Efficacy of splint therapy for the management of temporomandibular disorders: a meta-analysis

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ABSTRACT

Temporomandibular disorders (TMD) are a group of clinical problems affecting temporomandibular joint (TMJ), myofascial muscles and other related structures. Splint therapy is the most commonly used approach to treatment of TMD, but its effectiveness is remains unclear. We therefore conducted a meta-analysis to evaluate the effectiveness of splint therapy for TMD in adults. The electronic databases PubMed, EMBASE, Cochrane Library, and ClinicalTrials.gov were searched for reports published up to March 31, 2016. Thirteen eligible studies involving 538 patients were identified. The results indicated that splint therapy increased maximal mouth opening (MMO) for patients with a MMO <45mm and reduced pain intensity measured using the visual analogue scale (VAS) for patients with TMD without specific description (TMDSD). Splint therapy also reduced the frequency of painful episodes for patients with TMJ clicking. No publication bias was observed, as determined with Egger’s test for all outcomes. On the basis of this evidence, we recommend the use of splints for the treatment and control of TMD in adults.

INTRODUCTION

Temporomandibular disorders (TMD) are a group of clinical problems affecting the temporomandibular joint (TMJ), myofascial muscles and other related structures [1]. There is currently no unified standard for the classification of TMD, but research diagnostic criteria for temporomandibular disorders (RDC/TMD) are the most commonly applied criteria [2]. The main signs and symptoms involve TMJ pain and clicking, myofascial or oral masticatory muscle pain, and abnormal jaw movement [3]. TMD constitute a major public health problem, as they are a main source of chronic oral facial pain, interfering with daily activities [4]. These disorders are also commonly associated with other symptoms affecting the head and neck region, including headache, ear-related symptoms, cervical spine dysfunction, and altered head and cervical posture [5, 6]. Notably, signs of TMD are detected in about 60-70% of the general population, yet only about one in four people with signs are actually aware of any symptoms [7]. The etiologies of TMD are not yet clear, but contributing factors include occlusal abnormalities, psychological stress, orthodontic treatment, microtrauma, poor health and nutrition, joint laxity and exogenous estrogen [8, 9].
The main treatment options for TMD include occlusal therapy [10], psychotherapy [11], physical therapy [12], medication [13], manual therapy [14], and surgery [15]. In practice, the occlusal splint is a removable appliance, usually fabricated of resin and most often designed to cover all of the occlusal and incisal surfaces of the teeth in the upper or lower jaw. Occlusal splint therapy is most commonly used clinical approach because of its ease of use, low cost, and broad indications. A previous meta-analysis addressed the effectiveness of splint therapy for TMD, but why it is effective remains unclear [16]. To further explore the clinical effectiveness of splint therapy in the management of TMD in adults, we performed the present meta-analysis to elucidate the functional properties of splint therapy by comparing the clinical effects reported in all relevant randomized controlled trials (RCTs) [17].

RESULTS

Literature search outcome

We identified 2062 publications in the electronic databases (Figure 1). Employing the selection criteria summarized in Materials and methods section, we obtained quantitative data for our meta-analysis after reading all titles, abstracts and full texts. Thirteen eligible studies [10, 18-27] from 11 publications were included in our final analysis.

Study characteristics

The 13 eligible studies included a total of 538 patients. The change in maximal mouth opening (MMO) was determined for 122 patients from 5 studies [10, 18, 20, 23], and the initial scope of MMO was categorized to three levels: less than 37 mm, 37 mm to 45 mm, and greater than 45 mm. The change in the amount of pain experienced, as reported using the visual analogue scale (VAS), was assessed in 285 patients from 6 studies [19-22, 25]. Three types of disorder were assessed: TMD without specific description (TMDSD) [21, 22, 25], osteoarthritis [20], and TMJ clicking [19]. The rate of healing from TMJ clicking was recorded in 170 patients from 4 studies [19, 26, 27]. The frequency of pain relief was evaluated in 112 patients from 2 studies [22, 24]. There were 5 studies [18, 20, 22, 23, 25] missing standard deviations (SD), which were estimated from the P-values. Table 2 describes the clinical characteristics of patients in all 13 studies.

Quality of the included studies

The risk of bias in the included studies was strictly evaluated. Details of methodological approach are presented in Table 3.

Table 1: Search strategy and picots criteria for the systematic review

| PICOS criteria | Population | Intervention | Outcomes | Study design | Search combination | Language | Electronic database | Focused question |
|----------------|------------|--------------|----------|--------------|--------------------|----------|---------------------|-----------------|
| Population     | 1) MeSH term: (temporomandibular joint disorders) OR (temporomandibular joint disc) OR (temporomandibular joint) OR (temporomandibular disorders) 2) Text word: (temporomandibular joint dis*) OR (dis*, temporomandibular) OR (disc*, temporomandibular joint) OR (joint dis*, temporomandibular) OR (TMJ disorders) OR (disorder, TMJ) OR (disorders, TMJ) OR (TMJ dis*) OR (temporomandibular disorder*) OR TMD | 3) MeSH term: splints OR (occlusal splints) 4) Text word: splint* OR (splints, occlusal) OR (occlusal splint*) OR (splint, occlusal) | 7) MeSH term: pain OR (pain measurement) 8) Text word: (maximal mouth opening) OR (MMO) OR pain OR (pain measurement) OR (visual analogue scales of pain) OR (VAS of pain) OR (healing from TMJ clicking) OR (pain relief) | 9) MeSH term: randomized controlled trials AND controlled clinical trials | 1 AND 2 AND 3 AND 4 AND 5 AND 6 AND 7 AND 8 AND 9 | English | Electronic database Medline/PubMed, EMBASE, Cochrane Central Register of Controlled Trials (CENTRAL) and Clinical Trials.gov | Is using splint therapy helpful to improve clinical outcomes in the management of temporomandibular disorders (TMD)? |
Figure 1: Summary of trial identification and selection.

2062 publications obtained from electronic databases and websites (PubMed (n=972), EMBase (n=399), Cochrane Central Register of controlled Trials (n=534), Clinicaltrials.gov (n=157))

Duplicates (n=647)

Potential relevant publications by reading titles and abstracts (n=1415)

Excluded publications based on selection criteria (n=1179)

More potential relevant publications by reading full text (n=236)

Publications excluded (n=224):
- Intervention or outcome did not strictly accord with the inclusive criteria (n=151)
- Conference abstract/Letter (n=23)
- Not available data (n=23)
- Other (n=39)

13 studies included in meta-analysis from 11 publications:
- The change of maximal mouth opening (n=5)
- The change of visual analogue scales of pain (n=6)
- The rate of healing from temporomandibular joint clicks (n=4)
- The frequency of pain relief (n=2)
Table 2: Characteristics of the included studies

| Study                  | Year | Region | Age (mean±SD) | Gender (female %) | Sample                  | Diagnostic criteria | Classification of diseases | History (month) | Intervention group | Control group | Course (month) | Follow-up (month) |
|------------------------|------|--------|---------------|-------------------|-------------------------|---------------------|--------------------------|-----------------|-------------------|---------------|----------------|------------------|
| Conti PC[18]           | 2012 | Brazil | 38.09±38.14   | 80.9%/100%        | 17/9                    | RDC/TMD             | TMD                      | NA              | splint            | Self-care     | NA             | 3                |
| Madani AS[19]          | 2011 | Iran   | 27.20±12.43   | 75%/92.8%         | 20/14                   | RDC/TMD             | TMJ clicking            | 6               | splint            | Physical      | 4/5            | NA               |
| Alpaslan C a[20]       | 2008 | Turkey | 29.8±11.82    | NA                | 22/14                   | Clinical examination | Osteoarthritis          | 18              | Hard splint       | Arthrocentesis | NA             | 6                |
| Alpaslan C b[20]       | 2008 | Turkey | 31.6±10.5     | NA                | 9/14                    | Clinical examination | Osteoarthritis          | 18              | Soft splint       | Arthrocentesis | NA             | 6                |
| Al Quran FA[21]        | 2006 | Jordon | 31.8±36       | NA                | 38/38                   | Clinical examination | TMD                      | NA              | Splint            | Control appliance | NA             | 3                |
| Ekberg E[22]           | 2003 | Sweden | 31/28         | 83.3%/90%         | 30/30                   | RDC/TMD             | TMD                      | 6               | Stabilization splint | Control appliance | 2.5             | NA               |
| Maloney G[23]          | 2002 | USA    | NA            | NA                | 10/7                    | RDC/TMD             | TMD                      | NA              | splint            | Control appliance | NA             | NA               |
| Ekberg E[24]           | 1998 | Sweden | 13.7±15-17    | 86.6%/96.6%       | 30/30                   | Clinical examination | TMD                      | 36              | splint            | Control appliance | 2.5             | 2.5              |
| Wright EF[10]          | 1995 | USA    | 34/31         | NA                | 10/10                   | NA                   | TMD                      | NA              | Soft splint       | No treatment   | 1.5             | NA               |
| Turk DC[25]            | 1993 | USA    | 35.9±13.31    | 75%/80%           | 30/20                   | NA                   | TMD                      | 2               | splint            | BF/SM         | 1.5             | 6                |
| Lundh H[26]            | 1988 | Sweden | NA            | NA                | 21/22                   | NA                   | TMJ clicking            | NA              | splint            | Unerated controls | 6               | 6                |
| Lundh H a[27]          | 1985 | Sweden | NA            | NA                | 24/23                   | NA                   | TMJ clicking            | NA              | Anterior repositioning splint | Control appliance | NA             | NA               |
| Lundh H b[27]          | 1985 | Sweden | NA            | NA                | 23/23                   | NA                   | TMJ clicking            | NA              | Flat splint       | Control appliance | NA             | NA               |

Abbreviations: RDC/TMD: research diagnostic criteria for temporomandibular disorders; TMD: temporomandibular disorders; TMJ: temporomandibular joint; BF/SM: Biofeedback-assisted relaxation and stress management; NA: Not available.

Table 3: Risk of bias in the included studies

| Study                  | Year | Random sequence generation | Allocation concealment | Blinding of participants and personnel | Blinding of outcome assessment | Incomplete outcome data | Selective reporting | Other bias |
|------------------------|------|-----------------------------|------------------------|---------------------------------------|--------------------------------|------------------------|---------------------|------------|
| Conti PC[18]           | 2012 | unclear                     | high                   | high                                  | high                          | low                    | low                 | low        |
| Madani AS[19]          | 2011 | unclear                     | high                   | high                                  | high                          | low                    | low                 | high       |
| Alpaslan C a[20]       | 2008 | unclear                     | high                   | high                                  | high                          | low                    | low                 | high       |
| Alpaslan C b[20]       | 2008 | unclear                     | high                   | high                                  | high                          | low                    | low                 | low        |
| Al Quran FA[21]        | 2006 | high                        | high                   | high                                  | high                          | low                    | low                 | high       |
| Ekberg E[22]           | 2003 | low                         | low                    | high                                  | low                           | low                    | low                 | low        |
| Maloney G[23]          | 2002 | low                         | high                   | high                                  | high                          | low                    | low                 | high       |
| Ekberg E[24]           | 1998 | low                         | low                    | low                                   | low                           | low                   | low                 | low        |
| Wright EF[10]          | 1995 | low                         | low                    | low                                   | low                           | low                   | low                 | low        |
| Turk DC[25]            | 1993 | unclear                     | high                   | high                                  | high                          | high                   | high                 | low        |
| Lundh H[26]            | 1988 | unclear                     | high                   | high                                  | high                          | high                   | high                 | low        |
| Lundh H a[27]          | 1985 | unclear                     | high                   | high                                  | low                           | low                   | low                 | high       |
| Lundh H b[27]          | 1985 | unclear                     | high                   | high                                  | low                           | low                   | low                 | low        |

Results of individual outcome variables

Changes in MMO

Comparison of the splint therapy and control groups revealed a significant difference in the change of MMO (Figure 2) (MD = 5.39, 95% CI [3.96, 6.81], I² = 48.9%, P = 0.098]). Moreover, subgroup analysis showed that for patients with an initial MMO < 37mm (MD = 6.21, 95% CI [4.50, 7.92], I² = 34.0%, P = 0.220) or an initial MMO > 45mm (MD = 2.57, 95% CI [1.71, 3.61], I² = 59.7%, P = 0.008), a significant increase in MMO was detected for the subgroups with MMO > 45mm (MD = 1.57, 95% CI [-2.22, 5.36], F = NA, P = NA). In addition, for the MMO < 37mm group, meta-regression showed there was no significant difference between the control and splint therapy groups after adjusting for differences in baseline and possible confounding factors (Table 4).

Change in pain level (VAS)

Comparison of the splint therapy and control groups also revealed a significant difference in the changes in level of perceived pain, as determined using the VAS (Figure 3) (MD = 2.02, 95% CI [1.55, 2.49], F = 0%, P = 0.558). Subgroup analysis showed that patients
Figure 2: Forest plot of the changes in MMO.

Figure 3: Forest plot of the changes in VAS for pain.
with TMDSD in the splint therapy group experienced a significant decrease in pain as compared to control (MD = 2.00, 95% CI [1.50, 2.51], \( I^2 = 34.5\% \), \( P = 0.217 \)). This was also the case for patients with TMJ clicking (MD = 2.35, 95% CI [0.89, 3.81], \( I^2 = NA \), \( P = NA \)), but not for patients with osteoarthritis (MD = 1.41, 95% CI [-1.16, 3.97], \( I^2 = 0\% \), \( P = 0.494 \)). On the other hand, meta-regression revealed no significant differences in the change in VAS for pain after adjusting for baseline and possible confounding factors (Table 4).

### Rate of healing from TMJ clicking

Comparison of the splint therapy and control groups using a fixed-effects model showed that there was no significant difference in the rates of healing from TMJ clicking between the two groups (Figure 4) (RR = 1.17, 95% CI [0.69, 1.98], \( I^2 = 0\% \), \( P = 0.701 \)).

### Frequency of pain relief

Comparison of the splint therapy and control groups using a fixed-effects model showed that splint therapy significantly reduced the frequency of painful episodes in patients with TMJ clicking (Figure 5) (RR = 1.90, 95% CI [1.19, 3.02], \( I^2 = 0\% \), \( P = 0.442 \)).

### Publication Bias

The result of Egger’s test showed there was no significant difference between both the change in MMO (Bias = -1.915 [-4.50, 0.67], \( P = 0.100 \)) and the change in the VAS for pain (Bias = -0.072 [-2.13, 1.98], \( P = 0.927 \)).

### DISCUSSION

In this meta-analysis, we evaluated 13 clinical trials that included TMD patients older than 18 years of age. We found that patients with an initial MMO less than 45mm showed a statistically significant change with splint therapy, whereas patients with an initial MMO greater than 45mm did not. Furthermore, subgroup analysis confirmed that splint therapy was most effective with patients exhibiting a limited ability to open their mouths. It also appears splint therapy provides significant pain relief to patients with TMDSD and TMJ clicking, but splint therapy did not reduce the pain in patients with osteoarthritis. We speculate that this is because osteoarthritis occurs after damage to the tissue has already occurred, whereas TMDSD and TMJ clicking occur in the early stage of the disease, when the splint is more able to exert a beneficial effect. In addition, although we found that splint therapy

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**Table 4: Meta-regression results for the main outcomes: VAS for pain and MMO <37 mm**

| Confounding factors | MMO <37mm | VAS of pain |
|---------------------|-----------|-------------|
|                     | Number of study | Coef 95%CI | P  | Number of study | Coef 95%CI | P  |
| Age                 | 3          | 0.468(-1.941, 2.878) | 0.703 | 6          | -0.027 (-0.215, 0.162) | 0.782 |
| Gender              | 3          | NA          | NA  | 6          | 0.103 (-0.027, 0.234) | 0.12  |
| Region (Ref=Europe) | 0          | NA          | NA  | 1          | NA          | NA  |
| North America       | 1          | NA          | NA  | 1          | -0.300 (-2.969, 2.369) | 0.826 |
| Other               | 2          | -5.791(-12.409, 0.826) | 0.086 | 4          | 0.517 (-2.146, 3.179) | 0.704 |
| Diagnostic criteria (Ref=Clinical examination) | 2 | NA | NA | 3 | NA | NA |
| RDC                 | 1          | 5.791(-0.826, 12.409) | 0.086 | 2 | -0.193 (-1.664, 1.279) | 0.797 |
| Other               | 0          | NA          | NA  | 1          | -0.834 (-1.853, 0.185) | 0.109 |
| Classification of diseases (Ref=Osteoarthritis) | 2 | NA | NA | 2 | NA | NA |
| TMD                 | 1          | 5.791(-0.826, 12.409) | 0.086 | 3 | 0.615 (-2.130, 3.360) | 0.66  |
| TMJ clicking        | 0          | NA          | NA  | 1          | 0.935 (-2.196, 4.066) | 0.558 |
| Course              | 3          | NA          | NA  | 6          | -0.061 (-0.319, 0.197) | 0.643 |
| Follow-up           | 3          | -3.217(-6.894, 0.459) | 0.086 | 6 | -0.318 (-0.659, 0.024) | 0.069 |
| Sample              | 3          | -0.377(-0.0820, 0.067) | 0.096 | 6 | 0.019 (-0.014, 0.052) | 0.25  |
| Publish year        | 3          | -0.965(-2.068, 0.138) | 0.086 | 6 | 0.053 (-0.015, 0.122) | 0.131 |
| Sources of SD (Ref=Reported SD) | 1 | NA | NA | 2 | NA | NA |
| Estimate the SD from P value | 2 | 2.733(-10.521, 15.988) | 0.686 | 4 | -0.883 (-1.828, 0.062) | 0.067 |

Abbreviations: VAS: Visual analogue scales, MMO: Maximal mouth opening, RDC: Research diagnostic criteria, TMD: Temporomandibular disorders, TMJ: Temporomandibular joint, SD: Standard deviation, CI: Confidence interval, Coef: Coefficient, Ref: Reference, NA: Not available.
had no significant beneficial effect on the symptoms of TMJ clicking, the meta-analysis showed that splint therapy reduced the frequency of TMJ pain.

There are many types of splints for the treatment or control of TMD. These include the stabilization splint, soft splint, flat splint, and pivot splint [11]. Although the mechanisms of action of splints are not fully understood, Aleksandra et al [28] found that plasma levels of calcitonin gene related peptide (CGRP) were increased in TMD patients treated with an occlusal splint. CGRP is a potent neuropeptide thought to play an essential role in stretching and relaxing muscle, neurogenic vasodilatation

![Figure 4: Forest plot of the rate of healing of TMJ clicking.](image)

![Figure 5: Forest plot of the frequency of pain relief.](image)
and maintaining the functional integrity of peripheral tissues [29]. We therefore suggest that the increased CGRP reflects the decreased activity muscles responsible for MMO. In addition, Glaros et al [30] proposed that splints decrease pain by preventing tooth contact and reducing muscle tension. Seligman et al [31] suggested that function occlusal relationships reflect the balance of working occlusal contacts, length and symmetry of retruded contact position-intercuspal position (RCP-ICP) slides, occlusal guidance patterns, parafunction, and dental attrition. All these interacting factors play important roles during splint therapy, which explains in part why combination therapies are more effective for the treatment of TMD [32].

To our knowledge, four meta-analyses examining effectiveness of splint therapy have been published [11, 33-35]. Two of those studies [11, 34] indicated that splints reduce pain in patients with TMD. Ebrahim et al [34] included 11 studies involving 455 patients. Two outcomes, VAS for pain and the incidence of continued pain, were described without subgroup analysis. Fricton et al [11] only included the rate of pain reduction among 50 patients self-reporting pain, and Al-Ani et al [35] only included pain as an outcome. MMO was not examined in any of those studies. In the present meta-analysis, we included 13 studies and used two main outcomes (VAS for pain and MMO) and two secondary outcomes (rate of healing from TMJ clicking and frequency of pain) [36]. Moreover, comprehensive subgroup analyses were conducted based on the range of initial MMO and subclasses of VAS for pain. The clinical heterogeneity was relieved, and the results were largely consistent with the experience in clinical practice [37].

There are several limitations to this study that should be addressed. First, only a few clinical trials met the inclusion and exclusion criteria. Consequently, more clinical studies will be required to confirm our results [38]. Second, some of the clinical trials had missing data on basic characteristics, possibly falsely increasing heterogeneity due to failure to perform a meta-regression for confounding factors [39, 40]. Although we estimated the missing SD from P-values, this can lead to errors [41]. Finally, although all included studies were randomized controlled trials or parallel-group design clinical trials, we could not implement complete allocation concealment, blinding the participants and personnel to the outcome assessment [42, 43].

CONCLUSIONS

This study examined the effectiveness of splint therapy in TMD patients in a meta-analysis of published results. Our results indicate that splint therapy effectively reduces pain levels in TMJSD patients, and reduces the frequency of pain inpatients with TMJ clicking. Additionally, splint therapy increased mouth opening ability in patients with initial MMO < 45mm. On the basis of this evidence, we recommend the use of splints for the treatment and control of TMD in adults.

MATERIALS AND METHODS

This meta-analysis was conducted according to the Preferred Reporting Items for Systematic Reviews and Meta-analyses (PRISMA) statement [44]. No ethical issues were involved in this study, and all collected data were based on published studies.

Literature search strategy

We conducted a search of four electronic databases, PubMed, EMBASE, the Cochrane Central Register of Controlled Trials and Clinical Trials.gov, up to March 31, 2016 for eligible randomized or parallel-group design clinical trials that evaluated the effectiveness of splint therapy in patients suffering from the TMD. The electronic search and the PICO (population, intervention, comparator, outcomes) strategy are shown in Table 1.

Selection criteria

All studies were selected in accordance with the following inclusion criteria: 1) RCTs; 2) included only TMD patients older than 18 years; 3) compared the effectiveness of splint therapy using controls receiving no treatment or placebo; 4) included only patients who should have been diagnosed with TMD (e.g., osteoarthritis, TMJ clicking or anterior disc displacement with or without reduction); 5) included patients who had not been administered a TMD treatment prior to the study; and 6) investigated one of the following outcomes: i) changes in MMO without support, ii) changes in VAS for pain, iii) rate of healing from TMJ clicking or iv) change in the frequency of pain from more than once a week to less than once a week. The main outcomes in this study were defined as a change of MMO and change in VAS for pain. The secondary outcomes were the rate of healing from TMJ clicking and the change in the frequency of pain.

Exclusion criteria

Studies were excluded based on the following criteria: 1) pain at rest was used as the pain score; 2) the study was a duplicate; 3) the data could not be extracted or obtained through contact with the author; and 4) too little information to calculate the missing SD.

Data extraction

The relevant information, including study design, patient characteristics, interventions, comparisons, and
outcomes, were independently extracted and entered into a database by two investigators. When relevant research information was missing, particularly study design or outcome information, we contacted the original authors for clarification. The following information was extracted from each study: publication year, region, age, gender, sample, diagnostic criteria, classification of diseases, history, intervention and control groups, course, follow-up, and outcomes. Disagreements between the two investigators on data extraction or quality assessment were resolved by discussion. If the dispute persisted, other senior investigators were consulted to attain consensus.

Quality assessment of included studies

Two investigators independently evaluated the methodological quality of eligible trials using the Cochrane collaboration tool [38] for assessing risk of bias (random sequence generation, allocation concealment, blinding of participants and personnel, blinding of outcome assessment, incomplete outcome data, selective reporting and other sources of bias).

Statistical analysis

To describe the main outcomes based on continuous data, we used weighted mean differences (MD) [38], and 95% confidence intervals (CI). For the secondary outcomes, based on dichotomous data, we used relative risk (RR) [38, 45] and 95% CI. All the outcome data were processed using STATA 14.0 software. All missing SD were estimated from P-values [41]. We performed a statistical test for heterogeneity [37] and adopted I² > 50% and P≥0.1 as evidence for heterogeneity [38]. If the data were homogeneous under a fixed-effects model, the initial scope of the MMO and disease classification were identified as key sources of heterogeneity in the main outcomes [37]. Heterogeneity was then dealt with using subgroups based on these modifiers. If the data were still heterogeneous, we introduced a random-effects model [37]. In addition, the baseline and possible confounding factors, including age, gender, region, diagnostic criteria, classification of diseases, course, follow-up, sample, publish year, and sources of SD, were detected using meta-regression [46]. Finally, the Egger’s test was employed to address quantitative detection bias [47].

CONFLICTS OF INTEREST

The authors declare no conflict of interest.

Author contributors

CZ and MHD had full access to all of the data in the study, and took responsibility for the integrity of the data and the accuracy of the data analysis. JYW, YMN, and CZ designed the study. MHD and YT developed and tested the data collection forms. JYW, YMN, and DLD acquired the data. CZ and BYH conducted the analysis and interpreted the data. JYW and YMN drafted the manuscript. All authors critically revised the manuscript. MHD and CZ are guarantors.

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