Efficacy of horizontal muscle augmentation combined inferior oblique muscle shortening for pediatric strabismus
Study Protocol
Xiu-Mei Du, MB*

Abstract
Background: This study will explore the efficacy and safety of horizontal muscle augmentation (HMA) combined inferior oblique muscle shortening (IOMS) for the treatment of pediatric strabismus (PS).

Methods: Literature search for studies will be carried out in the following databases: Cochrane Library, MEDILINE, EMBASE, CINAHL, Web of Science, PsycINFO, CBM, and CNKI. We will search all these databases without language and publication status restrictions. Two independent authors will perform selection of studies, data collection and management, risk of bias evaluation. A third author will be consulted with the help of discrepancies.

Results: This study will provide a synthesis of existed evidence for HMA combined IOMS for the treatment of PS.

Conclusion: The results of this study will provide evidence to evaluate the efficacy and safety of HMA combined IOMS for the treatment of PS, which can help to guide clinical decision-making.

Systematic review registration: PROSPERO CRD42019149716.

Abbreviations: CIs = confidence intervals, HMA = horizontal muscle augmentation, IOMS = inferior oblique muscle shortening, PS = pediatric strabismus, RCTs = randomized controlled trials.

Keywords: efficacy, horizontal muscle augmentation, inferior oblique muscle shortening, pediatric strabismus, safety

1. Introduction
Pediatric strabismus (PS) is misalignment disorder of the eyes among children population.[1–3] It has been reported that about 3% children can experience such disorder.[4] If it is untreated, about 50% children with PS may cause visual loss due to the amblyopia.[4–6] Several factors are supposed to response for PS, such as refractive error, muscle imbalance, retinoblastoma, or other serious ocular defects or diseases.[7–9] Several managements can help to treat PS, including patching or atropine drops, contact lenses or eyeglasses, eye exercises, and surgical alignments (such as horizontal muscle augmentation (HMA), inferior oblique muscle shortening (IOMS)).[10–21] Previous studies have reported that HMA and IOMS can benefit children with PS.[22–31] However, no studies have addressed this topic systematically. Thus, this study will systematically assess the efficacy and safety of HMA and IOMS for the treatment of PS.

2. Methods

2.1. Dissemination and ethics
This study is a literature-based study; therefore, no ethical approval is required. This study is expected to be published at a peer-reviewed journal.

2.2. Study registration
This study protocol has been registered on PROSPERO CRD42019149716. We report this study based on the Cochrane Handbook for Systematic Reviews of Interventions and the Preferred Reporting Items for Systematic Reviews and Meta-Analysis Protocol statement guidelines.[32]

2.3. Inclusion and exclusion criteria
2.3.1. Types of studies. Any randomized controlled trials (RCTs) of HMA combined IOMS for the treatment of PS. We will exclude non-RCTs and quasi-RCTs.

2.3.2. Types of participants. Participants diagnosed with PS will be included without restrictions of ethnicity, gender, and age.

2.3.3. Types of interventions. The experimental group has used HMA combined IOMS for the treatment.

This study has supported by the Shaanxi Provincial Science and Technology Department (2018GXX-091).

The authors have no conflicts of interest to disclose.

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How to cite this article: Du XM. Efficacy of horizontal muscle augmentation combined inferior oblique muscle shortening for pediatric strabismus. Medicine 2019;98:46(e17941).
Received: 15 October 2019 / Accepted: 16 October 2019
http://dx.doi.org/10.1097/MD.0000000000017941
The control group has received any interventions, except the single therapy of HMA, IOMS, or combination of HMA and IOMS.

2.3.4. Types of outcome measurements. The primary outcome includes the incidence of postoperative nausea and vomiting. The secondary outcomes consist of pain intensity (as measured by any pain scores), incidence of occludocardiac reflex, incidence of wound healing, incidence of wound infection, quality of life (as measured by any related scales), and adverse events.

2.4. Search strategy
2.4.1. Electronic databases searching. We will systematically search Cochrane Library, MEDILINE, EMBASE, CINAHL, Web of Science, PsycINFO, CBM, and CNKI without language, time of publication, and publication status restrictions. All electronic databases will be searched from inceptions to the present. A detailed search strategy for Cochrane Library will be showed in Table 1. We will also build identical search strategy to the other electronic databases.

2.4.2. Other resources searching. We will also search dissertations, ongoing studies, clinical registry, and reference list of qualified studies to avoid missing any potential studies.

2.5. Data synthesis and statistical analysis
2.5.1. Study selection. Two authors will independently identify titles and abstracts of all literatures through the search strategy to check eligible studies. Any different opinions will be solved through discussion with a third author invited. We will exclude any irrelevant studies after initial screen, and full texts of remaining studies will be carefully read to further check their eligibility criteria. The process of study selection will be presented in diagram chart.

2.5.2. Data collection and management. We will use predefined data acquisition sheet to extract data from each eligible study by 2 independent authors. Any discrepancies noticed in the process of data collection will be solved via another author. The following extracted information includes study characteristics (such as time of publication, title, first author, et al); patient characteristics (such as sample size, gender, age, race, et al); study methods (such as randomization, blinding, allocation, et al); treatment details (such as duration, frequency, et al); outcomes (such as primary and secondary outcomes, et al); and safety.

2.5.3. Dealing with missing data. When the essential missing information occurs, primary authors will be contacted to require that. We will analyze available data only when that information is not achievable.

2.5.4. Risk of bias assessment. Two authors will independently evaluate the study quality by using Cochrane risk of bias. This tool includes selection bias, performance bias, detection bias, attrition bias, reporting bias, and other possible sources. Each one is further graded as low, unclear or high risk of bias. If there are inconsistent results occur between 2 authors, a third author will help to make final decision through discussion.

2.5.5. Measurement of treatment effect. For dichotomous outcome data, we will utilize risk ratio along with its 95% confidence intervals, whereas continuous data will be calculated as mean difference or standardized mean difference along with 95% confidence intervals.

2.5.6. Assessment of heterogeneity. Heterogeneity among included RCTs will be checked using $I^2$ test. The study is considered as low heterogeneity if $I^2 \leq 50\%$. Meanwhile, a meta-analysis will be conducted based on the same treatments, and outcome measurements. We will apply a random-effects model to pool the data if substantial heterogeneity exists among eligible RCTs ($I^2 > 50\%$). At the same, we will perform subgroup analysis, and meta-regression test to explore the possible reasons for the high heterogeneity.

2.5.7. Data synthesis. We will use RevMan 5.3 software to perform statistical analysis. We will use a fixed-effects model to pool the data if low heterogeneity occurs among included RCTs ($I^2 \leq 50\%$). Meanwhile, a meta-analysis will be conducted based on the same treatments, and outcome measurements. We will apply a random-effects model to pool the data if substantial heterogeneity exists among eligible RCTs ($I^2 > 50\%$). At the same, we will perform subgroup analysis, and meta-regression test to explore the possible reasons for the high heterogeneity.

2.5.8. Publication bias. We will use Funnel plot and Egger regression test to visually inspect reporting bias if a sufficient number of included RCTs (more than 10 studies) are available.[33]

2.5.9. Subgroup analysis. We will employ subgroup analysis to explore the source of heterogeneity according to the different treatments, comparators, and outcomes.

| Table 1 | Search strategy for Cochrane Library database. |
|---------|-----------------------------------------------|
| Number  | Search terms                                           |
| 1       | MeSH descriptor: (pediatrics) explode all trees     |
| 2       | MeSH descriptor: (strabismus) explode all trees     |
| 3       | MeSH descriptor: (children) or (cross-eyed) or (wall-eyed) or (visual problem) or (eye disorder) or (pediatrics) or (general surgery) or (operative) or (horizontal muscle augmentation) or (inferior oblique muscle shortening) |
| 4       | Or 1-3                                               |
| 5       | MeSH descriptor: (surgical procedures, operative) explode all trees |
| 6       | MeSH descriptor: (general surgery) explode all trees |
| 7       | (surgery) or (general) or (procedures) or (operative) or (operative surgical procedures) or (general) or (operation) or (horizontal muscle augmentation) |
| 8       | Or 5-7                                               |
| 9       | MeSH descriptor: (randomized controlled trials) explode all trees |
| 10      | (random) or (blind) or (allocation) or (placebo) or (control) or (clinical trials) or (controlled trials) |
| 11      | Or 9-10                                              |
| 12      | 4 and 8 and 11                                       |

7 ((surgery) or (general) or (procedures) or (operative) or (operative surgical procedures) or (general) or (operation) or (horizontal muscle augmentation) or (inferior oblique muscle shortening))

8 Or 5-7
9 MeSH descriptor: (randomized controlled trials) explode all trees
10 (random) or (blind) or (allocation) or (placebo) or (control) or (clinical trials) or (controlled trials)
11 Or 9-10
12 4 and 8 and 11
2.5.10. Sensitivity analysis. Sensitivity analysis will be carried out to evaluate the robustness and satiability of pooled outcomes according to the different methodological quality.

3. Discussion

Although previous studies have reported that HMA and IOMS can benefit patients with PS,[22–1] there is still no systematic study to investigate its efficacy and safety for the treatment of patients with PS. Therefore, this study will firstly assess the efficacy and safety of HMA and IOMS for PS comprehensively and systematically. We will summarize the current evidence on the efficacy and safety of HMA and IOMS compared to other controls in the treatment of PS. The findings of this study will provide helpful evidence for both clinical practice and further researches.

Author contributions

Conceptualization: Xiu-mei Du.
Data curation: Xiu-mei Du.
Formal analysis: Xiu-mei Du.
Investigation: Xiu-mei Du.
Methodology: Xiu-mei Du.
Project administration: Xiu-mei Du.
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Visualization: Xiu-mei Du.
Writing – original draft: Xiu-mei Du.
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References

[1] Donahue SP. Clinical practice. Pediatric strabismus. N Engl J Med 2007;356:1040–7.
[2] Williams AR, Conroy JM. The anesthetic management of the pediatric strabismus patient. J AAPOS 1998;2:113–5.
[3] Mills MD. Perianesthesia care of adult and pediatric strabismus surgery patients. J Perianesth Nurs 1998;13:16–23.
[4] Chou MR, Malik ANJ, Suleman M, et al. Time trends over five decades, and recent geographical variation, in rates of childhood’s quint surgery in England. Br J Ophthalmol 2013;97:746–51.
[5] Ricci B, Coppola G, Ricci V, et al. Nationwide study of hospitalization and surgical treatment for childhood strabismus in Italy between 1999 and 2004. Int Ophthalmol 2009;29:153–6.
[6] Dombrow M, Engel HM. Rates of strabismus surgery in the United States: implications for man power needs in pediatric ophthalmology. J AAPOS 2007;11:330–5.
[7] Matta NS, Singman EL, Brubaker C, et al. Auto-objective accommodative measurements as a valid and reliable new method of pediatric, strabismus and amblyopia, vision screening. Binocul Vis Strabolog Q Simms Romano 2011;26:140–5.
[8] Kacarzan L, Kocoglob H, Yilmaz F, et al. Relation between middle ear pressure changes and postoperative nausea and vomiting in pediatric strabismus surgery. J Clin Anesth 2007;19:101–4.
[9] Weinstock SM, Flynn JT. Brief hospital admissions for pediatric strabismus surgery. Am J Ophthalmol 1975;80(3 Pt 2):523–9.
[10] Tuzcu K, Coskun M, Tuzcu EA, et al. Effectiveness of sub-Tenon’s block in pediatric strabismus surgery. Rev Bras Anestesiol 2015;65:349–52.
[11] Apivor D, Ravi PK. Ketamine and the oculocardiac reflex. Dysrhythmia in pediatric strabismus surgery: the role of intravenous atropine. Anaesthesia 1976;31:18–22.
[12] Khan AO. Pediatric strabismus. N Engl J Med 2007;356:2750.
[13] Kothary AS, Kothari M, Parabh S. A double-masked randomized trial of postoperative local anaesthetic for pain control in pediatric strabismus surgery. J AAPOS 2017;21:346–7.
[14] Khurana R, Tiwari S, Mehta R, et al. A double-masked randomized trial of postoperative local anaesthetic for pain control in pediatric strabismus surgery. J AAPOS 2017;21:346–7.
[15] Szigiato AA, Caldwell M, Buys YM, et al. Trends in pediatric strabismus surgery in the new millennium: influence of funding and perceived benefits of surgery. Can J Ophthalmol 2017;52:243–9.
[16] Ibrahim AN, Shabana T. Sub-Tenon’s injection versus paracetamol in pediatric strabismus surgery. Saudi J Anaesth 2017;11:72–6.
[17] Younus A, Zahoor A, Andrey B, et al. Comparison of topical oxybuprocaine and intravenous fentanyl in pediatric strabismus surgery. Saudi J Anaesth 2017;11:67–71.
[18] Sayed JA, F. Riad MA, M. Ali MO. Comparison of dexamethasone or intravenous fluids or combination of both on postoperative nausea, vomiting and pain in pediatric strabismus surgery. J Clin Anesth 2016;34:136–42.
[19] Ben-Menachem E, Gargi Y, Berkenstadt H, et al. Percussion pacing as management of nonresponsive asystole during pediatric strabismus surgery. J Clin Anesth 2014;26:332–4.
[20] Oh JN, Lee SY, Lee JH, et al. Effect of ketamine and midazolam on oculocardiac reflex in pediatric strabismus surgery. Korean J Anesthesiol 2013;64:300–4.
[21] Kim YH. Comparison of proseal laryngeal mask and endotracheal tube for airway safety in pediatric strabismus surgery. Saudi Med J 2012;33:914.
[22] Li Q. Clinical observation of the treatment of pediatric strabismus with horizontal muscle augmentation and lower oblique muscle cutting. J Modern Med Health Res 2019;3:10–1.
[23] Chung H. Clinical observation of horizontal muscle augmentation combined with inferior oblique muscle shortening in the treatment of pediatric strabismus. Da Doctor 2019;4:38–9.
[24] Li P, Wang H, Sun XT. Clinical analysis of horizontal muscle augmentation combined with inferior oblique muscle shortening for children with strabismus. Clin Res 2018;26:26–7.
[25] Ma YT, Jiang MS. Observation on the effect of horizontal muscle augmentation combined with inferior oblique muscle migration in the treatment of strabismus in children. Med Theory Pract 2018;31:2316–7.
[26] Yang XZ, Zhao YH. Clinical observation of horizontal muscle augmentation combined with inferior oblique muscle surgery for strabismus in children. Chin Prescrip Drugs 2018;16:136–7.
[27] Li BD, Zhang P. Clinical observation of the treatment of pediatric strabismus with horizontal muscle augmentation and lower oblique muscle surgery. Chin Med Guide 2017;15:61–2.
[28] Xiao ZG. Clinical effect of horizontal muscle augmentation and ablation of lower oblique muscle in the treatment of strabismus in children. J Clin Anesth 2017;25:148–6.
[29] Lu YB, Sun XT, Sun S. Thpeutic effect of horizontal muscle augmentation combined with inferior oblique muscle shortening in the treatment of pediatric strabismus. Hualiai Med 2017;33:22–3.
[30] Wang W. Observing the clinical efficacy and characteristics of horizontal muscle augmentation combined with inferior oblique muscle surgery for pediatric strabismus. World Med Inform Digest 2016;16:68–72.
[31] Luan L, Lu QS, Deng DY. Clinical observation of 40 cases of pediatric strabismus treated by horizontal muscle augmentation and lower oblique muscle cutting. Chin Folk Med 2016;25:58–60.
[32] Shamsee L, Moher D, Clarke M, et al. PRISMA-P Group. Preferred reporting items for systematic review and meta-analysis protocols (PRISMA-P) 2015: elaboration and explanation. BMJ 2015;349:g7647.
[33] Egger M, Davey Smith G, Schneider M, et al. Bias in meta-analysis detected by a simple, graphical test. BMJ 1997;315:629–34.