Abstract. Infantile haemangioma (IH) is a benign vascular tumour type that occurs in 3-10% of infants. In the present meta-analysis, previous studies comparing clinical outcomes, including the recovery rate and haemangioma activity score (HAS), adverse effects and relapse rates, were compared between patients treated with atenolol and those treated with propranolol for IH. A systematic search in various databases, including Medline, Cochrane Controlled Register of Trials, ScienceDirect and Google Scholar from inception until July 2019 was performed. The Cochrane risk of bias tool was used to assess the quality of published trials. A meta-analysis with a random-effects model and reported pooled mean differences (MD) or odds ratios (OR) with 95% CIs was performed. In total, 8 studies including 608 participants were analyzed. Only 2 studies were randomized controlled trials, while the majority of studies had low or unclear bias risks. Except for the response to medication (pooled OR=1.49; 95% CI, 0.85-2.18), all other outcomes (HAS, adverse reactions and relapse rate) were better for the atenolol group than the propranolol group. Atenolol resulted in better HAS (pooled MD=0.16; 95% CI, -0.42 to 0.73). Propranolol had more adverse reactions (pooled OR=2.17; 95% CI, 0.93-5.06) and a higher relapse rate (pooled OR, 1.67; 95% CI, 0.44-6.41) when compared to atenolol. However, these findings were not statistically significant. The results of this analysis suggest that atenolol may be non-inferior to propranolol and may offer advantages, including lower adverse reactions and relapse rates.

Introduction

Infantile haemangioma (IH) is a type of benign vascular tumour that occurs in 3-10% of infants (1). While most of these lesions are asymptomatic and subside by the age of 5 years, complications may arise, including painful bleeding, ulceration or disfiguration (2,3). IH may also induce mental distress to the parents and children (4). An early intervention is required in such cases to prevent future complications.

Systematic steroids were used as a first-line medication for the treatment of IH. However, the long-term use of steroids may lead to serious adverse reactions, including growth delay, adrenal cortical insufficiency and/or hypertension (5). To overcome these adverse reactions, The Food Drug Administration (FDA) of the US approved beta (β) blockers as the first-line medications for the management of IH in 2014 (6). Propranolol is a non-selective lipophilic β blocker proven to be effective against His (7) by inhibiting the proliferation and inducing the regression of the lesion during the proliferative phase (8). However, propranolol treatment also has certain risks, including side effects of diarrhoea, hyperkalaemia, hypoglycaemia and bronchial hyperreactivity. Propranolol also affects the central nervous system (CNS) as it crosses the blood-brain barrier due to its lipophilic nature and may cause adverse reactions, including agitation and sleep disturbances. These undesired effects from propranolol have led to discontinuation of therapy and regrowth of the lesions (9-11).

Atenolol, a hydrophilic β blocker, has been used as an alternative to propranolol in the treatment of IH (12). Atenolol has minimal safety concerns, as it primarily acts on β1 receptors with minor effects on β2 receptors (13). As it does not act on pulmonary β2 receptors, it may be safely used in infants with pulmonary diseases (e.g. reactive airway conditions). Atenolol also does not affect the pancreatic β2 receptors and does not interfere with the glycogenolysis, gluconeogenesis or lipolysis (14). Due to its hydrophilic nature, it does not cross the blood-brain barrier and has limited adverse reactions when compared to propranolol (15). Studies have reported that atenolol is as effective as propranolol in reducing the size of the IH lesions (13,14).

Even though the morbidity profile of atenolol for the management of IH has been established, only a few systematic studies have compared the clinical outcomes and/or adverse
effects between these two different treatment methods (16,17). The purpose of the present meta-analysis was to compare clinical outcomes [recovery rate, haemangioma activity score (HAS), adverse effects and relapse rates] between patients treated with atenolol and those treated with propranolol in the management of IH.

Materials and methods

Search strategy. An extensive search was performed in the following databases: Medline (PubMed) (https://pubmed.ncbi.nlm.nih.gov/), Google Scholar (https://scholar.google.com/), ScienceDirect (https://www.sciencedirect.com/) and Cochrane Central Register of Controlled Trials (CENTRAL) (https://www.cochranelibrary.com/central). In addition, a search was performed in the following clinical trial registries: ClinicalTrials.gov (https://clinicaltrials.gov/) and the World Health Organization International Clinical Trials Registry Platform (WHO ICTRP) (https://www.who.int/ictrp/search/en/). A combination of medical subject headings and free text terms, including 'haemangioma', 'atenolol', 'propranolol', 'beta blockers', 'adverse events', 'infants', 'infantile haemangioma', 'haemangioma activity score' and 'randomized controlled trial' were searched and all publications in English language from the database's inception to July 2019 were retrieved.

The reference lists of primary trials obtained through electronic searches were also checked and relevant articles were included for review and analysis. In cases requiring clarification or additional information, the authors of the published trials were contacted.

Inclusion and exclusion criteria. For inclusion in the present analysis, studies were required to fulfill all of the following criteria: i) Parallel arm individually randomized, quasi randomized and cluster randomized controlled trials (RCTs), and prospective/retrospective cohort studies, ii) studies on patients with IH and iii) studies comparing the effectiveness of atenolol and propranolol for IH management.

All cross-over studies were excluded due to the possibility of carryover effects. Only full-text/abstract publications were included.

Outcome measures. The following outcome measures were assessed: HAS, response to medication (reduction in the lesion size), adverse events and relapse rate. Studies reporting any of the above-mentioned outcomes and that met the inclusion criteria were included.

Selection of studies. The literature search was performed by two independent investigators (CW and DS) who screened the titles, abstracts and keywords of all the retrieved citations and assessed them for possible inclusion in the present analysis. Full-text articles of relevant studies were obtained and further screening was performed independently by the primary and secondary investigators (CW and DS) to select the studies satisfying the eligibility criteria for the present analysis. Any disagreements between investigators during the selection process were resolved either through consensus or consultation with another investigator (LW). A third investigator monitored the quality of the overall review process (LW). The Preferred Reporting Items for Systematic Review and Meta-Analysis checklist was used for reporting in this review (18).

Data extraction and management. The primary investigator (CW) extracted the relevant study characteristics for review from the included studies and included general information, including the date of extraction, study title and authors; methods including the study design, participants and study setting; participant's characteristics, including the total number of participants in each arm, baseline and post-treatment outcome measures, and inclusion and exclusion criteria; intervention characteristics including details on the intervention and comparison group and follow-up duration; outcomes section, including primary and secondary outcomes, time taken for outcome assessment and other details necessary for assessing the risk of bias of included studies.

Primary and secondary investigators (CW and DS) independently extracted data associated with outcome measures from the studies included. Only extracted data from the relevant arms of studies reporting on multiple arms in a single trial were used for the present analyses. The primary investigator (CW) transferred the obtained data to the statistical software RevMan 5.3 (Cochrane). The third investigator (LW) double-checked data entries for correctness by comparing them to the data in the studies.

Risk of bias assessment for the studies included. The risk of bias of included RCTs was assessed by two independent investigators (CW and DS) using the Cochrane risk of bias tool (19). The following domains were assessed: Random sequence generation, allocation concealment, blinding of outcome assessment and study participants, incomplete outcome data, selective reporting of outcomes and other sources of bias.

For non-randomized studies, the risk of bias assessment tool for non-randomized studies (20) was used with the following domains: Selection of participants, confounding variables, intervention measures, blinding of outcome assessment, incomplete outcome data and selective outcome reporting.

For each of the above-mentioned domains, the risk of bias was graded as low (if adequate information was provided), as high (if the information was inadequate or not performed) or as unclear (if the information was missing).

Statistical analysis. For continuous outcome (HAS), the mean and standard deviation (SD) reported at baseline and follow-up were obtained. In studies where change in mean and SD scores from baseline were reported, they were extracted directly. If change scores were not reported, manual calculation was performed using the following method:

\[ \text{Mean (change)} = \text{mean (after)-mean (before)} \]

Since the data were paired, equal variances were assumed for baseline and follow-up data. \( n_1 \) and \( n_2 \) were the number of participants at baseline and follow-up, respectively, while \( s_1 \) and \( s_2 \) were the standard deviations of baseline and follow-up, respectively.

The square of the SD was multiplied with the degrees of freedom: \( (n_1-1)s_1^2 \). This was repeated for the outcome: \( (n_2-1)s_2^2 \).
The two equations were added together and divided by the total degrees of freedom:

$$s^2_p = \frac{(n_1 - 1)s_1^2 + (n_2 - 1)s_2^2}{n_1 + n_2 - 2}$$

The standard error (SE) of the difference between means was as follows:

$$SE = \sqrt{\frac{s_1^2}{n_1} + \frac{s_2^2}{n_2}}$$

SD (change) was calculated with the equation: SD=SE x √(sample size). Mean (change) and SD (change) of both groups were then entered into meta-analysis software. Finally, pooled estimates were reported as Mean Difference (MD) with 95% CI.

For dichotomous variables including the response rate, adverse effects and relapse rate, the numbers of events and of participants in each group were obtained to estimate the pooled effect size in terms of odds ratios (most studies included were retrospective in nature).

Appropriate analyses were performed based on the level of randomization (either individual or clustered). No cluster randomized trials satisfying the eligibility criteria were identified and therefore, they did not require appropriate clustering adjustments. A random-effects model with inverse variance was utilized (21).

χ² tests of heterogeneity to assess inter-study variance and I² statistics were applied to quantify inconsistencies (19). Heterogeneity was classified according to I² as mild (I²<25%) moderate (I² between 25 and 75%) or substantial (I²>75%). Study-specific and pooled estimates were represented graphically through forest plots with random-effects model. None of the outcomes exhibited significant heterogeneity. Hence, subgroup analysis or meta-regression was not required for the present study.

Publication bias was not assessed, as the outcomes did not have the required number of studies (minimum of 10 studies) to assess the publication bias.

Results

Study selection. A systematic search to retrieve studies that directly compared the effectiveness of atenolol or propranolol for the management of IH from the dates of database inception until July 2019 was performed. A total of 798 citations, 383 studies retrieved from Medline, 134 from ScienceDirect, 177 from CENTRAL, 85 from Google Scholar, 12 from ClinicalTrials.gov and 7 from WHO ICTRP were identified (Fig. 1). After the first screening stage (title, abstract and keywords), 22 relevant studies were retrieved and their full texts were reviewed for eligibility. Simultaneously, the bibliographies of the studies retrieved were reviewed and 4 more relevant studies were identified. Finally, data from 8 studies with 608 participants satisfying the inclusion criteria were analysed (13,14,22-27).

Characteristics of studies included. Table I lists the characteristics of the studies analysed. Two studies were RCTs (13,27), 3 were prospective (23,25,26) and the remaining studies were retrospective studies (14,22,24). Most of the studies were performed in Asian countries [China (2 studies) (23,26) and India (1 study)] (27) and the others were performed in American and European countries. The mean age of the study participants ranged from 2 to 6.4 months in the atenolol group and that in the propranolol group ranged from 3 to 6 months. Of the 608 participants, 250 were in the atenolol cohort and 358 were in the propranolol group. The sample sizes in the studies (both groups together) ranged from 23 to 173, while the sample size in the atenolol group ranged from 7 to 82 patients and that in the propranolol group ranged from 10 to 98. Among the 8 studies included, 6 reported on response to medication (reduction in the lesion size) (13,22-26), 2 reported on HAS (14,27), 2 reported on relapse rate (13,25) and 6 reported on adverse effects following treatment (13,14,23-25,27).

Methodological quality of the studies included. Assessments of risk of bias were performed separately for RCTs and non-randomized studies (Table IIA and B, respectively). There were no patients lost to follow-up reported in any of the studies included. The two RCTs (13,27) included in the present meta-analysis had low risks of bias in almost all of the domains. Among the non-randomized studies (14,22-26), all of the studies had a low risk of bias regarding selection of participants, intervention measures, incomplete outcome data and unclear risks of selective reporting of outcomes (protocols not published), or blinding of outcome assessment (not mentioned in the studies). Furthermore, three out of six studies had high risks of bias with respect to confounding variables.

HAS. A total of two studies reported on HAS for the two groups (atenolol and propranolol) (14,27). Fig. 2 presents the pooled MD in the HAS at 0.16 (95% CI, -0.42 to 0.73). This indicates that the evidence is not conclusive to determine which method results in a greater improvement in HAS. Furthermore, no significant heterogeneity was identified in the studies included reporting on HAS (I²=11%, P=0.29).

Response to medication. Among the studies included, 6 reported on the response rate or reduction in the lesion size following intake of the medication in the two groups (atenolol and propranolol) (13,22-26). Apart from the study by Wang et al (26), all of the other studies (13,14,22-25,27) indicated that propranolol was favoured, as it had a higher response rate when compared to the atenolol group. The overall pooled odds ratio (OR) in the propranolol arm was 1.36, indicating these infants had a 1.36 times greater odds of having complete response (reduction in lesion size) following the medication than those in the atenolol group (Fig. 3). However, the confidence of this pooled estimate crossed the null value of 1 (95% CI, 0.85-2.18) and the result was not statistically significant. Furthermore, no heterogeneity among the studies reporting the response rate with I²=0% was identified. The χ² test for heterogeneity also indicated the absence of significant heterogeneity among the studies reporting on the response rate (P=0.43).

Adverse effects. A total of 6 studies reported on adverse effects following the medication in the
Figure 1. Preferred Reporting Items for Systematic Review and Meta-Analysis flow chart depicting the selection process of studies for the current review (n=8). WHO, World Health Organization.
Table I. Characteristics of the studies included (n=8).

| First author (year) | Country   | Study design            | Sample size in atenolol arm | Sample size in propranolol arm | Interventions                                                                 | Follow-up                                                                 | Mean age of the study participants in atenolol arm | Mean age of the study participants in propranolol arm | (Refs.) |
|---------------------|-----------|-------------------------|----------------------------|-------------------------------|------------------------------------------------------------------------------|---------------------------------------------------------------------------|------------------------------------------------|------------------------------------------------|---------|
| Araya et al (2014)  | Chile     | Randomized controlled trial | 13                         | 10                            | Atenolol: 1 mg/kg/day for 6 months in a single daily dose; Propranolol: 2 mg/kg/day in 3 daily doses for 6 months | Follow-up at 2 weeks, 4 weeks, and then monthly until 6 months of treatment were completed | Not given separately (mean 5.2 months) | (13)                                           |
| Ashraf (2019)       | India     | Randomized controlled trial | 20                         | 20                            | Atenolol: 1 mg/kg/day for 9 months; Propranolol: 2 mg/kg/day for 9 months     | Follow-up at monthly intervals until end of treatment at 9 months          | 3.3 months | 4.8 months (27) |
| Bayart et al (2017) | US        | Retrospective non-inferiority study | 27                         | 53                            | Atenolol: 0.5 mg/kg/day until 12-15 months of age or until complete response; Propranolol: 2 mg/kg/day until 12-15 months of age or until complete response | Baseline and post-treatment assessment done                               | 2 months | 3 months (14) |
| Dakoutrou et al (2019) | Greece | Prospective               | 26                         | 28                            | Atenolol: 0.5 mg/kg/day, increased up to 2 mg/kg; Propranolol: 2 mg/kg/day divided into two doses Both continued until macroscopic regression of lesion or no response after 1 month of initiation of treatment | Follow-up was performed 1 week after the initial check and monthly thereafter | 3.63 months | 5.95 months (25) |
| De Graff et al (2013) | Netherlands | Retrospective                | 30                         | 28                            | Atenolol: Starting dose, 0.5 mg/kg/day (once daily). After 1 week of treatment, the atenolol dosage was increased to 1 mg/kg/day Propranolol: Average dosage 2 mg/kg/day | Follow-up at 2-8 weeks (t1) and 11-24 weeks | Provided only for atenolol group (median age, 6.4 months) | (24)                                           |
| Sharma et al (2013) | Canada    | Retrospective                | 7                          | 98                            | Atenolol: 1.6 mg/kg/day Propranolol: 1.5 mg/kg/day                           | Premature, follow-up weekly; 2-12 weeks, follow-up bi-weekly; >12 weeks, follow-up every 2-3 weeks | Not given separately (mean age of total participants, 3.3 months) | (22)                                           |
| Sun et al (2018)    | China     | Prospective                | 82                         | 91                            | Both drugs until 24 weeks; dosage not given                                  | First week was followed up each day, followed by once a month. At 6 months after commencement of treatment, the response was compared between the two groups | Not specified | (23)                                           |
two groups (13,14,23-25,27). Except for the studies by Ashraf et al (27) and Dakoutrou et al (25), all other studies reported that infants taking propranolol had a greater chance of developing adverse effects following medication when compared with infants taking atenolol. The overall pooled OR in the propranolol group was 2.17, indicating these infants had a 2.17 times higher odds of developing adverse reactions following the medication than those in the atenolol group (Fig. 4). However, the confidence of this pooled estimate crossed the null value (95% CI, 0.93-5.06) and the result was not statistically significant. Furthermore, moderate heterogeneity was present among the studies reporting on adverse effects with $I^2=51\%$. The $\chi^2$ test for heterogeneity indicated the absence of significant heterogeneity among the studies reporting on adverse effects ($P=0.07$).

Relapse rate. A total of 2 studies reported on the relapse rate following successful completion of treatment for the two groups (atenolol and propranolol) (13,25). The overall pooled OR in the propranolol arm was 1.67, indicating that these infants had a 1.67 times higher odds of relapse of IH following successful completion of medication than those in the atenolol group (Fig. 5). Similar to the above, the confidence of this pooled estimate crossed the null value (95% CI, 0.44-6.41) and the result was not statistically significant. Furthermore, mild heterogeneity among the studies reporting on adverse effects with $I^2=8\%$ was obtained. The $\chi^2$ test for heterogeneity indicated an absence of significant heterogeneity among the studies reporting on adverse effects ($P=0.45$).

**Discussion**

The management of IH has varied historically and $\beta$ blockers have been the mainstay of treatment with a complete response rate of ~60% (2). Each of the $\beta$ blockers used in the present analysis has its own advantages and disadvantages. There is a lack of systematic and high-quality studies assessing the effectiveness of these $\beta$ blockers directly. Hence, the present analysis was performed to compare the efficacies of atenolol and propranolol, in terms of clinical outcomes including the HAS and response rate, adverse effects and relapse rate among infants with haemangioma. The best possible evidence available to date was compiled in order to compare these medications.

A total of 8 studies comprising 608 participants were selected for inclusion in the present analysis. Of these, only 2 studies were RCTs and 3 were prospective studies, while the remainder were retrospective in nature. Most of the studies had either low or unclear bias risks. No substantial heterogeneity was identified among the reported outcomes in the studies. Hence, subgroup analysis or meta-regression was not performed to explore the source of heterogeneity. Except for the response to medication, all other outcomes (HAS, adverse reactions and relapse rate) were better for the atenolol group than the propranolol group. However, no conclusive or significant evidence for any of these outcomes was obtained, as the confidence limit crossed the null value in all of the outcomes assessed. The results from this analysis suggested
that atenolol may be non-inferior to propranolol treatment in the management of IH. In almost all of the studies, atenolol was used at a dose of 0.5-1 mg/kg/day and propranolol at a dose of 2 mg/kg/day. However, the optimal dose of atenolol and propranolol remains to be determined, as there is a lack of dose-response studies.

The major strengths of the present study include the comprehensive literature search and the broad search strategy to include all of the relevant up-to-date publications. To the best of our knowledge, the present study was the first review directly comparing the clinical outcomes and adverse reaction profile between atenolol and propranolol for the management of IH.

Table II. Risk of bias assessment.

| Study                        | Random sequence generation | Allocation concealment | Blinding of the participants, outcome assessment | Incomplete outcome data | Selective reporting of outcomes | Other risk of bias (Refs.) |
|------------------------------|----------------------------|------------------------|-------------------------------------------------|-------------------------|-------------------------------|----------------------------|
| A, Randomized studies (n=2)  |                            |                        |                                                 |                         |                               |                           |
| Araya et al (2014)           | Low risk                   | Low risk               | Low risk                                        | Low risk                | Unclear risk                   | Low risk                   | (13)                      |
| Ashraf et al (2019)          | Low risk                   | Low risk               | Low risk                                        | Low risk                | Unclear risk                   | Low risk                   | (27)                      |

B, Non-randomized studies (n=6)

| Study                        | Selection of participants | Confounding variables | Intervention measures | Blinding of the outcome assessment | Incomplete outcome data | Selective reporting of outcomes | Other risk of bias (Refs.) |
|------------------------------|----------------------------|-----------------------|-----------------------|-----------------------------------|-------------------------|-------------------------------|----------------------------|
| Bayart et al (2017)          | Low risk                   | High risk             | Low risk              | Unclear risk                      | Low risk                | Unclear risk                   | (14)                      |
| Dakoutrou et al (2019)       | Low risk                   | Low risk              | Low risk              | Unclear risk                      | Low risk                | Unclear risk                   | (25)                      |
| De Graff et al (2013)        | Low risk                   | High risk             | Low risk              | Unclear risk                      | Low risk                | Unclear risk                   | (24)                      |
| Sharma et al (2013)          | Low risk                   | High risk             | Low risk              | Unclear risk                      | Low risk                | Unclear risk                   | (22)                      |
| Sun et al (2018)             | Low risk                   | Low risk              | Low risk              | Unclear risk                      | Low risk                | Unclear risk                   | (23)                      |
| Wang et al (2016)            | Low risk                   | Low risk              | Low risk              | Unclear risk                      | Low risk                | Unclear risk                   | (26)                      |

Figure 2. Forest plot indicating the difference in haemangioma activity score between the atenolol and propranolol arm (n=2). SD, standard deviation; IV, inverse variance; df, degrees of freedom.

Figure 3. Forest plot analyzing the difference in response to medication between the atenolol and propranolol arm (n=6). SD, standard deviation; IV, inverse variance; df, degrees of freedom.
of IH. Two previous reviews (16,17) comparing propranolol with various other interventions have only included one study by Ábarzúa-Araya et al (13) to directly compare atenolol and propranolol.

The present review has certain limitations. Only 2 RCTs were included among the 8 studies. Since certain studies were retrospective in nature, no causal associations between the interventions and the outcomes can be inferred. Hence, more trials of adequate size are required to be performed to gather more evidence. It was not possible to assess publication bias, as the number of studies included in the review was <10 (minimum requirement to perform funnel plot or Egger’s test) (19). There was insufficient information to determine the optimal dose for propranolol or atenolol, the optimal schedule or factors responsible for regrowth of IH. Finally, most of the studies included in the present review were performed in high-income countries, which may limit the generalizability of the results to other geographical regions.

The results of the present study had certain implications towards clinical practice. Atenolol may be non-inferior to the propranolol treatment in the management of IH. Propranolol is widely used as a first-line drug in the management of complications. Previous evidence has indicated that propranolol has potential adverse effects on the development of the CNS of infants. It is known to negatively influence psychomotor function or the memory of infants. In addition, bronchial-associated adverse effects (e.g. bronchial hypersensitivity) have been reported propranolol users. With the current evidence, clinicians are able to use atenolol in place of propranolol depending on the patient's profile (i.e. if the infants require medication having a minimal effect on bronchus or CNS) or it may be used as an alternative if the infant on propranolol develops side effects. However, uncertainties regarding the efficacy and side effects persist, as most of the studies have an inadequate sample size that limits the power of the studies.

Apart from efficacy and safety concerns, questions regarding the dose-response association to determine the optimal dose, schedule and factors responsible for regrowth of IH following treatment require further exploration. To obtain conclusive evidence on these factors, more RCTs or prospective studies with larger sample sizes are required to strengthen the evidence for recommendations on how to best treat infants with haemangioma, as β blockers are the only FDA-approved drug for this condition.

To summarize, atenolol may be non-inferior to propranolol in the management of IH with respect to clinical outcomes and adverse reactions. However, more RCTs with larger sample sizes are required to derive conclusive evidence towards efficacy, safety and dose-response association of atenolol and propranolol.

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Availability of data and materials
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.
Authors' contributions

ZL designed the study, C Wu, DS, LW, JL, C Wang and LG were involved in literature search and data interpretation. C Wu and DS were responsible for the data analysis. ZL prepared the manuscript. LG edited the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate

Not applicable.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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