Vitalograph® copd-6™

User Manual

Model No. 4000

Medical Devices Directive
EN ISO 13485
FDA QSR 21 CFR 820/803

Manufacturer: Vitalograph (Ireland) Ltd, Ennis, Ireland
Table Of Contents

Warnings and Advisory Notices: ..................................................................................................3
Main Components of the Vitalograph copd-6™ ........................................................................4
What Is The Vitalograph copd-6™ Used For? ........................................................................4
What Is The Vitalograph copd-6™ Used For? ........................................................................5
How To Use The Vitalograph copd-6™ .................................................................................5
  Entering Subject Data ...........................................................................................................5
  Performing the Test ...............................................................................................................6
  Setting the Obstructive Index and COPD Classification Zones ...........................................7
  Reviewing the Last Session Test Results ............................................................................8
Care and Cleaning Of The Vitalograph copd-6™ ..................................................................9
  Daily Procedures ................................................................................................................9
  Monthly Procedures ...........................................................................................................9
  Annual Procedures .......................................................................................................... 9
  Table of Materials Used & Cleaning/Disinfection Methods ................................................9
  Removing the Flowhead for Cleaning and Disinfecting ......................................................10
Consumables, Accessories and Spare Parts ...........................................................................10
Explanation of Symbols .........................................................................................................11
Technical Specifications .........................................................................................................12
Warranty ..................................................................................................................................12
  Quick Start Guide ...............................................................................................................13
  Regulatory Notices for the Vitalograph copd-6 .................................................................14
    CE Notice ........................................................................................................................14
    FDA Notice ......................................................................................................................14
    Declaration of Conformity ...............................................................................................16

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Warnings and Advisory Notices:

Note: Please read all the information in this manual before using the Vitalograph copd-6. A full set of instructions, including cleaning instructions, is available at www.vitalograph.co.uk.

• This Vitalograph device is intended to measure lung function for use in clinic with disposable mouthpieces or filters or for patient use at home under medical supervision.
• Take care not to block the mouthpiece with the tongue or teeth. A ‘spitting’ action or coughing will give false readings.
• If used at home symptoms must take precedence over device measurements*.
• If the device is used for longer than its specified life, the accuracy of the device may deteriorate.
• Before use, ensure that the batteries do not exceed their shelf life, as indicated on the batteries.
• Store in a clean dry place.
* If you suspect that a defect has occurred you should check the Vitalograph copd-6 for operation and accuracy.
Main Components of the Vitalograph copd-6™

A  Flowhead
B  On / off button
C  Display
D  User buttons
What Is The Vitalograph copd-6™ Used For?

The Vitalograph copd-6 is a device intended for measuring lung function. It can be used by a healthcare professional or assistant as a rapid pre-spirometry test to screen-out at-risk individuals that do not have COPD and indicate those that may be at risk of COPD at the pre-symptomatic stage of the disease. This “screening” or “case selection” for full spirometry examinations may result in earlier medical intervention and facilitate better clinical outcomes.

The Vitalograph copd-6 displays some key parameters for clinical interpretation:

- FEV1 and FEV1 % Predicted
- FEV6 and FEV6 % Predicted
- FEV1/FEV6 and FEV1/FEV6 % Predicted
- FEV1/FVC ratio

If an abnormal result is indicated, arrows on the display show:

- Obstructive Index
- COPD Classification (stages I - IV)

How To Use The Vitalograph copd-6™

Entering Subject Data

The subject’s physical data can be entered into the Vitalograph copd-6 in order to calculate predicted data.

To enter the subject data, follow these steps:

1. Turn the device on, 

2. The age can now be set ( ). This is done by pressing the ▲ or ▼ button and releasing when the age is reached.
   The values will increase/decrease in values of 1. If the button is kept depressed, the values will scroll faster.
   Press the ← button to set the age.

3. The height can now be set ( ). This is done by pressing the ▲ or ▼ button and releasing when the height is reached.
   The values will increase/decrease in values of 1. If the button is kept depressed, the values will scroll faster.
   Press the ← button to set the height.
   Note: if height values are set below 100, the device assumes height is in inches and that Weight will be in Ibs rather than Kg.

4. The gender can now be set ( ). This is done by pressing the ▲ or ▼ button and releasing when the correct gender is showing. (Male (♂); Female (♀))
   Press the ← button to set the gender.
   After gender is selected, the Vitalograph copd-6 will go to Test Mode, where the Blow Icon is showing. On some variants, an additional data field comes first (see below)
5. (This step may be omitted, depending on the variant) The Population Group (On some variants this is Weight instead) can now be set (Ø). This is done by pressing the ▲ or ▼ button and releasing when the desired setting is reached.

Press the ← button to set.

<table>
<thead>
<tr>
<th>Set No.</th>
<th>Example Population Group</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NHANES III Caucasian</td>
</tr>
<tr>
<td>2</td>
<td>NHANES III African-American</td>
</tr>
<tr>
<td>3</td>
<td>NHANES III Hispanic-American</td>
</tr>
</tbody>
</table>

Performing the Test

1) The subject should sit down when blowing into the device (unless advised otherwise). Insert a new disposable SafeTway mouthpiece into the device.

2) Turn the device on, ⊙ (if not already ON). When the device is ready for a test the blow icon shows (⊙).

3) Instruct the subject and demonstrate (using a mouthpiece) as follows:
   a) “Hold your head up, breathe in as deeply as possible, hold the device in front of your mouth”.
   b) “Hold your breath, place the mouthpiece into your mouth, like this, bite the mouthpiece lightly, and seal your lips firmly around it, like this”. I will demonstrate.
   c) “Blow out as HARD, and FAST as you can, like this, until I tell you to stop” (the unit will beep at end of test - after 6 seconds).
   d) “Be careful not to block the mouthpiece with your tongue or teeth. A ‘spitting’ action will give false readings”.
   e) “Now you do it – deep breath in – bite and seal – blast the air out…keep going – keep blowing”
   f) “Well done!” “Now, we need to do that three times. Rest for a while until you are ready to blow again”.

4) To view the result (the best values in the session), press the ← button.
   a) The COPD Classification will show on the right hand arrow.
      i) Green is NORMAL, negative for COPD. No need to refer this subject for spirometry.
      ii) Any one of the blue zones, I, II, II or IV are not normal. This subject should be referred for spirometry.
   b) The Obstructive Index (OI) shows on the left hand arrow.
      i) 0 - Green is normal.
      ii) 1, 2 or 3 – Yellow, amber or red are not normal. Refer for spirometry, but this is unlikely to be COPD.
iii) Note: If the right hand (COPD) arrow is not green, nor will the left hand (OI) arrow be green.

5) This is the end of the test, but if desired some test parameters may be viewed.

6) Following each blow and at the end of the test session, the FEV1 value will be displayed and below that, FEV1 % Predicted results for that blow, or for the best in session if the E button has been pressed. Pressing E again will toggle between best and last blow.

7) Pressing the ▲ button will show the FEV6 and FEV6 % Predicted results.

8) Pressing the ▲ button again will show the FEV1/FEV6 and FEV1/FEV6 % Predicted results.

9) Press the ▲ button for a final time will show the estimated Lung Age (📅)

Notes on testing:

1. During testing, if an exclamation mark (!) appears, this means that the last blow was not a good quality blow and the user should blow again. Reasons for poor quality are:
   a. Slow Start of test: Vext (extrapolated volume) is > 5% or 150mL of FEV6
   b. Cough detected.

2. In the event that a test is < 3 seconds duration and has an abrupt end (change in volume is > 25ml in the last second of the test), the value FEV will be displayed instead of FEV6.

3. If the test subject experiences side effects such as dizziness or fatigue during the test procedure, stop testing until recovered.

### Setting the Obstructive Index and COPD Classification Zones

The Vitalograph copd-6 Obstructive Index and COPD Classification zones are set by the manufacturer to the GOLD (Global Initiative for Chronic Obstructive Lung Disease) standard and it is unlikely that a user will wish to change this – however if this is required please follow the following instructions;

The colour systems for each zone type are pre-set as follows;

<table>
<thead>
<tr>
<th>FEV1%Pred</th>
<th>Obstructive Index</th>
<th>COPD Classification</th>
<th>FEV1/FEV6 Ratio and FEV1%Pred.</th>
</tr>
</thead>
<tbody>
<tr>
<td>≥ 80%</td>
<td>0</td>
<td>Not COPD</td>
<td>FEV1/FEV6 &gt; 0.7</td>
</tr>
<tr>
<td>&lt; 80%</td>
<td>1</td>
<td>Stage I</td>
<td>FEV1/FEV6 &lt; 0.70 and FEV1 ≥ 80% Pred.</td>
</tr>
<tr>
<td>&lt; 50%</td>
<td>2</td>
<td>Stage II</td>
<td>FEV1/FEV6 &lt; 0.70 and FEV1 &lt; 80% Pred.</td>
</tr>
<tr>
<td>&lt; 30%</td>
<td>3</td>
<td>Stage III</td>
<td>FEV1/FEV6 &lt; 0.70 and FEV1 &lt; 50% Pred.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Stage IV</td>
<td>FEV1/FEV6 &lt; 0.70 and FEV1 &lt; 30% Pred.</td>
</tr>
</tbody>
</table>

To set the boundary percentage values for both the Obstructive Index and COPD Classification zones together, follow these steps;

1. Turn the device on, ⊙.
2. When the ![icon] icon appears, press and hold the ← and ▲ buttons for approximately 3 seconds.

3. The top boundary can now be set. This is done by pressing the ▲ or ▼ button and releasing when the value is reached.
   The values will increase/decrease in values of 1%. If the button is kept depressed, the values will scroll faster.

4. Press ← to set the top boundary value.

5. The middle boundary can now be set. This is done by pressing the ▲ or ▼ button and releasing when the value is reached.
   The values will increase/decrease in values of 1%. If the button is kept depressed, the values will scroll faster.

6. Press ← to set the middle boundary value.

7. The bottom boundary can now be set. This is done by pressing the ▲ or ▼ button and releasing when the value is reached.
   The values will increase/decrease in values of 1%. If the button is kept depressed, the values will scroll faster.

8. Press ← to set the bottom boundary value.

9. Press ←. The device will return to the age entry screen.

**Reviewing the Last Session Test Results**

The Vitalograph copd-6 will always store the last test session, even after the device has powered itself down or has been switched OFF. In order to view the last test session, follow these steps;

1. Turn the device on,  minX

2. When the device is ready for age entry (↑↓), press the ← button for approximately 3 seconds. The last test session (best results) data will now show again.

3. When you have finished reviewing the data, press the OFF button for 3s. OR

4. Press ←. The device will return to the age entry screen ready for entering the next test subject's data.
Care and Cleaning Of The Vitalograph copd-6™

Daily Procedures

A new mouthpiece (either SafeTway or BVF) should be used for each subject. A delay of at least 5 minutes should be allowed between subjects to allow settling of previously aerosolized particles in the measuring device.

Monthly Procedures

It is recommended that the device be regularly cleaned according to the guidelines of the user’s facility. The disinfection materials and procedures applied in the users’ facility may be more appropriate than the methods outlined below. In the event of visible contamination of the flowhead element, it should be cleaned or disinfected as described in the accompanying table. The device should be replaced in the event of damage, or if visibly contaminated.

The frequency of cleaning and disinfecting is dependent on the facility’s risk assessment, usage, and test environment, but it should be at least monthly or every 100 subjects (300 blows).

Annual Procedures

It is recommended that the device be replaced annually or test and calibration serviced at least annually. There is no planned preventive maintenance for this medical device.

Table of Materials Used & Cleaning/Disinfection Methods

This listing of materials used is given to provide clinical users with information to allow the assessment of other cleaning and disinfecting procedures available in the facility on this device.

<table>
<thead>
<tr>
<th>Part</th>
<th>Material</th>
<th>Clean/ Disinfect</th>
<th>Autoclave Possible?</th>
<th>Recommended Disinfectants</th>
</tr>
</thead>
<tbody>
<tr>
<td>SafeTway mouthpiece or BVF</td>
<td>Cardboard / ABS</td>
<td>Dispose – single use</td>
<td>No</td>
<td>Dispose – single use</td>
</tr>
<tr>
<td>Case Exterior</td>
<td>PC/ABS</td>
<td>Clean</td>
<td>No</td>
<td>Wiping with a 70% isopropyl alcohol impregnated cloth provides a suitable form of cleaning and low-level disinfection. This may be preceded by cleaning with an anti-static foam cleaner if necessary.</td>
</tr>
<tr>
<td>Fascia</td>
<td>PMMA/PET</td>
<td>Clean</td>
<td>No</td>
<td></td>
</tr>
<tr>
<td>Removable flowhead</td>
<td>PC/ABS</td>
<td>Clean</td>
<td>No</td>
<td>Disinfect by immersion in sodium dichloroisocyanurate solution at 1000 ppm concentration of free chlorine for 15 minutes (see following section for recommended cleaning/disinfection method for the flowhead)</td>
</tr>
</tbody>
</table>
All external parts of the Vitalograph asma-1 require **cleaning**, i.e. the removal of visible particulate contamination. The parts of the device that make up the flowhead, which comes into contact with the breath of the subjects being tested, also require **disinfecting**. This device is not designated as a ‘sterile’ device.

**Removing the Flowhead for Cleaning and Disinfecting**

1. Remove the flowhead from the body with a sharp pulling motion.
2. Clean the flowhead by washing in a mild detergent to remove particulate contamination, taking care not to touch the moving vanes. Swill vigorously in water with mild detergent. Do not attempt to “rub” or “scrub” in the area of the vanes. Rinse with clean water.
3. Disinfect by immersion in sodium dichloroisocyanurate solution at 1000 ppm concentration of free chlorine for 15 minutes. Prepare disinfectant solution as directed in the manufacturer’s guidelines. Rinse with warm water for faster drying.
4. Leave it to dry completely before reassembling. Drying the flowhead may require placing it in a warm place overnight. A drying cabinet is ideal.

Wiping with a 70% Isopropyl Alcohol impregnated cloth provides a suitable form of cleaning and low-level disinfection for the case exterior, display, screen surround and keys. Repeat this at least weekly to prevent a build-up of grime from normal handling and use.

Always follow the safety guidelines given by the manufacturer of cleaning and disinfectant chemicals or equipment.

Reassemble the flowhead by pushing back on until it ‘clicks’ into position. Ensure that the flowhead is pushed fully home.

When the flowhead is reassembled, it is good practice with any respiratory measuring device for an accuracy check be performed using a Precision Syringe, with the volume delivered in less than one second. An accuracy of +/- 3% should be achieved.

**Definitions of cleaning and disinfection**

* Definitions of cleaning and disinfection are as defined in “Sterilization, Disinfection and Cleaning of Medical Equipment: Guidance on Decontamination from the Microbiology Committee to Department of Health Medical Devices Directorate, 1996”
* Recommendations for chemical disinfectants are derived from the PHLS publication “Chemical Disinfection in Hospitals” 1993

**Consumables, Accessories and Spare Parts**

<table>
<thead>
<tr>
<th>Cat. No</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>20242</td>
<td>SafeTway Mouthpieces (200)</td>
</tr>
<tr>
<td>20303</td>
<td>Disposable Noseclips (200)</td>
</tr>
<tr>
<td>20980</td>
<td>Mini SafeTway® mouthpieces (50)</td>
</tr>
</tbody>
</table>
Explanation of Symbols

Device symbols;

- ▶ Type BF equipment
- □ Class II

- The device must be taken to separate collection at the product end-of-life. Do not dispose of these products as unsorted municipal waste.
- △ Attention (reference relevant section in manual)

User Interface Symbols;

- ▢ Battery status
  - Battery status Full
  - Battery status Half
  - Battery status Quarter
  - Battery status Empty (flashing)
- □ Blow Now Symbol
- ! Bad Test Symbol (Slow start or Cough)

- kg/lb Unit of weight measurement
- ▲ Lung Age Symbol
- △ Age Symbol
- ▶ Height Symbol
- ♂ Gender Symbol
- ○ Population Group Symbol
Technical Specifications

Material: PC/ABS
Accuracy: Better than ± 3%
Flow Impedance: Better than 0.15kPa/L/s at 14L/s
Measurement Range: 0 – 9.99 L BTPS
Performance And Safety Standards: ATS/ERS Guidelines 2005
Electromagnetic emissions: CISPR 11 Group 1 (battery operated)
Electromagnetic immunity: IEC 61000-4-2, IEC 61000-4-3 (battery operated)
Sensor: Stator/rotor
Power Supply: 2 x AAA batteries
Operating Temperature: 17 – 37°C

Bad Test Criteria
Slow start of test (Vext>5%) or a cough detected in the first second
FEV substituted for FEV6
When FET < 3s and abrupt end of test
Auto power down time
Set to 2 minutes as standard

Warranty

Your Vitalograph copd-6 is guaranteed for one year*. Replace if it is faulty, otherwise replace the unit every three years.

* Excepting accidental / transit damage or inappropriate use of the device.
## Appendices

### Quick Start Guide

#### Prepare Device
1. **Switch On.**

#### Set test subject's gender.

- **Press** or **to change gender.**
- **Press** to set.

#### Set test subject's age.

- **Press** or **to change age.**
- **Press** to set.

#### Set test subject's height.

- **Press** or **to change height.**
- **Press** to set.

#### Perform Test
1. **Insert disposable SafeTway mouthpiece.**

2. **Demonstrate to test subject.**
   - Fully breathe in, hold...
   - Insert and tightly bite the mouthpiece.
   - Press blow icon and blow for 5 seconds for repeat.
   - WELL done! Test subject for a short while and wait for blow icon to reappear.

3. **Repeat blow 2 more times.**

#### Test Result
1. **Press** to show result.

- **Negative**
  - Spirometry not required
  - COPD Classification given

- **Positive**
  - Refer for spirometry
  - COPD Classification I to IV

#### View Parameters
1. **Display shows FEV1 and % of predicted.**

2. **Press** to show FEV1 and % of predicted.

3. **Press** to show FEV1 Ratio and % of predicted.

4. **Press** to show Lung Age.

- [www.vitalograph.co.uk](http://www.vitalograph.co.uk)
Regulatory Notices for the Vitalograph copd-6

CE Notice

Marking by the symbol \( \bigcirc \) indicates compliance of the Vitalograph copd-6 to the Medical Devices Directive of the European Community. Such marking is indicative that the Vitalograph copd-6 meets or exceeds the referenced technical standards.

The Vitalograph copd-6 uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.

The Vitalograph copd-6 is battery operated and is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.

The Vitalograph copd-6 is intended for use in the electromagnetic environment specified below. The customer or the user of the system should assure that it is used in such an environment.

- Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.
- Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment
- Interference may occur in the vicinity of equipment marked with the following symbol - \( \bigcirc \).

FDA Notice

Caution: Federal Law restricts this device to sale by, or on the order of a physician.
The Vitalograph copd-6 is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the system can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the system as recommended below, according to the maximum output power of the communications equipment.

<table>
<thead>
<tr>
<th>Rated maximum output power of transmitter (W)</th>
<th>Separation distance according to frequency of transmitter (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>150 kHz to 80 MHz</td>
<td>80 MHz to 800 MHz</td>
</tr>
<tr>
<td>d = 1.2 \sqrt{P}</td>
<td>d = 1.2 \sqrt{P}</td>
</tr>
<tr>
<td>0.01</td>
<td>0.1m</td>
</tr>
<tr>
<td>0.1</td>
<td>0.4m</td>
</tr>
<tr>
<td>1</td>
<td>1.2m</td>
</tr>
<tr>
<td>10</td>
<td>3.7m</td>
</tr>
<tr>
<td>100</td>
<td>11.7m</td>
</tr>
</tbody>
</table>

For transmitters rated at a maximum output power not listed above, the recommended separation distance \( d \) in metres (m) can be estimated using the equation applicable to the frequency of the transmitter, where \( P \) is the maximum output power rating of the transmitter in watts (w) according to the transmitter manufacturer.

Note 1: at 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

Note 2: these guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.
Declaration of Conformity

Product: Vitalograph copd-6™

Vitalograph hereby ensures and declares that the above product associated with this user manual, is designed and manufactured in accordance with the following QMS regulations and standards:

European Medical Devices Directive {MDD} 93/42/EEC. This device, classified as 2a as per Annex IX of MDD 93/42/EEC, meets the following provisions of Annex II of the Medical Devices Directive as per Article 11, section 3a, excluding point 4 of Annex II. This device complies with the EMC Directive 89/336/EC, conformance demonstrated by following standard EN60601-1-2:2001. Equipment classification: Residential.

Canadian Medical Device Regulation {CMRD}

FDA Quality System Regulation {QSR} 21 CFR 820.


Certifying Body {for 93/42/EEC and CMDR}: British Standards Institute {BSI}

Certificate Nos. CE 00772, MD 82182, FM 83550