Different Dosage Regimens of Eptinezumab for the Treatment of Migraine: A Meta-Analysis from Randomized Controlled Trials

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Abstract

Background
Migraine is one of the most common neurological diseases around the world and calcitonin gene-related peptide (CGRP) plays an important role in its pathophysiology. Therefore, in the present study, we evaluated the efficacy of monoclonal antibodies blocking the CGRP ligand or receptor in episodic and chronic migraine.

Objective
The objective of our study is implementing a meta-analysis to systematically evaluate the efficacy and safety of eptinezumab for the treatment of migraine compared with placebo.

Method
We searched the Medline, Embase, Cochrane Library and Clinicaltrials.gov for randomized controlled trials (RCTs) which were performed to evaluate eptinezumab versus placebo for migraine up to September 2020. The data was assessed by Review Manager 5.3 software. The risk ratio (RR) and standard mean difference (SMD) were analyzed using dichotomous outcomes and continuous outcomes respectively with a random effect model.

Result
We collected 2,739 patients from 4 RCTs: the primary endpoint of efficacy was the change from baseline to week 12 in mean monthly migraine days (MMDs). We found that eptinezumab (30mg/100mg/300mg) led to a significant reduction in MMDs (P=0.0001, P < 0.00001, P < 0.00001) during 12 weeks compared with placebo, especially with 300mg. For the safety, we compared and concluded the treatment emergent adverse events (TEAEs) of the 4 RCTs. This indicated no evident statistical difference between eptinezumab and placebo.

Conclusions
In the present study, we found that eptinezumab is safe and has significant efficacy in the treatment of migraine, especially the dose of 300 mg.

Introduction
Migraine is one of the most prevalent disorder in neurologic field which can be divided into episodic and chronic migraine. Both types of migraine can disturb the common life of the patients [1]. Presently, more than 16% of the global population is affected by the attacks of migraine [2]. Unfortunately, since many drugs are available for the therapy of migraine, such as NSAIDS, triptans, ergots, they all have some contraindications or severe adverse effects in certain aspects [3]. Therefore, further studies are needed on the treatment of the migraine, which ultimately may help patients suffering from it by improving the adverse effects.

The release of calcitonin gene-related peptide (CGRP) plays an important role in migraine pathophysiology, which has been observed after the migraine attack [4, 5]. Over the past few years, the efficacy of monoclonal antibodies blocking the CGRP ligand or receptor including galcanezumab, fremanezumab and ubrogepant have been demonstrated in both episodic and chronic migraine [6]. Eptinezumab (ALD 403), a new monoclonal antibody that selectively inhibits both α-CGRP and β-CGRP, was available in the market since February 2019. However, there were no systematic review or meta-analysis comprehensively evaluating the efficacy and safety of eptinezumab in the treatment of migraine [7–9].

Therefore, in the present study, we performed a meta-analysis to discuss different dosage regimen of eptinezumab for the treatment of migraine. In the previous clinical trials, eptinezumab had exhibited flexible dosing regimens (10 mg, 30 mg, 100 mg, 300 mg, 1000 mg). During our study, we combined different doses of eptinezumab to analyze the efficacy and safety for the therapy of episodic and chronic migraine [10–13].

Methods

Study protocol
Before we started the project, we drafted a research protocol by following the Cochrane Collaboration format [14]. The meta-analysis was retrospectively registered.

Search strategy
Original researches in the MEDLINE, Embase, Cochrane Library and Clinicaltrials.gov were searched using the following terms: ["eptinezumab and migraine") ("ALD403 and migraine") until September 2020. Moreover, to make sure all relevant studies have been included, we screened reference lists of relevant articles manually.

Study selection
Studies were included as follows: (1) study type was randomized clinical trials; (2) enrolled participants diagnosed with migraine; (3) study used eptinezumab as intervention; (4) study period was over 12 months; (5) participants were over 18 years old. Studies were excluded as follows: (1) types of study: retrospective studies, cohort studies, case reviews and case reports; (2) control: active control (i.e. that a known, effective treatment as opposed to a placebo is compared with an experimental treatment).
Data extraction

All the data were extracted independently by 2 investigators (ZYY and TX) and any disagreements were settled through discussion. After several selections and assessments, the basic information of the included trails (first author, publication, country, centers, and treatment groups), patient characteristics (Age range, mean age and gender), study period and outcome events were used to extract the data (Table 1).

| Study             | Countries | Centers | Publication  | Type of migraine | Treatment group, (No. of participants) | Total number | Age range | Male (%) | Mean age ± SD (year) |
|-------------------|-----------|---------|--------------|------------------|----------------------------------------|--------------|-----------|-----------|---------------------|
| Dodick et al      | USA       | 26      | Lancet Neurol| episodic         | Eptinezumab1000mg (81) vs.PL (82)       | 163          | 18y-55y  |           | Eptinezumab 1000 mg:17 PL:20 |
| 2014 (NCT01772524)|           |         |              |                  |                                        |              |           |           | Eptinezumab 1000 mg:10.8 PL:38.6 ± 9.6 |
| Dodick et al      | 5         | 92      | Cephalalgia  | chronic          | Eptinezumab300mg (121) vs.100 mg (122) vs. 30 mg (122) vs.10 mg (130) vs. PL (121) | 616          | 18y-55y  |           | Eptinezumab 300 mg:19 PL:20 |
| 2019 (NCT02275117)|           |         |              |                  |                                        |              |           |           | Eptinezumab 300 mg:37.2 ± 10.0 |
| Ashina et al      | 2         | 84      | Cephalalgia  | episodic         | Eptinezumab300mg (224) vs.100 mg (223) vs. 30 mg (219) vs. PL (222) | 888          | 18y-75y  |           | Eptinezumab 300 mg:11.2 PL:10 |
| 2020 (NCT0255989)|           |         |              |                  |                                        |              |           |           | Eptinezumab 300 mg:40.2 ± 11.72 |
| Lipton et al      | 13        | 128     | Neurology    | chronic          | Eptinezumab300mg (350) vs.100 mg (356) vs. PL (366) | 1072         | 18y-65y  |           | Eptinezumab 300 mg:10.3 PL:10 |
| 2020 (NCT02974153)|           |         |              |                  |                                        |              |           |           | Eptinezumab 300 mg:41.0 ± 10.4 |

PLA: placebo; a: Adverse Events (AE) and Serious Adverse Events (SAE); b: monthly migraine days(MMDs); c: 50% responders rate; d: 75% responders rate; e: rate; f: Headache Impact Test(HIT-6) score; g: headache days; h: migraine hours; i: migraines with severe intensity; j: Percentage of patients with migraine, day

Outcomes

The primary efficacy outcome is mean monthly migraine days (MMDs), baseline to 12 week. Secondary efficacy endpoint included: patients with a 75% reduction in migraine days from baseline:75% responder rate, patients with a 50% reduction in migraine days from baseline:50% responder rate, and patients with migraine 1 day after dosing, baseline to 12 weeks. In addition, we choose the treatment emergent adverse events (TEAEs) as the safety endpoint.

Summary Measures and Synthesis of Results

Review manager 5.3 was used to assess the data. Estimated standard mean differences and estimated risk ratio (standard mean difference [SMD] or risk ratio [RR]; 95% confidence interval [CI]) were calculated using a random effects model. The $I^2$ statistic was used to estimate the statistical heterogeneity as follows: $I^2 < 30\%$ represents "low heterogeneity," $30\% < I^2 < 50\%$ means "moderate heterogeneity" and $I^2 > 50\%$ means "substantial heterogeneity." A $< 0.05$ P-value was considered to be significant for all analyses, and tests are two-tailed.

Risk of Bias
The risk-of-bias plot was assessed using Review Manager 5.3 software (The Cochrane Collaboration, Oxford, UK) for individual studies. The unified standard of the Cochrane Collaboration was applied to assess the risk of bias for RCTs, which included selection bias, performance bias, detection bias, attrition bias, reporting bias, and other potential biases.

Results

Search results

A total of 464 researches and abstracts from Medline, Embase, Cochrane library and Clinicaltrials.gov were identified. Among them, 190 studies were excluded due to duplicates. Further, 178 studies were excluded as they were irrelevant, such as research on other drugs or into the etiological analysis of migraine. After removing duplicates and uncorrelated titles, 96 of these articles were directly related to the topic of interest. Among them, 92 full text articles were excluded, which included 13 conferences, 4 comments, 46 reviews, 2 short survey and 27 summarizations. Finally, 4 RCTs containing 2,739 patients were included in our meta-analysis. The detailed process of screening is shown in Fig. 1.

Different dosage regimen for the efficacy and safety

The primary efficacy outcome was mean monthly migraine day (MMDs). From the perspective of primary efficacy outcome, treatment with 30mg (MD=0.29, 95% CI:0.45~0.14, P=0.0001) or 100mg (MD=0.31, 95% CI:0.42~0.21, P<0.00001) and 300mg (MD=0.41, 95% CI:0.52~0.30, P<0.00001) eptinezumab showed significant efficacy compared to the placebo (Figure 2). Whereas, for the secondary efficacy endpoint 75% responder rate, 50% responder rate and patients with migraine 1 day after dosing, the outcomes were abundant with the MMDs (Figure 3–5). Initially, for the 75% responder rate, treatment with 30mg (RR=1.46:95% CI:1.09~1.95, P=0.01), 100mg (RR=1.59:95% CI:1.29~1.96, P=0.0001), 300mg (RR=1.95:95% CI:1.60~2.39, P=0.0001) and 1000mg (RR=3.57:95% CI:1.63~7.81, P=0.001) indicated that they could increase the rate significantly. Further, for the 50% responder rate, treatment with 30mg (RR=1.35:95% CI:1.14~1.60, P=0.0004), 100mg (RR=1.41:95% CI:1.25~1.58, P<0.0001), 300mg (RR=1.52:95% CI:1.36~1.70, P<0.0001) and 1000mg (RR=1.86:95% CI:1.28~2.70, P=0.001) eptinezumab also showed improved efficacy compared to the placebo. Nevertheless, from the perspective of the data related to patients with migraine 1 day after dosing, treatment with 1000mg (RR=0.55:95% CI:0.46~0.67, P=0.00001) and 300mg (RR=0.65:95% CI:0.53~0.74, P<0.0001) appeared to be more effective than 30mg (RR=0.78:95% CI:0.53~1.13, P=0.19) compared with the placebo.

Of all the 2,739 patients receiving eptinezumab, no deaths occurred during the treatment period. The common adverse effects contained upper respiratory tract infection, nausea and sinus congestion. Therefore, we summarized the adverse events which showed the treatment with 10mg (RR=1.01:95% CI:0.82~1.26; p=0.91), 30mg (RR=0.92:95% CI:0.77~1.10; p=0.35), 100mg (RR=1.01:95% CI:0.91~1.11; p=0.92), 300mg (RR=1.06:95% CI:0.96~1.17; p=0.24), 1000mg (RR=1.08:95% CI:0.82~1.43; p=0.58) had no evident statistical difference between eptinezumab and placebo (Figure 6).

Dosage regimen of 100 mg vs. 300 mg

Further, study was carried out to compare the efficacy between the 100mg and 300mg (Figure 7a~d). Result from this comparison showed that the treatment with 300mg (MD=-0.10:95% CI:0.00~0.21; p=0.06, 75% responder rate; RR=0.81, 95% CI:0.69~0.96; p=0.01, 50% responder rate; RR=0.93, 95% CI:0.85~1.02; p=0.11) patients with migraine 1 day after dosing, RR=0.92, 95% CI:0.69~1.23, p=0.58) was more promising than 100mg. Meanwhile, as shown in Figure 7e, no differences were existed in TEAEs between the 100mg and 300mg (RR=0.94:95% CI:0.79~1.12; p=0.51).

Risk of bias

The independent risk of biases related to 4 RCTs are shown in Fig. 8. The risk for attrition bias is unclear in the studies carried out by Dodick (2019) and Lipton (2020). In addition to the measure, other studies had reported low risks of bias.

Discussion

Migraine is a prevalent neurological disease around the globe. However, previous therapies have some limitations or adverse effects, and are unresolved until now. As the importance of CGRP in the pathogenesis has been proved by the previous studies, its receptors are widely distributed in the central nervous system (CNS) and peripheral sensory neurons. Therefore, monoclonal antibodies blocking the CGRP ligand or receptor have a clear advantage in the treatment strategy for episodic and chronic migraine[15].

Our study is the first meta-analysis about different dosage regimens related to the safety and efficacy of eptinezumab in the treatment for migraine, and indicated eptinezumab as excellent therapeutic agent for the migraine. During our study, we pooled 2,739 participants from 4 randomized clinical trials (RCTs), which provided high clinical reliability in the research for the use of eptinezumab. Further, we gathered primary data from those articles and did not discover apparent heterogeneity in our outcomes as indicated by our statistical analysis. Subsequently, we found that eptinezumab had been divided into flexible dosage regimes in these RCTs, including 10 mg, 30 mg, 100 mg, 300 mg, 1000 mg. Further, by comparing the primary efficacy outcomes mean monthly migraine days (MMDs), baseline to 12 week, we proved that treatment with 30 mg, 100 mg, 300 mg can cause effective reduction in monthly migraine days (MMDs) compared with placebo. Whereas, for the secondary endpoint, all dosage regimens of eptinezumab increased the proportion of 75% responder rate except 10 mg. Similar results were observed in 50% responder rate. In addition, fewer patients suffered from migraine 1 day after 100 mg and 300 mg eptinezumab administration compared with 30 mg. Due to the lack of research and subsequent data, we could not continue further exploration of 10 mg (only in the study conducted by Dodick et al 2014) and 1000 mg (only in the study conducted by Dodick et al 2019) for the efficacy of eptinezumab. However, it doesn't mean that these dosage regimens were insignificant, probably research related to it needs more time for comprehensive outcome.

By analyzing the results of different dosage regimes of eptinezumab, we found that the dosage regimens of 100 mg and 300 mg were more significant in the efficacy of the treatment for migraine. Therefore, further study was carried out for these two dosage regimes. From the perspective of the outcome related to
the MMDs, baseline to 12 week, 300 mg (P = 0.06) eptinezumab showed no significant difference but potential tendency for the reduction of MMDs compared with 100 mg. Nevertheless, for the 75% responder rate, 300 mg eptinezumab has been proved more increasing proportion than 100 mg. The result of 50% responder rate and patients with migraine 1 day after dosing couldn't indicate the difference between 100 mg and 300 mg. Nonetheless, we can conclude that the dosage regimen of 300 mg has an advantage on the efficacy of the treatment for migraine.

During our study, the analysis of safety outcomes—TEAEs did not indicate existence of statistical difference between eptinezumab and placebo (P > 0.05). Therefore, generally the use of eptinezumab is safe for the treatment of migraine. The result was consistent with the meta-analysis conducted by Da Xu and Deng Chen [16] which demonstrated monoclonal antibodies blocking the CGRP ligand or receptor are safe. As reported in the previous studies, we observed that eptinezumab rarely causes serious adverse events or even death [17, 18]. Moreover, it only resulted in some mild adverse events such as upper respiratory tract infection, nausea and sinus congestion, just like the other monoclonal antibodies blocking the CGRP ligand or receptor[19]. Certainly, these studies on adverse events merely evaluated 12 weeks after the first dose. We cannot ensure whether eptinezumab will produce long lasting influence. Therefore, it still needs further comprehensive research.

After the analysis of our data, we found few limitations in our study which cannot be avoided through existing researches. First, numerous previous studies have concluded evidence to use other CGRP monoclonal antibodies such as ubrogepant, galcanezumab and rimegepant, for the treatment of migraine [20–22]. However, as interventions in our study were related to different dosage regimens of eptinezumab and placebo, we can only conclude the advantages of eptinezumab compared with placebo. Probably, our study needs more horizontal comparison of eptinezumab with other CGRP monoclonal antibodies in the future. Secondly, considering different dosage regimens in 4 RCTs, 1000 mg eptinezumab merely conducted by Dodick 2014, whereas, 10 mg merely conducted by Dodick 2019. Moreover, part of statistics from Dodick 2019, Ashina 2020 and Lipton 2020 did not indicate standard deviation (SD) clearly. However, in the present study, we ultimately achieved SD using statistical algorithm on our own. Therefore, the accuracy of the results needs further verification. Except for the limitation above, we also cannot ignore the lack of adherence in the therapy of migraine which occurred in our 4 RCTs in a way. This also encountered by few traditional treatments for migraine [23, 24].

**Conclusion**

In conclusion, eptinezumab showed outstanding efficacy for the treatment of migraine, especially dosage regimen of 300 mg. Meanwhile, no apparent differences existed when compared with placebo from the perspective of safety. Nonetheless, we are looking forward for more studies related to the eptinezumab so that it may have a promising future in the therapy strategy of migraine.

**Abbreviations**

CGRP: calcitonin gene-related peptide; RCTs: randomized controlled trials; RR: risk ratio; SMD: standard mean difference; MMDs: monthly migraine day; TEAEs: treatment emergent adverse events; CI: confidence interval; CNS: central nervous system.

**Declarations**

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**Conflict of Interest**

The authors declare that they have no competing interests.

**Ethics approval**

Not applicable.

**Consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Availability of data and material**

All data generated or analyzed during this study are included in this published article and its supplementary information files.

**Code availability**

Not applicable.

**Authors’ Contributions**
ZW and ZC were the principal investigators. ZYY and TX designed the study and developed the analysis plan; SJC analyzed the data and performed the meta-analysis; GJL and XYY contributed to the writing of the article. XW and SG revised the manuscript and polished the language. All authors read and approved the final submitted paper.

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Not applicable.

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