# Validation of a low cost manometer to evaluate upper airway oropharyngeal muscle strength

Submission date 04/05/2025	Recruitment status  No longer recruiting	<ul><li>Prospectively registered</li><li>Protocol</li></ul>
Registration date	Overall study status	Statistical analysis plan
06/05/2025	Ongoing	Results
Last Edited	Condition category	Individual participant data
06/05/2025	Ear. Nose and Throat	[X] Record updated in last year

## Plain English summary of protocol

Background and study aims

Obstructive Sleep Apnea-Hypopnea Syndrome (OSAHS) is a condition where people experience repeated episodes of blocked breathing during sleep. This can lead to snoring, poor sleep quality, daytime sleepiness, and increased risk of heart problems. The most effective treatment is Continuous Positive Airway Pressure (CPAP), but not everyone sticks with it. This study aims to find out if using certain tools to measure muscle strength in the mouth and throat can help improve treatment for OSAHS.

### Who can participate?

Adults aged 18 to 75 who have been diagnosed with moderate to severe OSAHS and have not received any prior treatment can participate. Healthy adults with good sleep habits and no complaints of snoring or daytime sleepiness can also participate as control subjects.

#### What does the study involve?

Participants will undergo various tests to measure the strength of their mouth and throat muscles using tools like the IOPI device, a digital manometer, and a digital spoon. They will also complete questionnaires about their sleep and muscle function. These tests will be done during a single visit to the hospital.

What are the possible benefits and risks of participating?

There are no expected side effects or complications from participating in this study. The tests are routine assessments commonly done in speech therapy and dysphagia units. Participants may benefit from gaining a better understanding of their condition and contributing to research that could improve treatment for OSAHS.

Where is the study run from?

Quirónsalud Hospital in Marbella and Hospital Campo de Gibraltar (Spain)

When is the study starting and how long is it expected to run for? January 2025 to April 2026

Who is funding the study? Investigator initiated and funded

Who is the main contact?

Dr Carlos O'Connor, carlos.oconnor@quironsalud.es

# Contact information

# Type(s)

Public, Scientific, Principal investigator

#### Contact name

Dr Carlos O'Connor-Reina

#### **ORCID ID**

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#### Contact details

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# Additional identifiers

#### Clinical Trials Information System (CTIS)

Nil known

# ClinicalTrials.gov (NCT)

Nil known

#### Protocol serial number

AWGAP-2024-2

# Study information

#### Scientific Title

Validation of the OMES questionnaire compared to objective measurement of oropharyngeal muscle tone using the IOPI manometer, Sandway manometer, and digital spoon

#### Acronym

**LCTM** 

# **Study objectives**

The use of the OMES questionnaire can be complemented by the values obtained through measurement instruments such as the IOPI device, digital manometer, and digital spoon.

#### Ethics approval required

Ethics approval required

#### Ethics approval(s)

approved 27/03/2025, CEIm Provincial de Málaga (Hospital Regional Universitario Carlos Haya. Avda Carlos Haya 84. Pabellon a.Hospital General 7 planta, Malaga, 29010, Spain; +34 951291977; eticainvestiga.hch.sspa@juntadeandalucia.es), ref: SICEIA-2024-002767

#### Study design

Prospective controlled quasi-experimental pilot study

## Primary study design

Observational

# Study type(s)

Prevention, Quality of life, Screening

## Health condition(s) or problem(s) studied

Obstructive sleep apnea

#### **Interventions**

Participants are enrolled after signing informed consent and screened to confirm eligibility (OSAHS diagnosis for cases; good sleep hygiene and absence of symptoms for controls). The study consists of a single evaluation visit taking place during the same week as the patient's sleep study. During this visit, participants complete the OMES questionnaire and undergo muscle tone assessments using the IOPI® manometer, Sandway® manometer, and a digital spoon. All tests are performed by a trained speech therapist and recorded for later blinded assessment. The total duration of observation is approximately 1 hour. There is no longitudinal follow-up; data collection is completed in a single session.

#### Intervention Type

Other

# Primary outcome(s)

Orofacial muscle function measured using the OMES questionnaire at baseline (single visit).

# Key secondary outcome(s))

- 1. Tongue and buccinator muscle strength measured using the IOPI® manometer at baseline (single visit). Measured in kilopascals (kps).
- 2. Tongue pressure measured using the Sandway® manometer at baseline (single visit). Measured in kilopascals (kps).
- 3. Tongue strength measured using a digital spoon at baseline (single visit). Measured in gr/cm<sup>2</sup>.
- 4. Correlation between OMES score and values obtained by IOPI®, Sandway®, and digital spoon, analyzed at baseline (single visit)

# Completion date

30/04/2026

# Eligibility

#### Key inclusion criteria

#### Cases:

- 1. Age between 18 and 75 years
- 2. Diagnosis of moderate to severe OSAHS (AHI >15) with no previous treatment experience
- 3. No prior treatment for OSAHS
- 4. Signed informed consent (IC)

#### Controls:

- 1. Adequate sleep hygiene
- 2. No complaints of snoring
- 3. No complaints of daytime sleepiness
- 4. Epworth Sleepiness Scale < 7 points

#### Participant type(s)

Healthy volunteer, Patient

#### Healthy volunteers allowed

No

#### Age group

Adult

#### Lower age limit

18 years

#### Upper age limit

75 years

#### Sex

All

#### Key exclusion criteria

- 1. Cognitive or neurological deficits
- 2. Inability to complete questionnaires
- 3. Severe alcoholism
- 4. Presence of craniofacial malformations
- 5. Active neoplastic disease
- 6. History of orofacial muscle rehabilitation or prior OSA treatment (surgery, MAD, CPAP)

### Date of first enrolment

05/05/2025

#### Date of final enrolment

05/12/2025

# Locations

#### Countries of recruitment

Spain

# Study participating centre Hospital Quironsalud Marbella

Avda Severo Ochoa 22 Marbella Spain 29603

# Study participating centre Hospital Quironsalud Palmones

Edificio Arttysur. Avda de los Empresarions s/n Parque Empresarial las Marisma de Palmones Palmones Spain 11379

# Sponsor information

# Organisation

Hospital Quirosalud Marbella

# Funder(s)

# Funder type

Other

#### **Funder Name**

Investigator initiated and funded

# **Results and Publications**

# Individual participant data (IPD) sharing plan

The dataset generated during and/or analysed during the current study will be available upon request from Dr Eduardo Correa educorreaorl@gmail.com

# IPD sharing plan summary

Available on request

# Study outputs

Output type Details
Participant information sheet

Participant information sheet Participant information sheet 11/11/2025 No Yes

Protocol file 06/05/2025 No No