
| (26)    | Tai 2016                      | USA     | gabapentin & placebo                   | 14 (7 / 7)                             | 12 / 2           | A-D          | 35.9 (8.96) / 35.9 (8.96)                            | 42.5 (88.8) / 42.5 (88.8)                                          | 10                      | NPS                    | moderate                 | NA                               |
| (27)    | Vranken 2008                  | Netherlands| pregabalin & placebo                  | 40 (20 / 20)                           | 21 / 19          | A-D          | 54.2 (9.4) / 54.7 (9.7)                             | ≥6                                                                         | 4                       | VAS (10)                | moderate                 | EQ-5D PDI                        |
| (28)    | Vranken 2011                  | Netherlands| duloxetine & placebo                  | 36 (18 / 18)                           | NA               | A-D          | NA                                                    | NA                                                                       | 8                       | VAS (10)                | high                     | EQ-5D PDI                        |
| (29)    | Yilmaz 2015                   | Turkey  | pregabalin & gabapentin                | 30 (15 / 15)                           | 25 / 5           | A-D          | 32.93 (11.87)                                        | 31.48 (11.08)                                                         | 18                      | VAS (10)                | moderate                 | PDI                              |
| (30)    | Chun 2019                     | USA     | BTX-A & placebo                        | 8 (5 / 3)                              | 6 / 2            | A            | 45 (32–61)                                            | ≥1                                                                       | 12                      | NPRS                   | moderate                 | Pain of sleep                    |
| (31)    | Finnerup 2009                 | Denmark | levetiracetam & placebo                | 36 (18 / 18)                           | 29 / 7           | A-D          | 51 (11.2)                                             | ≥3                                                                       | 5                       | NRS                    | high                     | Sleep interference             |
| (32)    | Han 2016                      | Korea   | BTX-A & placebo                        | 40 (20 / 20)                           | 29 / 11          | A-D          | 53.1 (9.1) / 48.9 (14.2)                             | ≥3                                                                       | 8                       | VAS (100)               | high                     | SF-MPQ scores                   |
| (33)    | Kaydok 2014                   | Turkey  | gabapentin & pregabalin                | 28 (14 / 14)                           | 21 / 7           | A-D          | 42.8 (12.3)                                           | 29.3 (25.8)                                                          | 4                       | VAS (100)               | moderate                 | NPS                              |
| (34)    | Levendoglu 2004               | Turkey  | gabapentin & placebo                   | 20 (10 / 10)                           | 13 / 7           | NA           | 35.9 (9.8)                                            | 15.8 (9.0)                                                         | 8                       | VAS (100)               | moderate                 | NPS                              |
| (35)    | Norrbrink 2009                | Sweden  | tramadol & placebo                     | 35 (23 / 12)                           | 28 / 7           | NA           | 51.3 (10.8)                                           | ≥6                                                                       | 4                       | MPI scales              | high                     | HAD                              |
| (36)    | Rintala 2007                  | USA     | amitriptyline & gabapentin & placebo  | 79 (28 / 26 / 25)                      | NA               | A-D          | 42.6 (12.6)                                           | ≥6                                                                       | 8                       | VAS / NRS               | moderate                 | NA                               |
| (37)    | Rintala 2010                  | USA     | cannabinoids & placebo                 | 14 (7 / 7)                             | 50 / 8            | NA           | 50.8 (3.8)                                            | ≥6                                                                       | 4                       | NRS                    | moderate                 | NA                               |

SCI, spinal cord injury; NP, neuropathic pain; ASIA, American Spinal Injury Association; SD, standard deviation; IQR, inter-quartile range; VAS, visual analog scale; NRS, numerical rating scale; SFMPQ2, short-form McGill pain questionnaire-2; NPS, numerical pain scale; NPRS, numerical pain rating scale; MPI, multidimensional pain inventory; MOS-SS, medical outcomes study sleep scale; SF-36 Scale, the short-form 36 item health survey questionnaire; EQ-5D, EuroQol five dimensions questionnaire; PDI, pain disability index; HAD, hospital anxiety, and depression.
dichotomous outcomes were estimated using both pairwise and network meta-analysis and 95% credible interval (CrI) calculated. Group-level data were used for network meta-analysis with the binomial likelihood for dichotomous outcomes and normal likelihood for continuous outcomes. Study effect sizes were synthesized using a random-effects network meta-analysis model.

Three Markov chains were used to set the initial value, and the number of iterations for the first model update was set to 50,000 and for the continuous update to 100,000. The first 50,000 annealing times were discarded to eliminate the effect of initial values and sampling started from 50,001 times. The data fitting effect is shown in Supplementary Figure 1. Primary outcome pain scores and adverse events were used as outcome indicators to draw an evidence network map. In the case of a closed-loop, inconsistencies between direct and indirect comparisons were assessed through node splitting, and a $p < 0.05$ demonstrated that the inconsistency did not occur by chance (40). The inconsistency factor (IF) is close to zero with the 95% CrI included of zero, indicating good consistency between direct and indirect evidence (41). Surfaces under the cumulative ranking curve (SUCRA) and mean ranks were used to rank treatments with respect to each outcome (42). The area under the cumulative ranking probability map allowed assessment of each intervention as the potential best treatment. Higher SUCRA scores demonstrated better efficacy or safety. Statistical evaluation of inconsistent network graphs and results figures were processed using the network and network graphs packages of Stata (MP 14.2).

The current study was registered in the International Platform of Registered Systematic Review and Meta-Analysis Protocols (INPLASY), No.: INPLASY202070061, doi: 10.37766/inplasy2020.7.0061. The original protocol is presented in the Supplementary Protocol.

RESULTS

Results were reported according to the preferred reporting items for systematic reviews and meta-analyses (PRISMA) extension statement (Supplementary Material) for network meta-analyses (43). A total of 1,133 citations were identified with 201 potentially eligible full texts. A flow chart outlining the selection process is shown in Figure 1. Between 1963 and 2021, there were 20 RCTs (1,198 patients), comparing 11 drugs or placebos [Table 1; (18–37)]. The mean number of participants was 60 ± 55. A total of 622 participants were randomly administered drugs, whereas 576 received placebos. Most patients suffered from moderate to severe pain. Drug-based (pain relief medication) and non-drug therapy (physical and psychological educational therapies) were included. Nineteen out of 20 trials (95%) were double-arm studies, and the remaining one (5%) was a three-arm study. Fourteen (70%) compared drugs with placebo and 5 (25%) compared two different drugs. Network estimates of the main outcomes were based on moderate to high certainty of evidence (Supplementary Figures 2, 3). Network analysis of primary and secondary outcomes conformed with the consistency model (Supplementary Figures 4, 5). All trials assessed during the current study involved the following 11 drugs: BTX-A, ketamine, amitriptyline, lamotrigine, pregabalin, duloxetine, gabapentin, tramadol, levetiracetam, carbamazepine, and cannabinoids. A network of eligible comparisons is presented for primary outcomes in Figure 2 and for secondary outcomes in Figure 3. The circle area represents the number of studies included in each treatment group, and the line width represents the number of comparative trials. All studies were RCTs, assessing drug action vs. a placebo, except for those involving ketamine and lamotrigine.

Pairwise efficacy and network meta-analysis of 11 drugs with respect to placebos from 20 RCTs are visualized as the forest plot shown in Figure 4. Pairwise and network meta-analysis of other outcomes are presented in Supplementary Figure 6. Inconsistent test node split analysis for primary and secondary outcomes was also performed, and a $p < 0.05$ or DIC (inconsistency) minus DIC (consistency) of $< 5$ was taken to indicate significant inconsistency. Results gave $p$-values of over 0.05 (Figure 4 and Supplementary Figure 6), indicating consistency between primary and secondary outcomes, and direct and indirect results. Detailed results of a pair of meta-analyses of primary and secondary outcomes of 11 drugs are presented in Figures 5, 6, respectively.

BTX-A, ketamine, amitriptyline, lamotrigine, pregabalin, and gabapentin were shown to be the most effective of the drugs tested for the primary outcomes of efficacy and safety (Figure 5). No significant differences were found among the 11 drugs in terms of adverse event rates. BTX-A was ranked as the most effective drug for pain relief at 4 weeks of follow-up. SMDs for BTX-A with respect to other drugs (gabapentin, tramadol, levetiracetam, carbamazepine, and cannabinoids) ranged from $-0.76$ to $-0.24$). Ketamine was also among the more effective drugs with SMDs compared with others (gabapentin, levetiracetam, carbamazepine, and cannabinoids), ranging from $-0.44$ to $-1.12$. No significant differences were found among lamotrigine, amitriptyline, and gabapentin, but their efficacy was higher than that of other drugs (carbamazepine and cannabinoids) with SMDs, ranging from $-0.88$ to $-2.81$. However, tramadol, levetiracetam, and cannabinoid did not produce significantly different outcomes when compared with one another or with a placebo. Lamotrigine had the best safety profile with respect to adverse events of the primary outcome when compared to pregabalin 0.02 (0.001–0.82), duloxetine 0.01 (0.001–0.32), and tramadol 0.02 (0.001–0.82).

Pain relief at 8-week follow-up and relief of mental or sleep-related symptoms represented the secondary outcome of efficacy, and network analysis results were in agreement with those for the primary outcome. BTX-A, ketamine, amitriptyline, gabapentin, and pregabalin showed greater long-term efficacy of pain relief (SMDs: $-0.90$ to $-1.20$; Figure 6A). Gabapentin, BTX-A, and pregabalin were more successful in relieving mental or sleep-related symptoms with SMDs, ranging from $-0.63$ to $-0.86$ (Figure 6B). The incidence of serious adverse events represented the secondary outcome for the assessment of safety (Figure 6C). No significant differences were found for any drug, with the exception of tramadol, which triggered more serious adverse events than any other (ORs: 0.09–0.11).
Rankings for all 11 drugs were based on SUCRA values for primary and secondary outcomes. Combined with network meta-analysis results on the efficacy of pain relief at 4-week follow-up (Supplementary Figure 6), BTX-A was ranked first, followed by ketamine, amitriptyline, lamotrigine, pregabalin, duloxetine, gabapentin, tramadol, levetiracetam, carbamazepine, and cannabinoids. SUCRA safety rankings based on adverse event incidence (Supplementary Figure 7) were as follows: lamotrigine, BTX-A, levetiracetam, amitriptyline, gabapentin, ketamine, carbamazepine, cannabinoids, pregabalin, tramadol, and duloxetine. SUCRA rankings of secondary outcomes were very similar to those for primary outcomes, except gabapentin was the most effective in mental or sleep-related symptom relief and tramadol had the worst safety profile (Supplementary Figures 8–11). A net funnel for primary outcomes is shown in Supplementary Figures 12, 13 and indicates that publication bias was under control. A heat map depicting the hierarchy of all 11 drugs according to mean SUCRA values across 5 primary and secondary outcomes is shown in Figure 7. In summary, the overall rankings are BTX-A, gabapentin, ketamine, amitriptyline, pregabalin, duloxetine, lamotrigine, levetiracetam, tramadol, carbamazepine, and cannabinoids. However, due to the paucity of data, quantitative synthesis was not performed on the remaining *priori*-defined outcomes.

**DISCUSSION**

Davis and Martin (44) have vividly delineated SCI-related NP: hot, burning pain would be replaced by severe crushing pressure, vise-like, pinching sensations, streams of fire running down the legs into the feet and out of the toes or a pain produced by the pressure of a knife being buried in the tissue, twisted around rapidly and simultaneously withdrawn. In stark contrast to the profound effectiveness of opioids in relieving nociceptive pain, there is, unfortunately, no similar panacea for the treatment of NP (45). To advance the evidence-based approach to drug management of SCI-related NP, the current study conducted a Bayesian network analysis and managed to make a discussion per drug based on the results of network analysis and etiology and/or pharmacology of NP as follows.

**BTX-A**

BTX-A is a complex proteinaceous neurotoxin with proteolytic activity, which affects both the synaptic and auxiliary proteins involved in the release of vesicle neurotransmitters (eg,
FIGURE 2 | Network meta-analysis of eligible comparisons for primary outcomes. (A) Pain relief for around 4 weeks, n = 1,207. (B) Any adverse events, n = 1,232.

Width of the lines is proportional to the number of trials, comparing every pair of treatments (numbers on the lines). Size of every circle is proportional to the number of randomly assigned participants (sample size). BTX-A, botulinum toxin-A; Ket, ketamine; Ami, amitriptyline; Lam, lamotrigine; Pre, pregabalin; Dul, duloxetine; Gab, gabapentin; Tra, tramadol; Lev, levetiracetam; Car, carbamazepine; Can, cannabinoids; Pla, placebo.
acetylcholine (Ach) and/or other local neuropeptides [e.g., calcitonin gene-related peptides (CGRP) and substance P] (46). However, BTX-A’s dual mechanisms of action are controversial (47). Favre-Guilmard et al. (48) and Park and Chung (49) have demonstrated that BTX-A inhibits the secretion of substance P and CGRP from dorsal root ganglion (DRG), restrains the expression of TRPV1 and P2X3 and induces a central effect through retrograde axonal transport. Those reports particularly emphasize BTX-A’s potential character in reducing NP. Although Finnerup et al. (50) have recommended BTX-A as a fourth-line treatment, there is still increasing interest in its use for the management of NP (47, 51, 52). Therefore, oral or injected administration of BTX-A has been shown to induce analgesia in neuro-related or neurogenic pain (53–55). Similarly, BTX-A (200 U) was administered subcutaneously during two RCTs (32, 56) referred to in the current study and alleviated intractable post-SCI NP as the previous report (57). More importantly, guidelines published by the American Academy of Neurology recommend the use of BTX-A, illustrating its effectiveness (Level A) in SCI-induced NP (58). The current study provides further evidence to support the efficacy of BTX-A via network meta-analysis within the Bayesian evidence framework. However, optimal routes of BTX-A administration and dosage remain unclear. The current findings may further knowledge of pain mechanisms and give insights into the potential for novel-engineered toxins that specifically target pain neurotransmitters.

**Ketamine**

The α2δ-1-bound NMDA receptors, in particular, are involved in the pathology of NP and present a potential target for NP therapy (59–61). Luo et al. (62) reported that repeated activation of presynaptic fibers leads to long-term potentiation (LTP), which represents a vital mechanism in producing central pain. In combination with activation of NMDA receptors, such algogenic mechanisms probably relate to reinforcing central sensitization (45, 63). Coincidentally, ketamine could antagonize NMDA receptor-mediated central sensitization by blocking harm signal inputs (64) and had been considered safe and effective in reducing chronic pain (65–67). Also, Amr (21, 22) studied the efficacy of long-term intravenous drip and epidural injection of ketamine for post-SCI NP. Despite the safety and effectiveness of the drug, its analgesic effects wore off 2 weeks after discontinuation of the infusion, although the epidural route produced more prolonged efficacy (1 month). The current study found ketamine suitable for the management of post-SCI NP, as it was effective with few adverse events (Figure 5 and Supplementary Figures 8, 11). Nevertheless, the necessity of administration via intravenous drip or epidural injection reduces its convenience compared with other drugs.

**Gabapentinoids**

Gabapentin and pregabalin are gabapentinoids that target the α2δ-1 auxiliary subunit of voltage-gated Ca2+ channels. Although structurally similar to GABA, gabapentin does not affect the uptake, synthesis, or metabolism of GABA (68) nor do either gabapentin or pregabalin bind to GABAA or GABAB receptors (69–71). Hence, gabapentin is commonly recommended as immediate therapy for the management of post-SCI NP (72, 73). Besides, pregabalin as a new generation of gabapentinoid is the only drug approved by the US Food and Drug Administration (FDA) for the treatment of post-SCI NP. Moreover, gabapentin and pregabalin were extensively compared in the systematic review and meta-analysis by Davari et al. (74). No significant difference between the two drugs was reported in terms of change of a pain score and safety. The current network meta-analysis produced findings consistent with those of previous studies (Figure 5). Although pregabalin resulted in more adverse events (Supplementary Figure 8), these tended to be minor and well tolerated. More importantly, gabapentin

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**FIGURE 3** | Network meta-analysis of eligible comparisons for secondary outcomes. (A) Pain relief more than 8 weeks, n = 615. (B) Mental or sleep-related symptom relief, n = 825. (C) Serious adverse events, n = 1,232. Width of the lines is proportional to the number of trials, comparing every pair of treatments for secondary outcomes (numbers on the lines). Size of every circle is proportional to the number of randomly assigned participants (sample size). BTX-A, botulinum toxin-A; Ket, ketamine; Ami, amitriptyline; Lam, lamotrigine; Pre, pregabalin; Dul, duloxetine; Gab, gabapentin; Tra, tramadol; Lev, levetiracetam; Car, carbamazepine; Can, cannabinoids; Pla, placebo.

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FIGURE 4 | Efficacy pairwise and network meta-analysis of 11 drugs and placebo from 20 RCTs. A forest plot (A) shows pairwise and network meta-analysis of different drugs on the target of neuropathic pain relief after SCI as compared with placebo. And the forest plot (B) shows pairwise and network meta-analysis among head-to-head drugs. The blue line in forest plots indicates the outcome analysis of corresponding studies, the green line refers to pooled within designed treatments, the red line means pooled overall. The test of consistency reveals all studies are according to the consistency model. BTX-A, botulinum toxin-A; Ket, ketamine; Ami, amitriptyline; Lam, lamotrigine; Pre, pregabalin; Dul, duloxetine; Gab, gabapentin; Tra, tramadol; Lev, levetiracetam; Car, carbamazepine; Can, cannabinoids; Pla, placebo.
proved to be particularly valuable for its relief of mental or sleep-related symptoms (Figure 6). The current findings reinforce the view that gabapentinoids are suitable first-line treatment options for NP. Novel information about their mechanism of action may lead to improved treatment management, and, thus, in combination with the previous findings (45, 75), we also support the perspective that combining gabapentinoids with other active drugs might be expected to improve overall therapeutic effectiveness in post-SCI NP.

Serotonin and Noradrenaline

Amitriptyline and duloxetine target serotonin or 5-hydroxytryptamine (5-HT) and noradrenaline (NA) uptake systems to have an impact on the nociceptive regulatory circuit (76). Amitriptyline is a tricyclic antidepressant with analgesic action in neuropathy and deafferentation-induced autotomy (77). Amitriptyline is associated with fewer adverse events and should be the first-choice drug for NP. This proposes lamotrigine as a favorable drug for NP treatment as in previous reports (50, 75). Similarly, carbamazepine presents an FDA-approved drug for epilepsy, bipolar disorder, and NP treatment (81, 82). A double-blind, placebo-controlled RCT plus parallel-group study conducted by Salinas et al. (24) found that early carbamazepine did not reduce overall NP incidence or intensity in the long run. Besides, many studies have indicated the suitability of the anticonvulsant carbamazepine.

Voltage-Gated Na⁺ Channels

Lamotrigine and carbamazepine act on voltage-sensitive sodium channels, stabilize neuronal membranes, and inhibit the pathological release of excitatory amino acid transmitters, such as glutamate, that are mediated by sodium influx. Agarwal and Joshi (20) found that lamotrigine is associated with fewer adverse events and should be the first-choice drug for NP. This proposes lamotrigine as a favorable drug for NP treatment as in previous reports (50, 75). Similarly, carbamazepine presents an FDA-approved drug for epilepsy, bipolar disorder, and NP treatment (81, 82). A double-blind, placebo-controlled RCT plus parallel-group study conducted by Salinas et al. (24) found that early carbamazepine did not reduce overall NP incidence or intensity in the long run. Besides, many studies have indicated the suitability of the anticonvulsant carbamazepine.

FIGURE 5 | Head-to-head comparisons for efficacy and safety (primary outcomes) of 11 drugs. Drugs are reported in efficacy order sorted from top left to bottom right. Data are SMD (95% CrI) and OR (95% CrI) in the column-defining treatment compared with the row-defining treatment. For efficacy (pain relief for around 4 weeks), SMD lower than 0 favors the column-defining treatment. For safety (adverse events), OR lower than 1 favors the top left one. To obtain OR for comparisons in the opposite direction, reciprocals should be taken. Significant results are in bold. SMD, standard mean deviation; OR, odds ratio; CrI, credible interval; BTXA, botulinum toxin-A.
### FIGURE 6
Head-to-head comparisons for secondary outcomes of drugs and placebo.

#### (A) Pain relief more than 8 weeks (8 drugs).

- **Pain relief more than 8 weeks (8 drugs).**
  - **BTXA**
    - ketamine: $-0.02$ (95% CrI: $-0.09, 0.85$)
    - gabapentin: $-0.35$ (95% CrI: $-0.99, 0.28$)
    - pregabalin: $-0.36$ (95% CrI: $-0.93, 0.27$)
  - **amitriptyline**
    - $-0.07$ (95% CrI: $-0.77, 0.63$)
  - **duloxetine**
    - $-0.40$ (-1.22, 0.42)
  - **carbamazepine**
    - $-1.01$ (-1.53, -0.48)
  - **placebo**
    - $-1.20$ (-1.90, -0.50)

#### (B) Mental or sleep-related symptom relief (8 drugs).

- **Mental or sleep-related symptom relief (8 drugs).**
  - **BTXA**
    - gabapentin: $-0.12$ (95% CrI: $-0.77, 0.53$)
    - pregabalin: $-0.44$ (-1.18, 0.30)
    - duloxetine: $-0.48$ (-1.26, 0.29)
  - **amitriptyline**
    - $-0.36$ (-1.14, 0.51)
  - **carbamazepine**
    - $-1.63$ (-1.16, -0.10)
  - **placebo**
    - $-0.86$ (-1.55, -0.16)

#### (C) Serious adverse events (11 drugs).

- **Serious adverse events (11 drugs).**
  - **BTXA**
    - carbinoids: $0.91$ (0.21, 1.59)
    - gabapentin: $0.91$ (0.32, 1.50)
    - pregabalin: $0.93$ (0.34, 1.53)
  - **amitriptyline**
    - $1.00$ (0.06, 2.12)
  - **carbamazepine**
    - $0.96$ (0.29, 1.63)
  - **placebo**
    - $0.98$ (0.21, 1.59)

Data are SMDs (95% CrI) and ORs (95% CrI) in the column-defining treatment compared with the row-defining treatment. For A and B, SMDs lower than 0 favor the column-defining treatment. For C, ORs lower than 1 favor the top left one. Significant results are in bold. SMD, standard mean deviation; OR, odds ratio; CrI, credible interval; BTXA, botulinum toxin-A.
Cannabinoids
Cannabinoids activate the CB1 and/or CB2 receptors, but any potential role in the treatment of SCI-related NP remains unclear (18). Cannabinoids produce analgesic effects due to actions in and around the brain and spinal cord, but use is limited by side effects and concerns over long-term risks (50, 87, 88). The current network meta-analysis suggested little effect of cannabinoids compared with a placebo, producing a SUCRA ranking in the last place. Thus, cannabinoids may not be appropriate for the treatment of post-SCI NP, mainly due to potential abuse, diversion, and long-term mental health risks, especially in susceptible populations (45, 50).

Tramadol and Levetiracetam
Tramadol and levetiracetam, an opioid analgesic and anticonvulsant, respectively, were generally reported to be minimally effective in the treatment of SCI-related NP (31, 35). On the one hand, tramadol produced more adverse effects than many other drugs, in agreement with the findings of Norrbrink and Lundeborg (35) and Finnerup et al. (86). These authors stressed the adjuvant role of tramadol and other opioids, indicating that gabapentin, pregabalin, serotonin, and norepinephrine reuptake inhibitors should first be tried for NP. Besides, numerous studies have recommended opioid titration with low initial and slow individual doses to mitigate adverse events. On the other hand, Finnerup et al. (31) indicated that levetiracetam in doses titrated up to 3,000 mg had no analgesic or other benefits in patients with post-SCI NP during a double-blind, placebo-controlled, crossover, multicenter RCT. The reason for the lack of efficacy of levetiracetam is unknown, but doses used may be insufficient to affect SV2A and have an impact on NP. Lastly, the current findings are to contra-indicate the use of tramadol and levetiracetam for SCI-related NP due to poor efficacy and high risk of adverse events (Figures 5–7).

**Recommendation and Insight**
To the best of our knowledge, the current comprehensive network meta-analysis is the first to address the pharmacological treatment of NP after SCI. Our findings are different from previous meta-analyses, which were limited to a single drug, a pair of drugs, or a class of drugs. Bayesian inference provides a statistical framework for the integration of current data, prior knowledge, and reasonable assumptions about the system to generate probability distributions (89). Hence, a Bayesian framework was used to implement the network meta-analysis and summarize the two primary and three secondary outcomes of drug efficacy and safety. In summary, gabapentin, BTX-A, amitriptyline, ketamine, and lamotrigine had relatively high pain relief efficacy with fewer adverse events than other drugs, which support them as first-line therapy for post-SCI NP. Pregabalin and duloxetine had some degree of efficacy in NP relief,
and they are proposed as second-line treatment because their safety should be carefully considered. Tramadol, levetiracetam, carbamazepine, and cannabinoids had lower efficacy and worse safety profiles than other drugs, making them less favorable in treating post-SCI NP. Although all drugs were accompanied by adverse effects to varying degrees, most of them were acceptable and well tolerated. Undoubtedly, head-to-head studies were prioritized, and the certainty of the retrieved evidence was confirmed within this finding. Totally, few differences were found among the scrutinized drugs, and differences in efficacy and safety were minimal. Anyway, the patient's individual perception of the disease and its symptoms proximately results in variable therapeutic efficacy and acceptability of side effects. Therefore, further multicenter, large sample RCTs, mechanistic and data-mining studies for each drug are required to extend evaluations and reduce the bias of significant innovation.

STUDY LIMITATIONS

Meta-analyses have some limitations due to the reliance on statistical assumptions and data-based simulation, and the rigor of the research methodology of individual studies is a potential confounding factor. We acknowledge some limitations to the present study. Firstly, some RCTs included in the current analysis had fewer than 50 participants. Secondly, SCI-related NP was generalized and not divided into subgroups (NP at the injured level and below the injured level). Thirdly, only drug efficacy and safety were compared without taking into account dosage and frequency or availability and cost – neither was the drug delivery route scrutinized. Fourthly, some studies included in the present meta-analysis were funded by drug companies, which may carry a risk of bias.

CONCLUSION

The management of post-SCI NP was analyzed within the framework of Bayesian network analysis to demonstrate that the efficacy and safety of BTX-A make it a suitable drug choice. Ketamine is also beneficial but has to be administered via intravenous drip or epidural injection. Gabapentin and amitriptyline remain appropriate first-line treatments for NP. Lamotrigine, pregabalin, and duloxetine have some utility as well. It is noteworthy that lamotrigine and gabapentin have fewer side effects and help relieve mental or sleep-related symptoms. Tramadol, levetiracetam, carbamazepine, and cannabinoids have poor safety or efficacy and are not recommended for SCI-related NP treatment, unless as adjuvants.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding authors.

AUTHOR CONTRIBUTIONS

H-QL and Z-HC conceptualized and designed the study, analyzed the data, and drafted and revised the manuscript. LH and FF designed the data collection instruments, mechanistic and data-mining studies for each drug are required to extend evaluations and reduce the bias of significant innovation.

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