Comparative Impact of Pharmacological Therapies on Cluster Headache Management: A Systematic Review and Network Meta-Analysis

Jae-Hee Kwon †, Ja-Young Han †, Ji-Woong Choi, Hye-Rim Park and Heeyoung Lee *

Abstract: It is important to find effective and safe pharmacological options for managing cluster headache (CH) because there is limited evidence from studies supporting the general efficacy and safety of pharmacological therapies. This systematic review and network meta-analysis (NMA) analyzed published randomized controlled trials (RCTs) to evaluate the efficacy and safety of pharmacological treatments in patients with CH. The PubMed and Embase databases were searched to identify RCTs that evaluated the efficacy and safety of pharmacological treatments for CH. Efficacy outcomes included frequency and duration of attacks, pain-free rate, and the use of rescue agents. Safety outcomes were evaluated based on the number of patients who experienced adverse events. A total of 23 studies were included in the analysis. The frequency of attacks was reduced (mean difference (MD) = −1.05, 95% confidence interval (CI) = −1.62 to −0.47; \( p = 0.0004 \)), and the pain-free rate was increased (odds ratio (OR) = 3.89, 95% CI = 2.76–5.48; \( p < 0.00001 \)) in the pharmacological treatment group, with a lower frequency of rescue agent use than the placebo group. Preventive, acute, and triptan or non-triptan therapies did not show significant differences in efficacy (\( p > 0.05 \)). In the NMA, different results were shown among the interventions; for example, zolmitriptan 5 mg was more effective than zolmitriptan 10 mg in the pain-free outcome (OR = 0.40, 95% CI = 0.19–0.82; \( p < 0.05 \)). Pharmacological treatment was shown to be more effective than placebo to manage CH with differences among types of therapies and individual interventions, and it was consistently shown to be associated with the development of adverse events. Thus, individualized therapy approaches should be applied to treat CH in real-world practice.

Keywords: pharmacological therapy; cluster headaches; systematic review; network meta-analysis

1. Introduction

Cluster headache (CH) is a primary headache disorder characterized by intense headaches occurring on one side of the head and the development of cranial autonomic symptoms, including agitation, nasal congestion, and conjunctival injection [1]. If severe CH attacks are not treated, symptoms can persist for weeks to months and may even trigger suicidal ideation [2]. Although CH is rare, the significant symptoms caused by the disease have been a public health issue and a personal burden to many individuals [3]. Jensen et al. reported that >90% of CH patients experienced a negative impact on the quality of their lives, including occupational and social disabilities, during the cluster period [4].

However, the current understanding of the pathophysiological mechanisms of CH remains far from complete in terms of neurovascular and chronobiological aspects despite many studies investigating pathophysiological mechanisms for developing therapies to treat CH [1,5–7]. Due to considerable limitations in non-drug treatment [2], the American Headache Society guidelines have recommended several pharmacological treatments [1,5,6].
Nevertheless, more evidence is needed to support the efficacy of some medications for treating CH in clinical practice [1]. Relevantly, Remahl et al. reported that the rates of placebo response, such as cessation of headache attacks, were 7% to 43% in previous trials involving CH patients [8]; thus, more empirical studies are needed to demonstrate the efficacy and safety of pharmacological treatments compared to placebo.

Furthermore, although Francis et al. previously offered a systematic review of pharmacological options to treat CH patients, the authors could not conduct quantitative analysis given the limited number of studies included and indicated that not all medications could be recommended to treat CH patients due to insufficient evidence [9]. Moreover, several current systematic reviews and meta-analyses could not investigate a wide range of pharmacological options or provide statistically powered evidence supporting CH treatment due to the scarcity of studies enrolling CH patients [10,11]. As Brandt et al. [12] reported that for clinicians, challenges in determining the relative merit of various pharmacological alternatives for treating CH and choosing the best treatments still exist, which could be solved by network meta-analysis with the same approaches as the current study. Along with controversy and limited evidence to support pharmacological treatments on CH, for the many different types of pharmacological treatments for CH, including preventive, acute, or triptans, sufficient evidence quantitatively evaluating efficacy and safety based on pharmacological treatment types has not been accumulated for treating CH patients [13]. Because, along with advantages of systematic reviews, combining the direct and indirect could provide refined estimates [14,15], we conducted a systematic review and network meta-analysis to evaluate the efficacy and safety of pharmacological options followed by subgroup analysis determining best treatments for CH care.

2. Materials and Methods

This systematic review and network meta-analysis was performed in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines [16]. The protocol was registered on the International Prospective Register of Systematic Reviews database under no. CRD42022301178.

2.1. Data Sources and Search Strategy

A comprehensive strategy was used to search PubMed and Embase literature databases for relevant systematic studies addressing pharmacological treatment in patients with CH. A database search was performed to identify relevant articles published up to 8 January 2021, using CH-related keywords and medical subject headings (MeSH) terms. The reference lists from other relevant articles were manually searched to identify additional potentially eligible studies. Titles and abstracts were screened using the following search terms to identify relevant texts: “cluster headache”, “histamine cephalalgia”, “ciliary neuralgia”, “Horton syndrome”, “Sluder’s neuralgia”, “sphenopalatine neuralgia”, “migraine”, “neuralgia”, “cephalgia and headache” and “RCT”. Two investigators independently searched and evaluated the articles retrieved from the databases. Discrepancies between investigators were resolved by a third investigator.

2.2. Study Selection

Two independent investigators evaluated the titles and abstracts of the articles retrieved in the literature search to assess their eligibility and inclusion. Randomized controlled trials (RCTs) investigating CH treatment in patients who underwent pharmacological therapy were included. All included RCTs compared pharmacological treatment with placebo. Supplements, review articles, studies published in languages other than English, and those with only a single arm were excluded. Studies with a sample size less than five and those published only as an abstract were also excluded.
2.3. Data Extraction and Quality Assessment

Data extracted from the included articles were as follows: year of publication, study design, type(s) of medications, aim(s) of therapy, mean age, male/female proportion, route of administration, frequency of attacks, duration of attacks, pain-free rate, and number of individuals needing rescue agents and experiencing adverse events (AEs). The risk-of-bias assessment tool developed by Cochrane Collaboration was used to evaluate the quality of the RCTs [17]. The quality of evidence was evaluated as high, normal, low, or very low according to the Grading of Recommendations, Assessment, Development and Evaluation (GRADE) approach to assess the level of confidence in each effect estimation [18].

2.4. Data Synthesis and Analysis

The current study assessed the efficacy and safety of pharmacological treatment in patients with CH compared with placebo. Individuals who underwent pharmacological treatment comprised the “intervention group” and those treated with placebo comprised the “placebo group”. Efficacy outcomes included frequency of attacks, duration of attacks, number of patients using rescue agents, and post-treatment pain-free rate. The safety outcome was the number of patients who experienced AEs. In addition, subgroup analysis was used to assess efficacy and safety according to the aim of therapy (i.e., preventive and acute) and compared. Depending on the aim of the treatment types, preventive treatment [13] was used to reduce the frequency of CH attacks or to restore patients to headache-free status. Acute treatment is used to provide rapid relief [13]. Another analysis in terms of treatment types as a subgroup analysis was performed to evaluate differences in the efficacy and safety between drugs with or without triptan ingredients.

2.4.1. Statistical Analysis

The data used in the direct meta-analysis were analyzed using Review Manager (RevMan, Version 5.3, The Nordic Cochrane Centre, The Cochrane Collaboration: Copenhagen, Denmark, 2014) and Collaborative Meta-Analysis version 3 (Biostat Inc., Englewood, NJ, USA). Network meta-analysis was performed by either the fixed-effect or random-effect model, using the “netmeta” and “gemtc” package of R software (version 4.1.1).

2.4.2. Pairwise Meta-Analysis

The overall effect size was expressed as odds ratio (OR), and continuous outcomes were expressed as mean difference (MD), with corresponding 95% confidence intervals (CI) for comparative studies and each intervention. The I² statistic was used to evaluate heterogeneity among studies, and the percentile statistics were classified as low (<25%), medium (25–50%), or high (>50%). If the resulting analysis included >10 studies, a linear regression test of the funnel plots and Egger’s test were performed to assess publication bias.

2.4.3. Network Meta-Analysis

In the Bayesian framework, we performed Markov Cain Monte Carlo with 10,000 simulations in each of the 4 chains. The first 5000 simulations were considered to be the burn-in period. In each Markov chain Monte Carlo cycle, the probabilities of each treatment ranking from first to last were estimated by effect size. According to the sum of probabilities for each treatment ranking, cumulative probabilities were defined. Each treatment’s ranking was based on the calculated SUCRA (the surface under the cumulative ranking curve) values. The value of the SUCRA ranged from 0% to 100%. A higher SUCRA value represents better treatment.

2.4.4. Assessment of Consistency and Heterogeneity

The net-splitting method was used to evaluate the inconsistencies between direct and indirect evidence. Differences were considered statistically significant at \( p < 0.05 \). Meta-regression was used to examine the quantitative influence of study characteristics on
the effect size. The overall effect size was analyzed using the mean age and proportion of males at baseline included as covariates.

3. Results

3.1. Study Selection

A comprehensive search of the PubMed and Embase databases retrieved 457 potentially relevant articles. After full-text review, however, this figure was narrowed to 40 articles, the reference lists of which were manually searched and screened to ultimately include a total of 23 studies in the present analysis (Figure 1).

Figure 1. Flowchart of study identification and selection.

3.2. Study Description

The basic characteristics of the 23 studies [19–41] included in the current investigation are summarized in Table 1. A total of 1559 patients were included in the current study. Regarding route of drug administration, twelve studies [19,23–28,30,35,37,38,41] were oral, seven [20,21,32,33,36,39,40] were injected, and four [22,29,31,34] were nasal. Zolmitriptan 5 mg (ZOL5) and zolmitriptan 10 mg (ZOL10) and placebo (PLA) were evaluated
by various studies [27,34]. Galcanezumab (GAL) and PLA were evaluated by various studies [39,40]. Cimetidine (CIM) and PLA were evaluated by one study [19]. Sumatriptan 6 mg (SUM6) and PLA were evaluated by one study [20]. SUM6 and sumatriptan 12 mg (SUM12) and PLA were evaluated by one study [21]. Capsaicin (CAP) and PLA were evaluated by one study [22]. Sumatriptan 100 mg (SUM100) and PLA were evaluated by one study [23]. Melatonin (MEL) and PLA were evaluated by one study [24]. Lithium carbonate (LCAR) and PLA were evaluated by one study [25]. Misoprostol (MIS) and PLA were evaluated by one study [26]. Verapamil (VER) and PLA were evaluated by one study [28]. Civamide (CIV) and PLA were evaluated by one study [29]. Valproate (VAL) and PLA were evaluated by one study [30]. Sumatriptan nasal spray (SUMS) and PLA were evaluated by one study [31]. Octreotide (OCT) and PLA were evaluated by one study [32]. Betamethasone (BET) and PLA were evaluated by one study [33]. Frovatriptan (FRO) and PLA were evaluated by one study [35]. Cortivazol (COR) and PLA were evaluated by one study [36]. Warfarin (WAR) and PLA were evaluated by one study [37]. Candesartan cilexetil (CAN) and PLA were evaluated by one study [38]. Prednisone (PRE) and PLA were evaluated by one study [41]. Of the included studies, nine [19–21,26,27,31,32,34,37] were crossover designs, and fourteen [22–25,28–30,33,35,36,38–41] were parallel studies. Regarding the types of therapy involved, seven trials [20,21,23,27,31,34,35] investigated triptans to treat CH, and sixteen [19,22,24–26,28–30,32,33,36–41] investigated non-triptans. The therapy type consisted of thirteen preventive [19,23,24,28–30,33,35,37–41] and ten acute therapies [20–22,25–27,31,32,34,36]. The baseline characteristics of the studies, including age and male/female proportions, are summarized in Table 2.

### Table 1. Characteristics of the included studies.

| Study Name          | Publication Year | No. of Patients | Aims of Therapy | Types of Medications Intervention | Comparator | Routes of Administration | Study Design |
|---------------------|------------------|-----------------|-----------------|----------------------------------|------------|--------------------------|--------------|
| Russell et al. [19] | 1979             | 12              | prevention     | cimetidine, chlorpheniramine     | placebo    | PO                       | crossover    |
| Ekbom et al. [20]  | 1991             | 39              | acute          | sumatriptan                      | placebo    | SC                       | crossover    |
| Ekbom et al. [21]  | 1993             | 134             | acute          | sumatriptan                      | placebo    | SC                       | crossover    |
| Marks et al. [22]  | 1993             | 13              | acute          | capsaicin                        | placebo    | nasal cream              | parallel    |
| Monstad et al. [23] | 1995             | 168             | prevention     | sumatriptan                      | placebo    | PO                       | parallel    |
| Leone et al. [24]  | 1996             | 20              | prevention     | melatonin                        | placebo    | PO                       | parallel    |
| Steiner et al. [25] | 1997             | 27              | acute          | lithium carbonate               | placebo    | PO                       | parallel    |
| Evers et al. [26]  | 1998             | 8               | acute          | misoprostol                      | placebo    | PO                       | crossover    |
| Bahra et al. [27]  | 2000             | 124             | acute          | zolmitriptan                     | placebo    | PO                       | crossover    |
| Leone et al. [28]  | 2000             | 30              | prevention     | verapamil                        | placebo    | PO                       | parallel    |
| Saper et al. [29]  | 2002             | 28              | prevention     | civamide                         | placebo    | nasal                    | parallel    |
| Amrani et al. [30] | 2002             | 95              | prevention     | sodium valproate                | placebo    | PO                       | parallel    |
| Vliet et al. [31]  | 2003             | 118             | acute          | sumatriptan                      | placebo    | nasal                    | crossover    |
| Matharu et al. [32] | 2004            | 57              | acute          | octreotide                       | placebo    | SC                       | crossover    |
| Ambrosini et al. [33] | 2005          | 23              | prevention     | betamethasone                    | placebo    | SC                       | parallel    |
| Cittadini et al. [34] | 2006          | 92              | acute          | zolmitriptan                     | placebo    | nasal                    | crossover    |
| Pager et al. [35]  | 2011             | 10              | prevention     | frovatriptan                     | placebo    | PO                       | parallel    |
| Leroux et al. [36] | 2011             | 43              | acute          | cortivazol                       | placebo    | SC                       | parallel    |
| Hakim et al. [37]  | 2011             | 34              | prevention     | warfarin                         | placebo    | PO                       | crossover    |
| Tronvik et al. [38] | 2013            | 32              | prevention     | candesartan cilexetil            | placebo    | PO                       | parallel    |
| Goodson et al. [39] | 2019            | 106             | prevention     | galcanezumab                     | placebo    | SC                       | parallel    |
| Dodick et al. [40] | 2020             | 237             | prevention     | galcanezumab                     | placebo    | SC                       | parallel    |
| Obermann et al. [41] | 2021           | 109             | prevention     | prednisone                       | placebo    | PO                       | parallel    |

PO, Per Oral; Nasal, nasal spray; SC, Subcutaneous injection.
### Table 2. Baseline characteristics of age and male proportion.

| Study Name        | Publication Year | Mean Age (Year) | Male Proportion (%) |
|-------------------|------------------|----------------|---------------------|
|                   |                  | Intervention Group | Control Group       |
| Russell et al. [19]| 1979             | 49              | -                   |
| Ekbom et al. [20] | 1991             | 42 ± 10         | 79.5                |
| Ekbom et al. [21] | 1993             | 41 ± 9          | 86.6                |
| Marks et al. [22] | 1993             | -               | 23.1                |
| Monstad et al. [23]| 1995            | 40 ± 10         | 88.7                |
| Leone et al. [24] | 1996             | 38              | 75                  |
| Steiner et al. [25]| 1997            | 34.5 ± 19       | 100                 |
| Evers et al. [26] | 1998             | 44.5            | 100                 |
| Bahra et al. [27] | 2000             | 43.8 ± 10.9     | 86.3                |
| Leone et al. [28] | 2000             | 44 ± 8          | 90                  |
| Saper et al. [29] | 2002             | 45.1 ± 10.5     | 89.3                |
| Amrani et al. [30]| 2002             | 47 ± 11.3       | 88.4                |
| Vliet et al. [31] | 2003             | 43 ± 11         | 82.2                |
| Matharu et al. [32]| 2004            | 40 ± 10         | 78.9                |
| Ambrosini et al. [33]| 2005       | 42              | 86.9                |
| Cittadini et al. [34]| 2006           | 40 ± 10         | 86.9                |
| Pageler et al. [35]| 2011            | -               | -                   |
| Leroux et al. [36]| 2011             | CCH: 41.3 ± 13.3| CCH: 42.8 ± 11.9    | 88.4              |
|                   |                  | ECH: 40.0 ± 7.8 | ECH: 41.9 ± 10.4    |
| Hakim et al. [37] | 2011             | 44.1 ± 3.1      | 76.5                |
| Tronvik et al. [38]| 2013            | 42 ± 10.1       | 84.4                |
| Goadsby et al. [39]| 2019            | 47 ± 11         | 83                  |
| Dodick et al. [40] | 2020             | 45.6 ± 11       | 72.6                |
| Obermann et al. [41]| 2021            | 42.4 ± 11.4     | 83.5                |

Age is presented as mean ± SD, CCH = chronic cluster headache, ECH = episodic cluster headache.

### 3.3. Efficacy Outcomes

#### 3.3.1. Frequency of Attacks

Six studies [19,24,26,28,35,38] reported the frequency of attacks. The overall reduction in the frequency of attacks was more significantly associated with pharmacological treatment in CH patients (MD = −1.05, 95% CI = −1.62 to −0.47; p = 0.0004) (Figure 2a) without significant heterogeneity. Regarding preventive treatment, five studies [19,24,28,35,38] were included in the analysis, and one study [26] was conducted with acute treatment. No difference was observed in the reduction of the frequency of attacks between preventive and acute treatments in CH patients (I² = 0%, p = 0.99) (Figure 2b). In addition, both triptan and non-triptan drugs were associated with a reduction in the frequency of attacks in CH patients, without significant differences between the triptan and non-triptan groups (Figure 2c). The network plots of each comparison about frequency of attacks are shown in Figure 3a. In this network meta-analysis, we observed CAN use was more associated with the reduction in the frequency of attack compared to five other treatments although it was not shown in significance (p > 0.05). On the other hand, FRO use was less associated with decreasing the frequency of attacks to manage CH (p > 0.05) (Figure 4a and Supplementary Materials Figure S1).
hand, FRO use was less associated with decreasing the frequency of attacks to manage CH ($p > 0.05$) (Figure 4a and Supplementary Materials Figure S1).

Figure 2. The effects of pharmacological treatments on frequency of attacks. (a) Overall; (b) comparison between preventive and acute; (c) comparison between triptan and non-triptan. Green squares indicated effect size for each of included studies and the size of green square indicates the weight assigned to that study in the meta-analysis. Black diamond suggested as meta-analyzed measure of effect. Bold letters represented a category or subtotal of each subgroup and overall outcome.

3.3.2. Pain-Free Rate

Nine studies [20,22,25,31–34,36,37] reported pain-free rates as outcomes. The pain-free rate was higher in the intervention group (OR = 3.89, 95% CI = 2.76–5.48; $p < 0.00001$) (Figure 5). Two studies [33,37] investigated preventive treatment; seven [20,22,25,31,32,34,36] investigated acute treatment. No differences were observed between the two types of therapy, the preventive and acute treatments ($I^2 = 56.5\%$, $p = 0.13$). Treatment with triptans or non-triptans demonstrated a correlation with pain-free rate (OR = 3.88, 95% CI = 2.55–5.90; $p < 0.00001$ and OR = 3.90, 95% CI = 2.14–7.11; $p < 0.00001$, respectively) without differences according to the type of therapy. Heterogeneity was not observed in subgroup analyses. The network plots of each comparison about the pain-free rate are shown in Figure 3b. In the result of network meta-analysis, ZOL10 had a significantly better pain-free rate compared to ZOL5 (ZOL5 vs. ZOL10: OR = 0.40, 95% CI = 0.19–0.82; $p < 0.05$). BET had a higher pain-free rate than nine other treatments. WAR showed a better pain-free rate than eight other treatments except the treatment BET (BET vs. WAR: OR = 2.43, 95% CI = 0.09–63.84; $p > 0.05$). Subsequently, SUM6 had a higher pain-free rate than seven other
treatments except BET (SUM6 vs. BET: OR = 0.37, 95% CI = 0.02–8.86; \( p > 0.05 \)) and WAR (SUM6 vs. WAR: OR = 0.89, 95% CI = 0.18–4.37; \( p > 0.05 \)). However, LCAR showed a lower pain-free rate than nine other treatments without showing statistical significance (\( p > 0.05 \)) (Figure 4b and Supplementary Materials Figure S2).

Figure 3. Network comparisons of studies included in the network meta-analysis. (a) Frequency of attacks; (b) pain-free rate; (c) duration of attacks; (d) number of patients using rescue agents; (e) adverse events. Betamethasone (BET), Candesartan cilexetil (CAN), Capsaicin (CAP), Cimetidine (CIM), Civamide (CIV), Cortivazol (COR), Frovatriptan (FRO), Galcanezumab (GAL), Lithium carbonate (LCAR), Melatonin (MEL), Misoprostol (MIS), Octreotide (OCT), Placebo (PLA), Prednisone (PRE), Sumatriptan 6 mg (SUM6), Sumatriptan 12 mg (SUM12), Sumatriptan 100 mg (SUM100), Sumatriptan spray (SUMS), Valproate (VAL), Verapamil (VER), Warfarin (WAR), Zolmitriptan 5 mg (ZOL5), Zolmitriptan 10 mg (ZOL10). The size of each circle represents the proportion of the number of patients for each treatment and the width of the lines represents the proportion of the number of studies.
Figure 4. Cont.
Figure 4. Overall network meta-analysis results of each outcome. (a) Frequency of attacks; (b) pain-free rate; (c) duration of attacks; (d) number of patients using rescue agents; (e) adverse events. * Statistical significance.
when compared with ZOL10 (ZOL10 vs. SUM6: OR = 2.09, 95% CI = 0.98–4.47; $p$ = 0.07) (Figure 8), without heterogeneity. Compared with preventive treatment, a greater number of individuals needing rescue agents was associated with preventive treatment (OR = 0.37, 95% CI = 0.27–0.51; $p$ < 0.00001) compared with placebo treatment (OR = 0.17, 95% CI = 0.06–0.45; $p$ = 0.0004) compared with acute treatment (OR = 0.41, 95% CI = 0.32–0.52; $p$ < 0.00001). Comparison of triptan and non-triptan groups revealed that triptan use was associated with a lower number of rescue agent use in the intervention group than in the placebo group (OR = 0.40, 95% CI = 0.29–0.54; $p$ < 0.00001) without significant differences observed in triptans ($p$ = 0.10). Figure 3d shows the network plots of each comparison about number of patients using rescue agents. Compared with SUM12, SUM6 increased the number of patients using rescue agents (SUM6 vs. SUM12: OR = 1.30, 95% CI = 0.54–3.09; $p$ > 0.05), but SUM6 decreased the number of patients using rescue agents when compared with ZOL10 (ZOL10 vs. SUM6: OR = 2.09, 95% CI = 0.98–4.47; $p$ > 0.05). SUM12 decreased the use of rescue agents compared to ZOL10 with significant differences (ZOL10 vs. SUM12: OR = 2.71, 95% CI = 1.08–6.80; $p$ < 0.05). ZOL5 increased the number of rescue agents used compared to other treatments (Figure 4d and Supplementary Materials Figure S4).

### 3.3.4. Number of Patients Using Rescue Agents

Analysis of the number of patients using rescue agents was performed in six studies [20,21,27,30,31,34]. An increased number of rescue agents used was associated with placebo treatment (OR = 0.37, 95% CI = 0.27–0.51; $p$ < 0.00001) (Figure 7). A greater number of individuals needing rescue agents was associated with preventive treatment (OR = 0.17, 95% CI = 0.06–0.45; $p$ = 0.0004) compared with acute treatment (OR = 0.41, 95% CI = 0.32–0.52; $p$ < 0.00001). Comparison of triptan and non-triptan groups revealed that triptan use was associated with a lower number of rescue agent use in the intervention group than in the placebo group (OR = 0.40, 95% CI = 0.29–0.54; $p$ < 0.00001) without significant differences observed in triptans ($p$ = 0.10). Figure 3d shows the network plots of each comparison about number of patients using rescue agents. Compared with SUM12, SUM6 increased the number of patients using rescue agents (SUM6 vs. SUM12: OR = 1.30, 95% CI = 0.54–3.09; $p$ > 0.05), but SUM6 decreased the number of patients using rescue agents when compared with ZOL10 (ZOL10 vs. SUM6: OR = 2.09, 95% CI = 0.98–4.47; $p$ > 0.05). SUM12 decreased the use of rescue agents compared to ZOL10 with significant differences (ZOL10 vs. SUM12: OR = 2.71, 95% CI = 1.08–6.80; $p$ < 0.05). ZOL5 increased the number of rescue agents used compared to other treatments (Figure 4d and Supplementary Materials Figure S4).

### 3.4. Adverse Events

Analysis of the number of patients who experienced AEs was performed in 15 studies [21,23–30,33,36,38–41]. In the intervention group, a greater number of patients experienced AEs compared with the placebo group (OR = 2.28, 95% CI = 1.73–3.00; $p$ < 0.00001) (Figure 8), without heterogeneity. Compared with preventive treatment, a greater number
of patients treated with acute pharmacological therapies experienced AEs ($I^2 = 88.6\%$, $p = 0.003$). In the subgroup analysis, the triptan group had a greater number of patients with AEs in the treatment group than in the control group (OR $= 2.79$, 95% CI $= 2.07$–$3.76$; $p < 0.00001$). Similarly, in the non-triptan group, the number of patients who experienced AEs in the treatment group was greater than that in the control group (OR $= 1.78$, 95% CI $= 1.30$–$2.44$; $p = 0.0003$). A greater number of patients experienced AEs in the triptan group than in the non-triptan group ($p = 0.04$). Once again, the analysis revealed no heterogeneity. The network plots of each comparison about adverse events are shown in Figure 3e. According to the network meta-analysis, SUM6 decreased the number of patients who experienced AEs compared with ZOL10 with significant differences (ZOL10 vs. SUM6: OR $= 2.53$, 95% CI $= 1.40$–$6.15$; $p < 0.05$). PRE experienced a smaller number of patients with adverse events than fourteen other treatments except MIS (MIS vs. PRE: OR $= 0.99$, 95% CI $= 0.05$–$20.37$; $p > 0.05$). On the other hand, BET indicated a greater number of patients who experienced adverse events than other treatments (Figure 4e and Supplementary Materials Figure S5).

Figure 8. The overall effect of pharmacological treatment for the number of patients with adverse events compared to placebo. Blue squares indicated effect size for each of included studies and the size of blue square indicates the weight assigned to that study in the meta-analysis. Black diamond suggested as meta-analyzed measure of effect. Bold letters represented a category or subtotal of overall outcome.

3.5. Rank Probability and SUCRA

The cumulative rank probabilities of each treatment are ranked in Figure 9, and SUCRA results based on the efficacy and safety outcomes are provided in Supplementary Tables S1–S5. Using the Bayesian network analysis, the SUCRA results showed the rank probabilities of all treatments from best treatment effect to the worst. Treatments with a larger area in Figure 9 were associated with larger probabilities of better outcomes. The results suggested that in the aspect of efficacy, CAN reduced the frequency of attacks the most, with a SUCRA value of 84%, while FRO reduced the frequency of attacks the least, with a SUCRA value of 18% (Supplementary Materials Table S1). BET had the highest pain-free rate, with a SUCRA value of 99%, while LCAR had the lowest, with a SUCRA value of 19%, except placebo (Supplementary Materials Table S2). MIS decreased the duration of attacks the most, with a SUCRA value of 77%, and SUM12 showed the highest value of SUCRA for the number of patients using rescue agents (83%) (Supplementary Materials Tables S3 and S4). In the aspect of safety, MEL had the smallest number of patients with adverse events and had the highest SUCRA value (76%), while BET had the greatest number of patients with adverse events and had the lowest SUCRA value (1%), except placebo (Supplementary Materials Table S5).
Figure 9. Cumulative rank with SUCRA (the surface under the cumulative ranking curve) for each treatment. The probability of each treatment distribution is displayed on the graph. (a) Frequency of attacks; (b) pain-free rate; (c) duration of attacks; (d) rescue agents; (e) adverse events.
3.6. Risk of Bias and Quality of Evidence

The assessment of the risk of bias among the included studies is shown in Figure 10. The risk of selection bias was not clear in ten studies [19,22,24–26,28,31,32,34,35] and one study demonstrated a high risk of bias in terms of reporting [21]. A low risk of bias in terms of selection, performance, detection, consumption, and reporting was found in the remaining studies [20,23,27,29,30,33,36–41]. Egger’s regression test showed no evidence of publication bias ($p = 0.75$) (Figure 11). The quality of evidence according to the GRADE approach with regard to the effects of interventions for CH is summarized in Table 3.

![Figure 10. Risk-of-bias assessment of the studies. High risk of bias was marked as red (−), low risk of bias was marked as green (+) and unclear risk of bias was marked as yellow (?).]
Figure 10. Risk-of-bias assessment of the studies. High risk of bias was marked as red (-), low risk of bias was marked as green (+) and unclear risk of bias was marked as yellow (?).

Figure 11. Publication bias of the studies analyzed.

Table 3. Summary of findings for efficacy and safety outcomes comparing interventions to comparators based on the GRADE approach.

| Outcomes                        | No. of Participants (Studies) | Limitation | Inconsistency | Indirection | Imprecision | Publication Bias | Pooled Estimates | Quality of Evidence |
|---------------------------------|-----------------------------|-----------|---------------|-------------|-------------|-----------------|------------------|-------------------|
| **Efficacy outcomes**           |                             |           |               |             |             |                 |                  |                   |
| Frequency of attacks            |                             |           |               |             |             |                 |                  |                   |
| Overall                         | 133 (6)                     | No serious| No serious    | No serious  | No serious  | No serious      | $-1.05$ ($-1.62, -0.47$) | ⚫⚫⚫⚫ High          |
| Acute                           | 117 (5)                     | No serious| No serious    | No serious  | No serious  | No serious      | $-1.05$ ($-1.62, -0.47$) | ⚫⚫.flip High         |
| Transitional                    | 133 (1)                     | No serious| No serious    | No serious  | No serious  | No serious      | $-1.00$ ($-12.28, 10.28$) | ⚫⚫.flip High       |
| Triptans                        | 11 (1)                      | No serious| No serious    | No serious  | No serious  | No serious      | $4.00$ ($-6.04, 14.04$) | ⚫⚫⚫ High            |
| Non-triptans                    | 122 (5)                     | No serious| No serious    | No serious  | No serious  | No serious      | $-1.06$ ($-1.64, -0.49$) | ⚫⚫ High             |
| Duration of attacks             |                             |           |               |             |             |                 |                  |                   |
| Overall                         | 143 (3)                     | No serious| No serious    | No serious  | No serious  | No serious      | $-1.08$ ($-13.60, 11.44$) | ⚫⚫⚫⚫ High          |
| Pain-free rate                  |                             |           |               |             |             |                 |                  |                   |
| Overall                         | 747 (9)                     | No serious| No serious    | No serious  | No serious  | No serious      | $3.89$ ($2.76, 5.48$) | ⚫⚫⚫ High            |
| Preventive                      | 91 (2)                      | No serious| No serious    | No serious  | No serious  | No serious      | $8.90$ ($2.85, 27.79$) | ⚫ High             |
| Acute                           | 656 (7)                     | No serious| No serious    | No serious  | No serious  | No serious      | $3.53$ ($2.46, 5.07$) | ⚫ High             |
| Triptans                        | 482 (3)                     | No serious| No serious    | No serious  | No serious  | No serious      | $3.58$ ($2.55, 5.90$) | ⚫ High             |
| Non-triptans                    | 747 (6)                     | No serious| No serious    | No serious  | No serious  | No serious      | $3.50$ ($2.14, 7.11$) | ⚫ High             |
| Number of people needing rescue agents |                     |           |               |             |             |                 |                  |                   |
| Overall                         | 1388 (6)                    | No serious| No serious    | No serious  | No serious  | No serious      | $0.37$ ($0.27, 0.51$) | ⚫⚫ High            |
| Preventive                      | 846 (10)                    | No serious| No serious    | No serious  | No serious  | No serious      | $0.17$ ($0.06, 0.45$) | ⚫ High             |
| Acute                           | 1293 (5)                    | No serious| No serious    | No serious  | No serious  | No serious      | $0.41$ ($0.32, 0.52$) | ⚫ High             |
| Triptans                        | 1293 (3)                    | No serious| No serious    | No serious  | No serious  | No serious      | $0.40$ ($0.29, 0.54$) | ⚫ High             |
| Non-triptans                    | 95 (1)                      | No serious| No serious    | No serious  | No serious  | No serious      | $0.17$ ($0.06, 0.45$) | ⚫ High             |
| **Safety outcomes**             |                             |           |               |             |             |                 |                  |                   |
| Adverse events                  |                             |           |               |             |             |                 |                  |                   |
| Overall                         | 1822 (16)                   | No serious| No serious    | No serious  | No serious  | No serious      | $2.16$ ($1.59, 2.94$) | ⚫⚫ High            |
| Preventive                      | 846 (10)                    | No serious| No serious    | No serious  | No serious  | No serious      | $1.66$ ($1.23, 2.23$) | ⚫ High             |
| Acute                           | 928 (5)                     | No serious| No serious    | No serious  | No serious  | No serious      | $3.19$ ($2.33, 4.39$) | ⚫ High             |
| Triptans                        | 1010 (3)                    | No serious| No serious    | No serious  | No serious  | No serious      | $2.79$ ($2.07, 3.76$) | ⚫ High             |
| Non-triptans                    | 812 (13)                    | No serious| No serious    | No serious  | No serious  | No serious      | $1.64$ ($1.21, 2.23$) | ⚫ High             |

3.7. Meta-Regression Analysis

Male proportion (coefficient = $-6.71$, 95% CI = $-14.21$ to $0.79$; $p = 0.08$) and age (coefficient = $0.17$, 95% CI = $-0.02$ to $0.36$; $p = 0.08$) did not significantly affect the pain-free rate. Moreover, male proportion (coefficient = $1.35$, 95% CI = $-13.47$ to $16.17$; $p = 0.89$) and age (coefficient = $-0.016$, 95% CI = $-0.19$ to $0.15$; $p = 0.85$) were not associated with
an increase or decrease in the use of rescue agents. Male proportion (coefficient = 3.65, 95% CI = −0.41 to 7.71; \( p = 0.08 \)) and age (coefficient = −0.08, 95% CI = −0.17 to 0.01; \( p = 0.09 \)) did not significantly influence the effect of pharmacological treatment on CH in AEs (Supplementary Materials Figure S6).

4. Discussion

The current systematic review and meta-analysis aimed to compare pharmacological treatment with placebo in CH patient care. According to our results, the use of pharmacological therapies is an effective option to treat CH. Compared to placebo use, the present study demonstrated that the use of pharmacological treatment in CH patients was associated with reduced frequency and duration of attacks. Frequent severe headache attacks—likely associated with severe CH pain—impaired patient quality of life or restricted activities of daily living, which led to losses of employment and economic burdens [3]. According to Palacios Ceña et al. [42], frequent headaches negatively impact patient health, resulting in increased headache intensity and psychiatric disturbances, sometimes causing depression. Sohn et al. consistently reported that clinical factors, such as increased duration and number of pain attacks, were closely associated with disability in CH patients [43]. Considering frequency of attacks as one of important measures for prevention in CH treatment, as the present study showed, pharmacological therapy should be first engaged in the treatment of CH patients [12,44]. Moreover, previously, a systematic review and meta-analysis also demonstrated the efficacy of pharmacologic treatment, using galcanezumab, in migraine and CH in reducing headache frequency compared with placebo [11]. Although the systematic review and meta-analysis provided quantitative evidence for using pharmacological treatment instead of placebo, the study included too limited a number of CH patients to provide confidence in the evidence in practice. Furthermore, Probyn et al. also showed non-pharmacological self-management was not associated with reducing headache frequency (standard mean difference = −0.07, 95% CI = −0.22 to 0.08) [45]. Individually, the current study showed that CAN was more effective than other treatments in reducing the frequency of attacks. Etminan et al. also reported that angiotensin II receptor antagonists including candesartan reduce the frequency of headache [46]. Without comparison between medicines in the previous study and a limited number of studies analyzed, the current results should be cautiously interpreted. Because considering evidence from active-controlled trials without comparison to placebo verifying the rationality of accepted criteria is regarded as secondary evidence [12], the current study might be a supportive suggestion of pharmacological treatment for CH patient care in real-world practice.

In addition, the current study demonstrated that pharmacological therapies reduced the need for rescue agents in CH patients, providing information regarding adequate pain control [47,48]. Overuse of rescue medications for relieving headache has been frequently reported in previous studies [49,50], but increased numbers of acute medications as rescue therapy are rather associated with developing chronic headache or medication overuse headache [51]. With prior research investigating methods to reduce acute medication use in headache treatment, decreased need for rescue agents resulting from proper pharmacological treatments may represent an improvement in adherence to CH therapy as well; this displays a close association with the discontinuation of therapy [11]. Seeing as the United States Food and Drug Administration also indicated that rescue medication use should be considered an endpoint in studies investigating pain management, appropriate pharmacological therapies might contribute to reducing the number of rescue agents used by CH patients [47,48].

For more effective individualized medication regimens, in the current analysis, pharmacological treatment is subdivided according to the aims of therapy or active ingredients. Along with analyzing discrepancies among therapies subdivided with aims or active ingredients to treat CH patients, the current study did not demonstrate significant differences of efficacy according to the subdivisions. For recommending appropriate pharmacological therapies, prior studies reported different levels of evidence for the efficacy of therapeutics
based on the aims of studies, which could not draw consistent conclusions through literature reviews [12,13]. Furthermore, although a prior meta-analysis demonstrated the efficacy of therapies including triptans as active ingredients to treat CH, the analysis only compared to placebo without comparison between therapies [52]. Although comparison between triptans and non-triptans showed no differences, among types of triptans, different efficacies were observed in types such as sumatriptans and zolmitriptans for reducing rescue agents or increasing the pain-free rate. As Pomeroy et al. also reported different efficacies among medicines including triptans, we need to consider different triptan use according to the patients’ statuses [53]. In contrast, the occurrence of AEs was differently associated with classifications depending on the treatment aims and active ingredients in the current study. Dodick et al. also reported, based on the aims of treatment, that acute treatment was more associated with the occurrence of AEs than prevention or transitional treatments with insistent needs to control symptoms due to the rapid onset [13]. Nevertheless, frequent daily dosing for treating attacks may lead to overmedication or toxicity [13], which may be associated with an increased number of individuals experiencing AEs. However, as the majority of CH patients received both preventive and acute types of medications [44], more patient-specific therapies should be applied in the practical realm. Furthermore, Law et al. showed AEs were more common with triptans than with placebo in the care of CH [52] consistent with the current study; therefore, depending on the patient’s clinical needs, a trial-and-error approach should be attempted using available pharmacological therapies in CH management [12,44].

The current study had several limitations. First, it did not evaluate the cost-effectiveness of pharmacological treatments compared with placebo in treating patients with CH. Although cost-effectiveness evaluation of therapies is an important issue, it was beyond the scope of the present study; therefore, further studies are needed. Second, the main issue of the current study is the lack of studies included to analyze differences of subgroups among aims and active ingredients. Although CH is a rare disease, complicating the ability to enroll patients to accomplish trials, less than two studies in one subgroup in the analysis could limit clinical utility. Thus, results evaluated according to subgroup classifications based on aims or active ingredients in the present study should be cautiously applied in practice; therefore, future studies in various clinical settings are expected to evaluate various clinical parameters. Third, the present study did not evaluate the sex differences related to pharmacological efficacy and safety. Although previous research has indicated different tendencies of CH development according to sex, studies included in the current analysis did not provide outcomes according to sex. Despite the scarcity of information, the current study revealed an insignificant correlation between efficacy or safety outcomes and male sex, which is consistent with reports describing a decreasing male predominance in CH development [12,54,55]. Fourth, the current study did not include head-to-head comparative trials among interventions. Considering advantages of systematic review and network meta-analysis such as generalizability and providing estimates [14,15], the current study could provide supportive evidence in clinical practice. However, to provide more precise outcomes in decision making, more future comparative studies among interventions are expected. Finally, studies that evaluated oxygen to manage CH patients were not included in the current analysis. As previous literature provided evidence related to the efficacy of oxygen treatment in CH patients [56], oxygen should be considered as an additional treatment for CH care. However, because of the controversies surrounding the inclusion of oxygen as a drug [57–59], we could not include studies with oxygen, and we expect more future studies evaluating oxygen as CH treatment.

5. Conclusions

The results of the present systematic review and network meta-analysis demonstrated that pharmacological treatments were significantly associated with a reduction in the frequency and duration of attacks and need for rescue agents and an increased pain-free rate in CH patients compared with placebo. In the subgroup analysis, there were
no differences of efficacy according to treatment aims and active ingredients. However, pharmacological treatments were associated with an increase in AEs in patients with CH, especially in acute treatment and medications containing triptans. Based on the efficacy and safety of pharmacological therapies, individualized therapies should be applied to treat CH in real-world practice.

**Supplementary Materials:** The following supporting information can be downloaded at https://www.mdpi.com/article/10.3390/jcm11051411/s1, Figure S1: Direct and indirect comparison in terms of frequency of attacks; Figure S2: Direct and indirect comparison in terms of pain-free rate; Figure S3: Direct and indirect comparison in terms of duration of attacks; Figure S4: Direct and indirect comparison in terms of adverse events; Figure S5: Meta-regression results with male proportion and age. Table S1: The result of surface under the cumulative ranking curve (SUCRA) for frequency of attacks; Table S2: The result of surface under the cumulative ranking curve (SUCRA) for pain-free rate; Table S3: The result of surface under the cumulative ranking curve (SUCRA) for duration of attacks; Table S4: The result of surface under the cumulative ranking curve (SUCRA) for number of patients using rescue agents; Table S5: The result of surface under the cumulative ranking curve (SUCRA) for frequency of attacks.

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