Research paper

Detachable string magnetically controlled capsule endoscopy for detecting high-risk varices in compensated advanced chronic liver disease (CHESS1801): A prospective multicenter study

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A B S T R A C T

Background: Gastroesophageal varices is a serious complication of compensated advanced chronic liver disease (cACLD). Primary prophylaxis to reduce the risk of variceal hemorrhage is recommended if high-risk varices (HRV) are detected. We performed this study to compare the accuracy, patients’ satisfaction and safety of detection of HRV by detachable string magnetically controlled capsule endoscopy (DS-MCCE) with esophagogastroduodenoscopy (EGD) as the reference.

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Methods: We prospectively recruited participants with cACLD from 12 university hospitals (11 in China and one in the United Kingdom) between November 2018 and December 2019 (ClinicalTrials.gov, NCT03749954). All participants underwent DS-MCCE, followed by EGD within a week in a blinded fashion. Following endoscopy, and on the same day, participants were asked to fill in a satisfaction questionnaire regarding their experience.

Findings: A total of 105 eligible participants were enrolled. With EGD as the reference standard, the concordance index, sensitivity, specificity, positive predictive value, negative predictive value, positive likelihood ratio, and negative likelihood ratio of DS-MCCE in diagnosis of HRV were 0.90 (95% confidence interval [CI]: 0.83–0.95), 92% (95% CI: 78–98%), 88% (95% CI: 78–95%), 80% (95% CI: 70–92%), 95% (95% CI: 90–100%), 7.91 (95% CI: 4.10–15.30), and 0.09 (95% CI: 0.03–0.30), respectively. The kappa score of 0.78 (95% CI: 0.65–0.90) suggested substantial agreement between DS-MCCE and EGD. Moreover, in participants undergoing EGD without sedation, the satisfaction of DS-MCCE was significantly better than that of EGD ($p < 0.0001$, $d = 1.15$ [95%CI: 0.48–1.42]). All participants confirmed the excretion of the capsule, and no adverse events occurred.

Interpretation: DS-MCCE is an accurate alternative to EGD for detecting HRV in cACLD, which is safe and associated with better satisfaction.

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Introduction

The presence of gastroesophageal varices (GEV) is a common and serious complication of compensated advanced chronic liver disease (cACLD) [1-3]. GEV hemorrhage is associated with a six-week mortality rate of between 15% and 25% [2,3]. In order to prevent variceal hemorrhage, screening and surveillance aims to detect high-risk varices (HRV) and determine the need for primary preventative therapy [2,3]. Esophagogastroduodenoscopy (EGD) is therefore an important part of the diagnostic work-up in patients with cACLD, serving as the gold standard to diagnose HRV [2,3]. However, EGD is invasive and poorly tolerated, with many patients needing intravenous sedatives or general anesthesia. Although EGD with sedation relieves patients’ anxiety and discomfort and reduces the potential for physical injury during the procedure, it incurs additional risks of cardiopulmonary adverse events [4]. Consequently, patients may decline a screening procedure if they are stable and asymptomatic.

Non-invasive methods for detection of cACLD are being explored [5-13]. Although preliminary research is encouraging, these techniques predict the presence, rather than confirm or assess the size, of GEV. By contrast, capsule endoscopy is a non-invasive alternative which also allows direct visualization of GEV [14].

A number of well-conducted trials [15-19] suggest that EGD and capsule endoscopy may be equivalent in terms of accuracy in the identification and grading of varices. However, concerns remain that the quality of examination may be adversely affected by the uncontrolled and sometimes rapid transit of the capsule through the esophagus [20]. String capsule endoscopy was introduced to address this concern by providing control of capsule movement as well as real-time visualization [21]. However, this technique is limited by the inability to detach the string from the capsule. Thus, observation of the fundus, one of the predilection sites of varices, is impossible and retrieval of the capsule from the esophagus causes discomfort.

Magnetically controlled capsule endoscopy (MCCE), a novel modality, was developed and approved by the China Food and Drug Administration in 2013 [22]. Our previous studies initially demonstrated that MCCE was comparable in accuracy to EGD for gastric examination [22,23]. Furthermore, it has several strengths including non-invasiveness, no sedation requirement, and easy op-
eration [22,23]. We have since combined an innovative detachable string system with the MCCE (detachable string magnetically controlled capsule endoscopy (DS-MCCE)) and carried out a pilot study showing that DS-MCCE was safe and feasible both in healthy volunteers and patients with suspected esophageal disease [24]. Moreover, successful detachment of the capsule from the string avoids the discomfort of retrieving the entire capsule from the mouth and allows subsequent investigation of the gastric cardia and fundus. To our knowledge, the diagnostic accuracy of HRV, comfort and safety of DS-MCCE in patients with cAUCD have not been explored in a large-scale trial. This prospective, multicenter study aimed to assess the accuracy, patient's satisfaction, and safety of DS-MCCE for detecting HRV in well-characterized patients with cAUCD.

Methods

Study design and participants

This study (CHESS1801 trial) was a multicenter blinded comparison trial (ClinicalTrials.gov identifier: NCT03749954), in which we prospectively recruited well-defined participants with cAUCD from 12 university hospitals (11 in China and one in the United Kingdom) between November 2018 and December 2019. Inclusion criteria were: 1) clinically evident or biopsy-confirmed cAUCD; 2) a schedule to undergo an EGD; 3) age between 18 and 75 years; 4) a life expectancy of at least 24 months without liver transplantation; 5) Model for End Stage Liver Disease score of 29 or less; 6) written informed consent. Participants with any contraindications to MCCE were excluded [22]. The study protocol conformed to the ethical guidelines of the Declaration of Helsinki, and was approved by the institutional review board of all institutions.

Procedure

The DS-MCCE system (Ankon Technologies Co, Ltd, Wuhan, China) consists of a thin latex sleeve attached to a hollow string, an endoscopic capsule, a magnetic guidance robot with five-degree-of-freedom (two rotational and three translational) robotic arm, a data recorder, and a computer workstation with the NaviCam® magnetic capsule guidance software for real-time viewing and control. The string is 80 cm in length, attached to the sleeve on the caudal end of the capsule, and a sterile syringe for single use. The capsule is 27 mm in length and 11-8 mm in diameter, with a permanent magnet inside its dome and a camera at one end (Figure S1A). It is partially enclosed within the sleeve and can be separated from the string system by using the syringe to inject air into the hollow string after exploring the whole esophagus. Images are captured at a rate of two frames per second. The angle of view is up to 140 degree. A CMOS image sensor is used in DS-MCCE. The LED light exposure time and signal gain of CMOS sensor are automatically adjusted by measuring the histogram of the image to optimize brightness and contrast of the images. The operator uses two joysticks to control capsule movement by varying the strength of the magnetic field (by altering the distance of the magnet from the patient) and the polarity of the magnet (Figure S1B). Recording and downloading data are similar to other capsule endoscopies.

An overnight fast (> 8 h) was needed before the DS-MCCE examination. At 30 min before examination, study participants ingested 2-5 g of dimethicone powder (Honghe Medicine, Zigong, China) dissolved in 50 mL of water or 400 mg of simethicone suspension (Espumisan, Berlin-Chemie, Germany) dissolved in 50 mL of water as a defoaming agent, followed by 500–1000 mL of water to fill the stomach cavity to provide a better view. Participants were positioned in the left lateral decubitus and swallowed the capsules with a small amount of water. The capsule was permitted to move passively as far as the cardia, from where the string was slowly pulled up to inspect the esophagus in real time. Regions of interest could be observed repeatedly. When the capsule reached the stomach after the esophageal observation, the string could be detached from the capsule by injecting 5–10 mL of air using the syringe. The string was then removed from the mouth and discarded. The capsule was controlled magnetically to observe the whole stomach surface area. A common protocol involved observation in the order of the fundus, cardiac regions, body, anulus, antrum, and pylorus, with the patients' body position changing from left lateral, supine to right lateral [22]. Meanwhile, water ingestion was repeated if there was insufficient gastric distention. All participants were followed during two weeks to check for capsule excretion and for any adverse events. DS-MCCE was performed by a dedicated operator at each center. All operators had been trained in a standardized manner before the enrollment.

EGD examinations were performed by experienced endoscopists using conventional standard forward-viewing upper gastrointestinal video endoscopes at individual centers and were blinded to the DS-MCCE findings. EGD was performed within a week of capsule ingestion, either without or with sedation according to the standard procedure of the center and to the preference of the patient. All major findings were recorded on digital pictures and then reported as the EGD results.

Conventional non-invasive models including image-based indexes including portal venous velocity, portal diameter, liver stiffness (LS), and LS-spleen size-to-platelet ratio score (LSPS), [13] and serum-based parameters including serum aspartate aminotransferase to alanine aminotransferase ratio (AAR), [6] aspartate aminotransferase to platelet count ratio index (APRI), [7] fibrosis index based on four factors (FIB-4), [8] Fibrosis Index, [9] gamma glutamyl transpeptidase to platelet count ratio (GPR), [10] King’s score, [11] and Lok score [12] were collected. LS was measured by transient elastography (FibroScan, Echosens, France; or FibroTouch, Hisky, China). Portal diameter, portal venous velocity, and spleen size were measured with Doppler ultrasound. Calculation formulas for LSPS, AAR, APRI, FIB-4, Fibrosis Index, GPR, King’s score, and Lok score were summarized in Table S1.

Outcomes

The primary outcome was the diagnostic performance of DS-MCCE for HRV, identified by either large varices (varices size > 5 mm) or small varices with presence of red signs according to the Baveno VI consensus [25]. In this study, EGD served as the gold standard for HRV. Two independent experienced interpreters, blinded to the patients’ EGD results, reviewed the coded images of all participants captured by DS-MCCE and rendered their opinions as to whether the patients had evidence of HRV. When there was a disagreement, a decision was made by a third independent reader. As for EGD results, saved pictures of all participants were transferred to two independent and experienced endoscopists who were blinded to each patient’s history for evaluation. In the event of discordant judgements, the opinion of a third independent interpreter was sought.

The secondary outcomes included patients’ satisfaction and safety of DS-MCCE. A satisfaction questionnaire (Appendix) regarding patients’ preoperative perception and postoperative satisfaction was developed referring to a previous research [15]. The questionnaire was submitted to a panel comprising three professionals with expertise in gastroenterology, outcome measurement or psychology for evaluation before the questionnaire was put into the trial. Participants were asked to fill in a satisfaction questionnaire after each procedure (DS-MCCE and EGD). The questionnaire consisted...
of 11 questions with total score ranged from 0 (poor) to 41 (excellent).

### Statistical analysis

The sample size calculation was based on a reported sensitivity, specificity, and prevalence of HRV in capsule endoscopy of 78%, 96%, and 27%, respectively [15]. A sample size of 101 participants was calculated by PASS (version 11.0, NCSS, 2012) to be required to provide a power of 90% at a statistic significance level of 0.05, allowing for a 5% dropout rate.

Categorical data were expressed as numbers (percentages), and continuous variables were expressed as mean (standard deviation) or median (interquartile range). With EGD as the reference standard, the diagnostic performance for detecting HRV of DS-MCCE was assessed by concordance index (C-index), sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV), positive likelihood ratio (+LR), and negative likelihood ratio (-LR). The beyond-chance agreement also known as the kappa score, between the DS-MCCE and the EGD was also calculated. Kappa values between 0–40 and 0–60 were considered of moderate agreement whereas values between 0–60 and 0–80 were substantial [26]. Diagnostic performance of other conventional non-invasive models were compared using the area under the receiver operating characteristic curve (AUC), sensitivity, specificity, PPV, and NPV. Comparison of satisfaction and safety between DS-MCCE and EGD was tested using a student’s t-test on paired data. A p value of < 0.05 was considered significant. An effect size (d value) > 0.80 was considered high effect. All statistical calculations were performed with R language (version 3.6.2, R Core Team, 2019).

### Role of the funding source

The funders had no role in study design; the collection, analysis, and interpretation of data; writing of the report; or the decision to submit for publication. The corresponding author had full access to all of the data and the final responsibility to submit for publication.

### Results

Between November 9, 2018, and December 20, 2019, a total of 113 consecutive eligible participants with well-characterized cA-CLD from 12 university hospitals in China and the United Kingdom were involved in this prospective trial. Among them, two participants were excluded because of failure to swallow the capsule. In addition, six were excluded during evaluation due to insufficient recordings of EGD (n = 5) and DS-MCCE (n = 1). Therefore, 105 eligible participants were enrolled in the final analysis (Fig. 1). The most common cause of cA-CLD was hepatitis B virus infection (66, 63%). Baseline characteristics of the enrolled participants were summarized in Table 1.

DS-MCCE operation procedure and representatives of high-risk and low-risk varices sampled by DS-MCCE and EGD were illustrated in Fig. 2. EGD detected HRV in 36 (34%) patients and DS-MCCE identified HRV in 33 (31%) of these. In eight cases, the diagnosis of HRV by the DS-MCCE was not confirmed by EGD. The C-index, specificity, PPV, NPV, +LR, -LR of DS-MCCE for detection of HRV were 0.90 (95% confidence interval [CI]: 0.83–0.95), 92% (95% CI: 78–98%), 89% (95% CI: 78–95%), 80% (95% CI: 70–92%), 95% (95% CI: 90–100%), 7–91 (95% CI: 4–10–153), and 0–09 (95% CI: 0–03–039), respectively. The kappa score between EGD and DS-MCCE on HRV diagnosis was 0.78 (95% CI: 0–65–0–90), demonstrating a substantial agreement.

Compared to conventional non-invasive tools, DS-MCCE showed the best diagnostic performance with an AUC of 0.90 (95% CI: 0–83–0–95). As for the image-based indexes including portal velocity, LSPS, portal diameter and LS, the AUCs were 0.74 (95% CI: 0.59–0.88), 0.73 (95% CI: 0.61–0.86), 0.64 (95% CI: 0.50–0.77), 0.59 (95% CI: 0.45–0.72), respectively (Fig. 3A). The AUCs of the serum-based parameters, FIB-4, Fibrosis index, Lok score, King’s score, APRI, AAR, and GPR were 0.77 (95% CI: 0.67–0.86), 0.73 (95% CI: 0.63–0.83), 0.70 (95% CI: 0.59–0.80), 0.67 (95% CI: 0.56–0.77), 0.63 (95% CI: 0.53–0.74), and 0.56 (95% CI: 0.44–0.67), respectively (Fig. 3B). Table 2 presented the AUCs, sensitivity, specificity, PPV and NPV of all non-invasive models for identifying HRV.

Ninety-nine participants’ questionnaires were collected. We classified the participants as those undergoing EGD without sedation (n = 89, 90%) or with sedation (n = 10, 10%). Calculation suggested that DS-MCCE fared significantly better than EGD without sedation for the satisfaction of the participants (average satisfaction score: 33–99 (4–23) vs. 27–84 (5–50), p < 0.0001, d = 1.15 [95%CI: 0.88–1.421]) (Fig. 4A). The respective average scores of DS-MCCE and EGD with sedation were 33–80 (3–88) and 31–80 (4–66), which showed no significant difference (p = 0.42, Fig. 4B). All par-
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Fig. 2. Examination procedure and representative examples of detachable string magnetically controlled capsule endoscopy and esophagogastroduodenoscopy.  
I. A thin and hollow string was attached on the capsule to allow the observation of esophagus by controlling the string. II. The capsule was released from the string after inspecting esophagus. III. The capsule was controlled to observe the stomach by the guidance of magnet robot. DS-MCCE, detachable string magnetically controlled capsule endoscopy; EGD, esophagogastroduodenoscopy.

Fig. 3. Diagnostic performance of detachable string magnetically controlled capsule endoscopy and other non-invasive tools.  
A. Receiver operating characteristic curves of detachable string magnetically controlled capsule endoscopy (DS-MCCE) and image-based indexes; AUC, area under receiver operating characteristic curve; LSPS, liver stiffness-spleen size-to-platelet ratio score. B. Receiver operating characteristic curves of DS-MCCE and serum-based indexes. FIB-4, fibrosis index based on four factors; APRI, aspartate aminotransferase to platelet count ratio index; AAR, aspartate aminotransferase to alanine aminotransferase ratio; GPR, gamma glutamyl trans-peptidase to platelet count ratio.

Participants confirmed excretion of the capsule and no adverse events occurred during the two-week’s follow-up.

**Discussion**

GEV comprises one of the most serious complications of portal hypertension in cACLD. Endoscopic screening of patients with cACLD to assess GEV is now recommended [1–3]. However, in view of the invasiveness and poor tolerance of EGD, non-invasive tools for HRV detection have been highlighted in recent years. This prospective blinded study showed that DS-MCCE was an accurate, comfortable, and safe, non-invasive method for inspecting HRV. It was possible to enroll a substantial population of patients with cACLD, the characteristics of whom were representative of patients with early cirrhosis needing endoscopic screening in routine practice.

In deciding whether the DS-MCCE is a valid alternative to screening EGD for patients with cACLD, three issues should be considered: diagnostic performance, patients’ satisfaction, and safety.
identify the MCCE.

Portal vein velocity, cm/s

Portal diameter, mm

AAR

LS, kpa

GPR

Table 2
Diagnostic performance of DS-MCCE and other non-invasive tools for high-risk varices.

|                | AUC (95%CI) | Sensitivity (95%CI) | Specificity (95%CI) | PPV (95%CI) | NPV (95%CI) |
|----------------|-------------|---------------------|---------------------|-------------|-------------|
| DS-MCCE        | 0.90 (0.83–0.95) | 92% (78–98%) | 88% (78–95%) | 80% (70–92%) | 95% (90–100%) |
| FIB-4          | 0.77 (0.67–0.86) | 69% (56–83%) | 75% (64–85%) | 60% (50–71%) | 82% (75–90%) |
| Portal vein velocity, cm/s | 0.74 (0.59–0.88) | 81% (62–95%) | 64% (45–79%) | 59% (48–72%) | 84% (72–96%) |
| LSPS           | 0.73 (0.61–0.86) | 77% (59–91%) | 66% (50–82%) | 57% (45–71%) | 83% (72–94%) |
| Fibrosis index | 0.73 (0.63–0.83) | 83% (69–94%) | 61% (49–73%) | 54% (46–62%) | 87% (79–95%) |
| Lok score      | 0.73 (0.63–0.83) | 92% (81–100%) | 48% (35–62%) | 52% (45–59%) | 91% (80–100%) |
| King’s score   | 0.70 (0.59–0.80) | 92% (81–100%) | 48% (35–62%) | 52% (45–59%) | 91% (80–100%) |
| APRI           | 0.67 (0.56–0.77) | 56% (39–72%) | 73% (63–84%) | 53% (41–67%) | 75% (69–83%) |
| Portal diameter, mm | 0.64 (0.52–0.77) | 79% (66–93%) | 49% (34–64%) | 49% (41–58%) | 79% (67–92%) |
| AAR            | 0.63 (0.53–0.74) | 86% (75–97%) | 48% (36–60%) | 47% (41–54%) | 86% (76–97%) |
| LS, kpa        | 0.59 (0.45–0.72) | 68% (50–86%) | 57% (45–72%) | 49% (38–60%) | 75% (64–86%) |
| GPR            | 0.56 (0.44–0.67) | 83% (72–94%) | 31% (20–42%) | 40% (35–46%) | 77% (62–92%) |

CI, confidence interval; AUC, the area under the receiver operating characteristic curve; PPV, positive predictive value; NPV, negative predictive value; DS-MCCE, detachable string magnetically controlled capsule endoscopy; FIB-4, fibrosis index based on 4 factors; LSPS, liver stiffness-spleen size-to-platelet ratio score; APRI, aspartate aminotransferase to platelet count ratio index; AAR, aspartate aminotransferase to alanine aminotransferase ratio; LS, liver stiffness; GPR, gamma-glutamyl transpeptidase to platelet count ratio.

As expected, the DS-MCCE showed a good performance for HRV detection with a C-index of 0.90, and a substantial overall agreement with EGD with a kappa score of 0.78. With EGD as the reference standard, the results showed eleven (10%) patients were misclassified, of whom only three (3%) had a missed diagnosis of HRV. A low false negative value means that suitable patients were rarely denied the opportunity of being offered primary prevention therapy for variceal hemorrhage. During EGD, air insufflation will increase the volume of the oesophageal lumen relative to the varices and perhaps make them appear smaller than is evident during DS-MCCE.

The present study evaluated several non-invasive modalities for identifying HRV in cACLD. Among them, the LSPS algorithm was the most validated non-invasive surrogate measurement of EGD to identify HRV in cirrhosis with an AUC of 0.95 (95% CI: 0.93–0.97) in hepatitis B cirrhosis [13]. However, the diagnostic accuracy of LSPS for HRV was moderate in our study with an AUC of 0.73 (95% CI: 0.61–0.86). The wider range of etiology of cACLD might explain this discrepancy. With evolved higher number of centers, we again validated that the performance of FIB-4, Fibrosis index, Lok score, King’s score were just fair and performance of AAR, APRI for assessing GEV were insufficient, as previous researches demonstrated [27]. In addition, GPR, which was originally proposed as a predictor for fibrosis stage in patients with chronic hepatitis B virus infection in west Africa, [10] did not perform well in diagnosing HRV with an AUC of 0.56 (95% CI: 0.44–0.67). This still requires further studies to confirm. Overall, direct, real-time visualization and assessment of varices by DS-MCCE outperformed reported image-based parameters and serum-based scores in terms of accuracy. Compared with previous research grading GEV by capsule endoscopy, in the largest study to date, de Franchis et al. reported a sensitivity of 78% and specificity of 96% for detecting HRV [15]. Other studies have reported a wide range of sensitivity (63–92%) and specificity (82–96%) for the detection of HRV [16–19]. However, no previous study demonstrated the diagnostic performance of capsule endoscopy for HRV in well-characterized patients with cACLD. The high sensitivity (92%) and specificity (88%) of our study for HRV suggested that DS-MCCE, in its current form, performed
well in this purpose. In addition, the patients' satisfaction for DS-MCCE was significantly better than that for EGD without sedation. DS-MCCE appeared to be a safe procedure since no adverse events occurred during follow-up. Moreover, DS-MCCE was safe for patients with dysphagia, as our previous study demonstrated the feasibility of pulling the capsule out if necessary [24]. Taken together, our results suggest that DS-MCCE used to detect varices was safe and associated with high patient satisfaction.

To the best of our knowledge, this is the first study to evaluate the DS-MCCE in assessing GEV in cACLD. The string allowed controllable movement of the capsule to observe regions of interest carefully and thoroughly. The examination scope of this technique matched that of EGD, making DS-MCCE a potential alternative to EGD especially for patients with contraindications to EGD or for those who refuse to undergo the procedure. More importantly, DS-MCCE could be performed under the direct supervision of the endoscopist from a computer workstation in an adjacent but separate room via real-time voice communication. As the COVID-19 continues to spread throughout the world, non-contact DS-MCCE can minimize risk of cross-infection and protect health-care providers better [28].

There are still some limitations. Firstly, DS-MCCE is not appropriate for patients with any contradictions to magnetic resonance imaging [22]. Secondly, the current cost of DS-MCCE is slightly higher than EGD but the cost will be decreased when it is widely used in the near future. Meanwhile, DS-MCCE takes longer than standard EGD. The median times of DS-MCCE for esophagus and stomach examinations were 6-2 and 14-3 min, respectively [29]. It should be noted that poor patient tolerance of EGD is likely to contribute to more rapid examinations, yet we know that procedures of less than seven minutes duration have half the diagnostic yield of high-risk lesions than longer procedures [30]. Therefore, a longer mean DS-MCCE examination time compared to EGD might be advantageous. Furthermore, the patient satisfaction questionnaire has not been validated. However, the questionnaire was based on one previously developed for use in assessing experience of esophageal capsule endoscopy and evaluated following the input of a panel of professionals from relevant disciplines. In addition, as DS-MCCE is a non-invasive procedure, biopsy or treatment of HRV is not possible such that a subsequent EGD is necessary for patients who need endoscopic intervention. A larger multicenter cohort (e.g. NCT03748563) is needed to further validate the accuracy of the DS-MCCE for HRV as well as portal hypertensive gastrospy.

In conclusion, DS-MCCE was an accurate, safe alternative to EGD for detecting HRV in patients with cACLD. In addition, DS-MCCE was better tolerated and thus may be indicated for those unwilling or unable to undergo EGD screening for HRV.

Declaration of Interests

The authors have declared no conflict of interests related to the study.

Author contributions

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Data sharing

The authors declare that all data supporting the findings of this study are available within the article and its supplementary. Researchers can apply for data by submitting a proposal to the corresponding author.

Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.lanwpc.2020.100072.

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