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1. Main Components

Figure 1 The main components of the Vitalograph micro.

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>PC Software</td>
</tr>
<tr>
<td>2</td>
<td>Flowhead</td>
</tr>
<tr>
<td>3</td>
<td>Flowhead Release Button</td>
</tr>
<tr>
<td>4</td>
<td>LCD/Touch Panel Display</td>
</tr>
<tr>
<td>5</td>
<td>Mini USB Port</td>
</tr>
<tr>
<td>6</td>
<td>On/Off Button</td>
</tr>
<tr>
<td>7</td>
<td>LED</td>
</tr>
<tr>
<td>8</td>
<td>Battery compartment (4 x 1.5V AAA Batteries)</td>
</tr>
</tbody>
</table>

**Note:** Computer not supplied.
1.1. Features

- Fleisch Pneumotachograph
- Removable flowhead
- Touch screen color display
- Choice of predicted values
- Report generation through Vitalograph Reports software
- Storage of tests and demographic information

2. Set Up

1. Insert 4 x 1.5V AAA batteries into the battery compartment. Alternatively the micro may be powered from the user’s computer via the mini USB port in the base using the USB cable supplied. Press the On/Off button on the front face of the instrument.

2. When first turned on, the date and time setup screen is presented. Use the up/down buttons to enter the current date and time, and press enter to save these settings.

3. The Vitalograph® micro is now ready for use.

4. To use with the Vitalograph Reports Application, install the software provided with the micro following the instructions supplied with the software.

5. Connect the micro to the computer using the USB cable (via ports marked with the symbol).

If the device has just been unpacked or transported, ensure that it is left sitting, fully powered so that it is at room temperature prior to testing.
3. Operating the Vitalograph micro

The Main Menu screen includes the following options - New Subject, VC Test, FVC Test and Post Test (for bronchodilator responsiveness testing).

When turning the device on for the first time the test screen icons will appear greyed out and can’t be selected until a subject is created. The Post Test icon will stay greyed out until an FVC pre-test is performed.

In addition to displaying the time and the battery icon the status bar at the top on the screen will show various icons to indicate the following:

1. V - indicates a VC test has been performed
2. F - indicates an FVC has been performed
3. P - indicates post mode (for bronchodilator responsiveness testing)

The icons will only appear after the test has been completed.

3.1. Entering Subject Data

1. Select the New Subject button on the Main Menu, to open the New Subject screen.
2. The Subject information fields available are as follows:

   - Age
   - Height
   - Gender
   - Weight
   - Ethnicity
3. Age, Height and Gender are on the first screen and are enabled by default. Weight and Ethnicity are on the second screen and are not enabled by default. Enable options by selecting:

- Configuration menu
- Subject Options

4. To enter information for Age, Height and Weight select the appropriate icon and type in the information using the touch panel keypad.
   - Units automatically switch between cm/in and kg/lbs.
   - Gender is selected by pressing either Female or Male

   - Ethnicity is selected from a list by pressing the appropriate option on screen. To access additional Ethnicities select the arrow on the right of the screen.

5. To save the subject details and return to the Main menu press the Enter button

6. If a value is not entered for Age, Height or Gender, an Error Icon will appear next to the empty field when the enter button is pressed, this is to indicate that the predicted values will not appear in the results of any testing done.

7. To exit the new subject screen press the enter button again.

**Note:** The micro retains data for the last subject until a new subject is created. Therefore, with the exception of the first use of the device or after clearing the memory, when the user enters the New Subject Screen the fields appear greyed out to indicate that data is retained. Selecting age, height or weight boxes will move the last subject data to the device memory to allow the user to enter new subject information. All subject details need to be completed in order to continue.
3.2. Conducting a Test

Before starting a test session:

1. Ensure that the accuracy of the device was checked recently. (Refer to the section on Calibration Verification).
2. Select a subject and ensure the required demographic information is entered.
3. Wash hands (operator and subject).
4. Fit a new Bacterial Viral Filter (BVF™) to the flowhead for each test subject. The use of a disposable noseclip is recommended.
5. Instruct and demonstrate the test.

3.2.1. Testing

1. From the Main Menu select either

- VC Test
- FVC Test

2. Wait for the ‘Blow Now’ icon to appear
   The device is now ready to accept a blow.

Example script¹:

- Sit upright with back straight and feet flat on floor. Fit the nose clip and relax.
- Place BVF in mouth and close lips around the mouthpiece.
- Seal your lips around the mouthpiece and keep your tongue down.
- Breathe normally

VC test session (with V/T graph display selected):

1. Inhale completely with a brief pause when your lungs are completely full (≤ 2 secs).
2. Exhale with no hesitation until no more air can be expelled while maintaining an upright posture.

¹ Derived from terminology and guidance taken from ATS/ERS Standardisation of Spirometry 2019 Update Am J Respir Crit Care Med 2019 Vol 200, Iss 8 pp e70-e88
It is vital that the operator encourages the subject to keep exhaling to ensure all air is expelled (when a plateau has been reached or forced expiratory time (FET) reaches 15 seconds). The operator should repeat instructions as necessary, coaching vigorously.

3. Listen for two beeps. This indicates that device is ready for the next blow.
4. Repeat for a minimum of three manoeuvres, usually no more than eight for adults.
5. Check VC repeatability and perform more manoeuvres as necessary.

Note: A single-breath VC technique may also be performed on the device.

FVC test session:

1. Inhale completely and rapidly with a brief pause when your lungs are completely full (≤ 2 secs).
2. Exhale with maximal effort until no more air can be expelled while maintaining an upright posture.

It is vital that the operator encourages the subject to keep exhaling to ensure all air is expelled (when a plateau has been reached or forced expiratory time (FET) reaches 15 seconds). The operator should repeat instructions as necessary, coaching vigorously.

3. Breathe in with maximal effort until completely full. The manoeuvre is now complete and the BVF is removed from the mouth.
4. Listen for two beeps. This indicates that device is ready for the next blow.
5. Repeat for a minimum of three manoeuvres, usually no more than eight for adults.
6. Check FEV1 and FVC repeatability and perform more manoeuvres as necessary.

Note: A single-breath FVC technique may also be performed on the device.

When testing is complete, press the enter button to exit the test screen and return to the Main Menu.
3.2.2. Saving the Test Session

The Vitalograph micro has capacity to store 750 subject entries with corresponding session data. Only the best 3 blows will be stored with each session. Stored session information includes subject details and best pre-test if it is a bronchodilator responsiveness testing session.

The Vitalograph micro is intended to be used to store test data temporarily. When the device is connected to Vitalograph Reports to produce pdf reports of the session data, all subject/sessions are moved to Vitalograph Reports and cleared from the device with the exception of the last FVC Pre-test done.

If more than 750 subject/session entries are stored on the device the existing subject/sessions entries will be deleted on a First In First Out (FIFO) basis i.e. the first session entered will be the first to be deleted.

3.2.3. Bronchodilator Responsiveness Testing

Bronchodilator responsiveness testing can be performed on the most recent FVC pre-test session performed. The device will retain the last pre-test even when it is turned off and on again and/or the data has been transmitted to Vitalograph Reports.

To perform a bronchodilator responsiveness test:

1. From the Main Menu select ‘Post Mode’

2. Perform the Post FVC Test Session following the example script for ‘FVC test session’ in section 3.2.1.

Note: Post Mode may only be selected if a FVC Pre-test has been completed. When returning to the main menu from the Post FVC test screen it is not possible to select either the VC or FVC test as the device is still in Post mode. These options will be greyed out.

3.2.4. Viewing VC Test Results

Results may be viewed as either a Volume/time (V/t) or Volume Bar graph by pressing the graph button on the side of the test screen. It is not possible to change the view of results during testing.
Volume/time (V/t) Volume bar graph

1. The graph may be changed to a full screen graph by using the zoom button on the side of the test screen. To return to normal mode select the zoom in button.

2. The results summary on the bottom of the screen shows the VC of the last blow. The number of blows is shown in a separate box next to the last test VC.

3.2.5. Viewing FVC Test Results

The results may be viewed as either a Volume/time (V/t) or Flow/Volume (F/V) graph by pressing the graph button on the side of the test screen. It is not possible to change the view of results during testing.

Volume/time (V/t) Flow/Volume (F/V)

1. The graph may be changed to a full screen graph by using the zoom button on the side of the test screen. To return to normal mode select the zoom in button.

2. The results summary on the bottom of the screen shows the FVC and FEV1 of the last blow.

3. The number of usable blows and bad blow indicator (!) are shown in a separate box next to the last test FVC and FEV1.

4. The best three tests are shown on the graph in order of rank (best 1, 2, 3...). A key is shown at the top of the graph to help identify the tests.
5. To view results as a table, select the results button on the side of the test screen.
   a. Select the test results for viewing by using the left/right arrows.
   b. Scroll through the results for each test by using the up/down arrows. The number of parameters displayed will depend on the configured parameters.
   c. Tests are shown in order of rank (best is ranked number 1 then 2, 3,...).

6. The results screen has several different columns:
   - Parameter name
   - Units
   - Test value (Pre and Post for bronchodilator responsiveness testing)
   - %Pred or Z-value (depending on the configuration)

3.2.6. Deleting Test Results

To delete the current blow:

1. From the menu on the side of the test screen, select the delete button.

2. To confirm the deletion of the blow press the Delete icon with a green tick.

3. To cancel the deletion select the Delete icon with a red cross X.

3.2.7. Ending a Test Session

A session ends and is saved when one of the following occur:
   - The device is turned off
   - A new subject is created
   - The device is connected to Vitalograph Reports

3.3. Reporting

Generating PDF reports from the micro requires a computer running Vitalograph Reports. Different tests conducted during the same session ie VC, FVC,
Post are treated as a single session and are printed as one report. If more than one test report is required for the same subject, the device should be switched off and on again between tests so that they are registered as separate sessions and separate reports can be generated.

1. To produce PDF reports from the micro, connect it to a computer using the USB cable supplied with the device.
2. Run Vitalograph Reports on the computer.
3. Ensure the micro is switched on and in the home screen.
4. Guidance on using Vitalograph Reports can be found in the Vitalograph Reports instruction for Use and in the software help menu.
5. Connect to Vitalograph Reports and select to move all data to clear/delete all the sessions from the device.

**Note:** When the micro is connected to Vitalograph Reports it will move, not copy, the stored sessions with the exception of the latest FVC Pre-session.

### 3.4. Calibration Verification

1. Select Configuration button at the top right corner of the main menu.

![Configuration button](image)

2. Select Calibration Verification

![Calibration Verification](image)

3. Enter the Syringe volume and reference using the touch panel keypad.
4. Enter the ambient temperature using the touch panel keypad.
5. Attach the flowhead to the syringe with a BVF fitted as shown in figure 2.
6. Pump air through the flowhead to bring it to ambient temperature. If the flowhead has very recently been used for testing or has come from a cold environment, its temperature should be equilibrated by pumping air through it from the syringe several times.

7. Press the Enter key to open the Calibration Verification screen and follow the on-screen instructions.

8. The calibration verification result is shown in % in the top right corner of the screen. If it is reproducible and within 3% a green tick will show. Press Enter to return to the main Configuration menu. The verification pass is recorded.

9. If the Calibration verification result is outside 3% the error icon will show. Consult the micro fault finding guide at section 7. Press the Enter key to proceed to the calibration update routine.

10. The Calibration Update screen shows the volume (L) at the top left corner of the screen, next to the number of strokes.

11. The procedure is the same as for the Accuracy Check. If two strokes are within 3% of the reference volume, press Enter to return to the Configuration menu. The Calibration factor is not updated and a pass is recorded. If outside 3%, the error
The icon is shown. Press Enter to return to the Configuration menu. The Calibration factor is updated and the Calibration update is recorded.

**Note:** To exit the Calibration verification screen without performing a check, press the Enter key to return to the Configuration Menu screen. The calibration verification will not be logged where the calibration verification routine has not been completed.

When to check accuracy:
- In accordance with establishment procedures
- After service checks
- After cleaning or disassembling spirometer for any reason
- After adjusting calibration
- If the flowhead or device has been dropped
- If a new flowhead has been fitted

### 3.5. Setting up a New Flowhead

After fitting a New Flowhead to the Vitalograph micro it is necessary to set up the flowhead with the device.

Follow these steps when setting up a new flowhead.

1. Perform a full calibration verification as per section 3.4. above.
2. Select the Accuracy/Calibration icon again and repeat steps from section 3.4. up to step 7.
3. Instead of continuing to follow on-screen instructions, withdraw the syringe fully and follow instructions below:
4. Select the New Flowhead icon.

5. Slowly push in the syringe fully and then withdraw fully, keeping the flow rate below the 0.75 L/sec limit lines on the graph. If this is performed correctly new limit lines of 2.50 L/sec will appear.
6. Using a medium speed push in the syringe fully and then withdraw fully, keeping the flow rate between the 0.75 L/sec and 2.50 L/sec limit lines on the graph. If this is performed correctly new limit lines of 10.00 L/sec will appear.
7. Using a fast stroke push in the syringe fully and then withdraw fully, keeping the flow rate between the 2.50 L/sec and 10.00 L/sec limit lines on the graph.

8. The result is shown in % on the bottom of the screen. If it is within 3% a green tick icon will be shown to indicate successful setup of the new flowhead. Press the Enter key to return to the main Configuration menu.

9. If the result is outside 3% the error icon will be shown. Press Enter to proceed to the new flowhead set-up screen. 

Only complete the next step where result is outside 3%

10. Repeat steps 5-8. At the end of the procedure a green tick icon will appear. Press the enter key to return to the Configuration menu.

Note: To exit the New Flowhead screen, press the Enter key again to return to the Configuration Menu screen. The result will not be logged to the Vitalograph micro memory in this case.

3.6. Configuration Options

To access the Configuration menu select the configuration icon on the top right corner of the main screen.

In the Configuration menu there are four options:

• Subject options (top left icon)
• Device settings (top right icon)
• Accuracy and Calibration (bottom left icon)
• About box (bottom right icon)
1. **Subject Options**

- **Posture** - sets Posture to be recorded as sitting or standing.
- **Weight** - turn on to enable the option to enter Weight in the subject screen.
- **Population Group** - turn on to enable the option to enter Ethnicity in the subject screen.

*Note: Default Posture is sitting. Default for Weight and Ethnicity is Off.*

2. **Device Settings**

**Results Options**

- Display % Predicted or Z-score in the results screen.
- Turn Sounds on or off.
- Device lock on or off. Turning on requires the user to enter a passcode.
- Set Temperature (up to 2 decimal places). The default setting is 23°C.
Parameters (See Parameters section)

- Select the parameters to be displayed in results.
- Use the left/right arrows to navigate between the screens.
- A maximum of 8 parameters can be selected.

Date/Time

- Select this option to set or change the Date and/or Time.
- Use the up/down arrows to edit these fields.
- Date format is YYYY/MM/DD, Time is in 24 hour format.

Service/Technician Mode

- An 8 digit passcode is required to enter this mode.

3. Accuracy/Calibration

- The user may perform calibration verification and calibration by selecting this option (See calibration verification section for detail).
- Spirometry standards recommend checking the accuracy of lung function measuring devices at least daily with a 3-L syringe to validate that the instrument is measuring accurately.
4. About Box

- Information about the software can be obtained from the About box. This information is required for queries to Vitalograph or a service agent.
- Information includes the Model number (6300), Serial number of the device, the Software reference number, date of the last Calibration verification and the Service Due date.

3.6.1. Parameters

Parameters available in the Vitalograph micro:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>VC</td>
<td>Vital capacity (L)</td>
</tr>
<tr>
<td>FVC</td>
<td>Forced vital capacity (L)</td>
</tr>
<tr>
<td>FEV1</td>
<td>Forced expiratory volume after 1 second (L)</td>
</tr>
<tr>
<td>FEV1R</td>
<td>FEV1 divided by the largest VC from the VC or FVC manoeuvre.</td>
</tr>
<tr>
<td>PEF L/s</td>
<td>Peak expiratory flow (L/sec)</td>
</tr>
<tr>
<td>PEF L/min</td>
<td>Peak expiratory flow (L/min)</td>
</tr>
<tr>
<td>FEF25-75</td>
<td>Maximal mid expiratory flow: the mean FEF in the time interval between 25% and 75% of the FVC (L/sec)</td>
</tr>
<tr>
<td>FEF75-85</td>
<td>Forced late expiratory flow: the mean FEF in the time interval between 75% and 85% of the FVC (L/sec)</td>
</tr>
<tr>
<td>EVC</td>
<td>Expiratory vital capacity (L)</td>
</tr>
<tr>
<td>IVC</td>
<td>Inspiratory vital capacity (L)</td>
</tr>
<tr>
<td>Term</td>
<td>Description</td>
</tr>
<tr>
<td>------</td>
<td>-------------</td>
</tr>
<tr>
<td>FIVC</td>
<td>Forced inspiratory vital capacity (L)</td>
</tr>
<tr>
<td>FIVC/FVC</td>
<td>Ratio FIVC of FVC</td>
</tr>
<tr>
<td>FEV.5</td>
<td>Forced expiratory volume after 0.5 seconds (L)</td>
</tr>
<tr>
<td>PIF L/s</td>
<td>Peak inspiratory flow (L/sec)</td>
</tr>
<tr>
<td>FMFT</td>
<td>Forced mid-expiratory flow time (sec)</td>
</tr>
<tr>
<td>FET</td>
<td>Forced expiratory time (sec)</td>
</tr>
<tr>
<td>FEV.5/FVC</td>
<td>Ratio FEV 0.5 of FVC</td>
</tr>
<tr>
<td>FEV.75</td>
<td>Forced expiratory volume after 0.75 seconds (L)</td>
</tr>
<tr>
<td>FEV.75/FVC</td>
<td>Ratio FEV 0.75 of FVC</td>
</tr>
<tr>
<td>FEV1/VC</td>
<td>Ratio FEV1 of VC</td>
</tr>
<tr>
<td>FEV1/IVC</td>
<td>Ratio FEV1 of IVC</td>
</tr>
<tr>
<td>FEV1/FVC</td>
<td>Ratio FEV1 of FVC</td>
</tr>
<tr>
<td>FEV1/FIVC</td>
<td>Ratio FEV1 of FIVC</td>
</tr>
<tr>
<td>FEV1/FEV6</td>
<td>Ratio FEV1 of FEV6</td>
</tr>
<tr>
<td>FEV1/PEF</td>
<td>Ratio FEV1 of PEF</td>
</tr>
<tr>
<td>FEV3</td>
<td>Forced expiratory volume after 3 seconds (L)</td>
</tr>
<tr>
<td>FEV3/VC</td>
<td>Ratio FEV3 of VC</td>
</tr>
<tr>
<td>FEV3/FVC</td>
<td>Ratio FEV3 of FVC</td>
</tr>
<tr>
<td>FEV6</td>
<td>Forced expiratory volume after 6 seconds (L)</td>
</tr>
<tr>
<td>FEF25</td>
<td>Forced expiratory flow at 25% of the FVC (L/sec)</td>
</tr>
<tr>
<td>FEF50</td>
<td>Forced expiratory flow at 50% of the FVC (L/sec)</td>
</tr>
<tr>
<td>FEF75</td>
<td>Forced expiratory flow at 75% of the FVC (L/sec)</td>
</tr>
<tr>
<td>FEF0.2-1.2</td>
<td>Mean forced expiratory flow in the volume interval between 0.2 and 1.2 L of the test (L/sec)</td>
</tr>
<tr>
<td>FEF 25-75/FVC</td>
<td>Ratio FEF25-75 of FVC</td>
</tr>
<tr>
<td>FIV1</td>
<td>Forced inspiratory volume after 1 second (L)</td>
</tr>
<tr>
<td>FIV1/FVC</td>
<td>Ratio FIV1 of FVC</td>
</tr>
<tr>
<td>FIV1/FIVC</td>
<td>Ratio FIV1 of FIVC</td>
</tr>
<tr>
<td>PIF L/min</td>
<td>Peak inspiratory flow (L/min)</td>
</tr>
<tr>
<td>FIF25</td>
<td>Forced inspiratory flow at 25% of the FVC (L/sec)</td>
</tr>
<tr>
<td>FIF50</td>
<td>Forced inspiratory flow at 50% of the FVC (L/sec)</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>FIF75</td>
<td>Forced inspiratory flow at 75% of the FVC (L/sec)</td>
</tr>
<tr>
<td>FIF50/FEF50</td>
<td>Ratio FIF 50% of FEF 50%</td>
</tr>
<tr>
<td>FEF50/FIF50</td>
<td>Ratio FEF 50% of FIF 50%</td>
</tr>
<tr>
<td>MVVind</td>
<td>Maximum voluntary ventilation indirectly calculated from the FEV1 (L/min)</td>
</tr>
<tr>
<td>Rind</td>
<td>Airways Resistance Indirect measurement.</td>
</tr>
<tr>
<td>Vext</td>
<td>Extrapolated volume (L)</td>
</tr>
<tr>
<td>Vext/FVC</td>
<td>Ratio Vext to FVC</td>
</tr>
<tr>
<td>FEV1/EVC</td>
<td>Ratio FEV1 to EVC</td>
</tr>
</tbody>
</table>

4. **Power Management in the Vitalograph micro**

The Vitalograph micro can be powered from a computer via the USB cable or from its internal batteries. The LED on the front face of the device and the battery power icon show the power status of the device.

When powered from USB power a power supply icon will be displayed on the status bar at the top of the screen and the LED on the device shows green.

4.1. **Batteries**

- **When the batteries are full**, a Green “Battery full” Icon is displayed on the Main Menu screen device. The LED will show green when the device battery is full.

- **When the batteries start to run low**, an orange “Battery Low” Icon is displayed and the LED will show orange. It is possible to continue to use the device. It is advised that the batteries are changed or USB cable connected to a computer in order to continue testing.
When the batteries are approaching fully discharged the Battery Discharged icon will be displayed on the full screen on power up and on the main screen status bar and both it and the LED will turn red. It is advised that you change the batteries or attach to a PC using a USB cable to continue testing.

The Vitalograph micro uses 4 x 1.5V non-rechargeable IEC60086 certified AAA alkaline batteries. This allows the device to be used without the USB cable connected to the device.

4.2. Power Save Mode

During battery only use the screen on the Vitalograph micro device will dim after 30 seconds, go blank after 60 seconds and the device will automatically power down after 2 minutes if left unused. When powered via USB the screen on the Vitalograph micro device will go blank if left unused for 5 minutes. The device will not automatically power down if powered by USB. Pressing the On/Off button will bring the device out of power save mode.

5. Cleaning & Hygiene

5.1. Preventing Cross-Contamination of Subjects

A spirometer is not designed or supplied as a ‘sterile’ device. Vitalograph intends that a new Bacterial Viral Filter (BVF) be used for every subject to prevent cross contamination. Using a new BVF provides a significant level of protection for the subject, the device and the user against cross contamination during spirometry manoeuvres.

The interior of a Vitalograph flowhead does not require decontamination where a new BVF is used for each subject. The outside surfaces of the device and flowhead tube may be cleaned with a 70% isopropyl alcohol impregnated cloth to remove any visible soiling and for low level disinfection. If you suspect the flowhead has become contaminated or where user risk assessment identifies a need for higher level of
decontamination, then it should be cleaned as per the instructions on ‘Cleaning and Hygiene’ on the Vitalograph website.

5.2. Inspection of the Vitalograph micro

A visual inspection is recommended on a routine basis. Remove the flowhead cone from the flowhead. Examine the mesh part of the flowhead cone for damage or contamination. If it is damaged or blocked, discard and replace with a new part. Re-assemble the cone and flowhead body.

It is recommended that calibration verification is carried out following cleaning and re-assembly as recommended in the ATS/ERS 2019 guidelines².

6. Remote Flowhead

The micro flowhead can be set up to work remotely from the device. This may be useful if the display need to be monitored, while the patient uses the device.

² ATS/ERS Standardisation of Spirometry 2019 Update Am J Respir Crit Care Med 2019 Vol 200, Iss 8 pp e70-e88
1. Press and hold the Flowhead release button and slide the flowhead away from the device from left to right.
2. From the remote flowhead adaptor kit, attached the device cap in the space that was occupied by the flowhead.
3. Attach the remote flowhead adaptor to the flowhead. This is done by sliding the flowhead into the grooves in the remote flowhead adaptor. Ensure this is fully pushed in.
4. Attach the remote flowhead to the port on the micro device cap using the Flowhead Connection Tube.
5. It is recommended that an accuracy check is carried out before the flowhead is used remotely or refitted to verify correct operation and accuracy.

7. Fault Finding Guide

| Problem Fault Symptoms: | • Calibration verification variations > +/-3%  
|                        | • Error at last Calibration Verification  
|                        | • Accuracy / Calibration Fail  
|                        | • False readings suspected |

<table>
<thead>
<tr>
<th>Possible Solutions: (In probable order)</th>
</tr>
</thead>
</table>
| • Recheck Accuracy/Calibration  
| • Was the correct syringe volume selected?  
| • Calibration verification is required after cleaning/disinfecting the flowhead assembly.  
| • Flowhead body pressure port holes/ grommets blocked.  
| • Flowhead Fleisch element not dried thoroughly.  
| • Flowhead Fleisch element assembly blocked.  
| • Main PCB failure – contact support. |

<table>
<thead>
<tr>
<th>Problem Fault Symptoms:</th>
</tr>
</thead>
</table>
| • Test begins automatically  
| • Volume accumulates automatically without the subject blowing.  
<p>| • Very small VC or FVC test displayed |</p>
<table>
<thead>
<tr>
<th>Possible Solutions: (In probable order)</th>
<th>• Flowhead and/or tubing when using remote flowhead not stationary at the start of test. Hold them steady until the 'Blow Icon' appears. • Return to Main Menu and re-enter the test routine.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Problem Fault Symptoms:</strong></td>
<td>• No volume measurements.</td>
</tr>
<tr>
<td>Possible Solutions: (In probable order)</td>
<td>• Ensure that the grommets on flowhead are not pinched or trapped.</td>
</tr>
<tr>
<td><strong>Problem Fault Symptoms:</strong></td>
<td>• Cannot print to PC (Vitalograph Reports Application). • Corrupt or missing data on printout.</td>
</tr>
<tr>
<td>Possible Solutions: (In probable order)</td>
<td>• Check USB cable is connected between Vitalograph micro and the PC. • Check to ensure the Vitalograph Reports Application is correctly installed. • Check to ensure the required software drivers are installed on the PC. • Main PCB failure – contact support.</td>
</tr>
<tr>
<td><strong>Problem Fault Symptoms:</strong></td>
<td>• Cannot read screen.</td>
</tr>
<tr>
<td>Possible Solutions: (In probable order)</td>
<td>• The batteries may be low. Plug in the USB cable and switch on the device. • LCD failure – contact support. • Main PCB failure – contact support.</td>
</tr>
</tbody>
</table>

1. Medical Devices may be affected by mobile RF communications equipment including cellular telephones and other personal or household devices not intended for medical facilities. It is recommended that all equipment used near the Vitalograph product comply with the medical electromagnetic compatibility standard and to check before use that no interference is evident or possible. If interference is suspected or possible, switching off the offending device is the normal solution, as is required in aircraft and medical facilities.
2. Portable RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm (12 inches) to any part of the micro, including cables specified by Vitalograph. Otherwise, degradation of the performance of this equipment could result.

3. Use of this equipment adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, this equipment and the other equipment should be observed to verify that they are operating normally.

8. **Customer Service**

Service and repairs should be carried out only by the manufacturer, or by Service Agents specifically approved by Vitalograph. Vitalograph makes instructions and parts available to designated service agents as required. For the names and addresses of approved Vitalograph Service Agents or to arrange spirometry workshops, please refer to the contact information at the start of this manual.

Any serious incident that has occurred in relation to the device should be reported to Vitalograph or its Authorized Representative and the Regulatory Authorities of the country. Refer to the Vitalograph contact information at the start of this manual.

9. **Consumables and Accessories**

<table>
<thead>
<tr>
<th>Cat. No</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>28350</td>
<td>BVF - Bacterial/Viral Filters (50)</td>
</tr>
<tr>
<td>28501</td>
<td>Eco BVF – Bacterial/Viral Filters (100)</td>
</tr>
<tr>
<td>28572</td>
<td>Eco BVF and Disposable Nose Clip (80)</td>
</tr>
<tr>
<td>28554</td>
<td>Eco BVF with Bite Lip (75)</td>
</tr>
<tr>
<td>28553</td>
<td>Eco BVF with Bite Lip and Disposable Nose Clip (75)</td>
</tr>
<tr>
<td>36020</td>
<td>3-L Precision Syringe</td>
</tr>
<tr>
<td>79158</td>
<td>Flow Cone (10)</td>
</tr>
<tr>
<td>79191</td>
<td>Flowhead Complete</td>
</tr>
</tbody>
</table>
10. **Disposal**

The device must be taken to separate collection at the product end-of-life. Do not dispose of these products as unsorted municipal waste.

Used BVFs constitute minimally soiled waste from human healthcare and should be disposed of in line with local requirements. BVFs are made from polypropylene.

11. **Explanation of Symbols**

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Type B equipment" /></td>
<td>Type B equipment</td>
</tr>
<tr>
<td><img src="image" alt="Class II" /></td>
<td>Class II</td>
</tr>
<tr>
<td><img src="image" alt="Power rating" /></td>
<td>Power rating</td>
</tr>
<tr>
<td><img src="image" alt="Direct current" /></td>
<td>Direct current</td>
</tr>
<tr>
<td><img src="image" alt="Instruction for Use; operating instructions" /></td>
<td>Instructions for Use; operating instructions</td>
</tr>
<tr>
<td><img src="image" alt="Manufacturer" /></td>
<td>Manufacturer</td>
</tr>
<tr>
<td><img src="image" alt="Date of Manufacture (include date in format yyyy-mm-dd)" /></td>
<td>Date of Manufacture (include date in format yyyy-mm-dd)</td>
</tr>
<tr>
<td><img src="image" alt="USB connector" /></td>
<td>USB connector</td>
</tr>
<tr>
<td><img src="image" alt="The device must be taken to separate collection at the product end-of-life. Do not dispose of these products as unsorted municipal waste" /></td>
<td>The device must be taken to separate collection at the product end-of-life. Do not dispose of these products as unsorted municipal waste</td>
</tr>
</tbody>
</table>
Fragile, handle with care

Keep Dry

Do not re-use

Non sterile

Recycle

QR code - matrix bar code. All information in the bar code is included in the text under it

Use by Date (Date format yyyy-mm-dd)

Device Order Number

Lot/Batch Number

Serial Number

On/Off Button

Battery Positive

### 11.1. Icons used in the Vitalograph micro

<table>
<thead>
<tr>
<th>Icon</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Subject</td>
<td>VC Test</td>
</tr>
<tr>
<td>FVC Test</td>
<td>Post Test</td>
</tr>
<tr>
<td>Settings</td>
<td>Enter</td>
</tr>
<tr>
<td>----------------------------------------------</td>
<td>---------------------------</td>
</tr>
<tr>
<td>Subject Options</td>
<td>Device Settings</td>
</tr>
<tr>
<td>Accuracy / Calibration</td>
<td>Device and software information</td>
</tr>
<tr>
<td>Age</td>
<td>Height</td>
</tr>
<tr>
<td>Gender- Male</td>
<td>Gender- Female</td>
</tr>
<tr>
<td>Posture - Sitting</td>
<td>Posture - Standing</td>
</tr>
<tr>
<td>Weight On</td>
<td>Weight Off</td>
</tr>
<tr>
<td>Results Options</td>
<td>Parameters</td>
</tr>
<tr>
<td>Time/Date</td>
<td>Service Mode</td>
</tr>
<tr>
<td>VC Volume-Time Graph</td>
<td>VC Volume Graph</td>
</tr>
<tr>
<td></td>
<td>Zoom Out</td>
</tr>
<tr>
<td>---</td>
<td>----------</td>
</tr>
<tr>
<td></td>
<td>Test Results</td>
</tr>
<tr>
<td></td>
<td>FVC Volume-Time Graph</td>
</tr>
<tr>
<td></td>
<td>Delete</td>
</tr>
<tr>
<td></td>
<td>Serial Number</td>
</tr>
<tr>
<td></td>
<td>Syringe Volume</td>
</tr>
<tr>
<td></td>
<td>USB Power</td>
</tr>
<tr>
<td></td>
<td>Battery Low</td>
</tr>
<tr>
<td></td>
<td>Z Score</td>
</tr>
<tr>
<td></td>
<td>Sound On</td>
</tr>
<tr>
<td></td>
<td>User Passcode - On/Locked</td>
</tr>
</tbody>
</table>
12. Description of the Vitalograph micro

The Vitalograph micro is a handheld spirometer which measures subject respiratory parameters including but not limited to VC, FVC, FEV1, PEF and MVV. The Vitalograph micro is designed for portable spirometry. The Fleisch flowhead is used for testing and is integral to the device, although it may be removed to use with an adaptor for remote testing.

12.1. Indications for Use

The indications for use of the Vitalograph micro is in the assessment of lung function through the measurement of dynamic lung volumes, i.e. spirometry.

The Vitalograph micro is designed to be operated by medical professionals trained in respiratory and lung function testing on adults and paediatrics, 2.5 years and older, in a variety of professional healthcare environments, e.g. primary care, hospitals and occupational health centres. The measurements obtained from a lung function test provide objective information used in the diagnosis of lung diseases and monitoring lung health.
## 13. Technical Specification

<table>
<thead>
<tr>
<th>Product</th>
<th>Vitalograph micro, Model 6300</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flow Detection Principal</td>
<td>Fleisch type pneumotachograph</td>
</tr>
<tr>
<td>Volume detection</td>
<td>Flow integration sampling at 100Hz</td>
</tr>
<tr>
<td>Maximum test duration</td>
<td>90 seconds</td>
</tr>
<tr>
<td>Maximum displayed volume</td>
<td>10 L</td>
</tr>
<tr>
<td>Volume Accuracy</td>
<td>Better than ±3% or ±0.05L of the reading. (ISO 26782:2009)</td>
</tr>
<tr>
<td>Flow Measurement Range</td>
<td>Max. flow rate ±960 L/min (±16 L/s) Min. flow rate ±1.2 L/min (±0.02 L/s)</td>
</tr>
<tr>
<td>PEF Accuracy</td>
<td>±10% or ±10L/min of the reading (ISO 23747:2015)</td>
</tr>
<tr>
<td>Back pressure</td>
<td>Less than 0.1kPa/L/sec at 14L/sec (ATS/ERS 2005)</td>
</tr>
<tr>
<td>Operating temperature range</td>
<td>ISO 26782 limits: 17–35°C</td>
</tr>
<tr>
<td>Operating humidity range</td>
<td>30%–75%</td>
</tr>
<tr>
<td>Ambient pressure range</td>
<td>850hPa–1060hPa</td>
</tr>
<tr>
<td>Vitalograph micro meets or</td>
<td></td>
</tr>
<tr>
<td>exceeds</td>
<td></td>
</tr>
<tr>
<td>EMC Standards</td>
<td>EN 60601-1-2: 2015</td>
</tr>
<tr>
<td>QA/GMP standards</td>
<td>EN ISO 13485, FDA 21 CFR 820, CMDR SOR/98-282 &amp; JPAL, MDSAP</td>
</tr>
<tr>
<td>Dimensions</td>
<td>83mm x 91mm x 32mm</td>
</tr>
<tr>
<td>Weight</td>
<td>250g</td>
</tr>
</tbody>
</table>
| **Communications** | USB & Bluetooth option available  
*(Note: Vitalograph reports is not enabled for Bluetooth communications with this device)* |
|---------------------|--------------------------------------------------------------------------------|
| **Power Supply**    | 4 x 1.5V AAA batteries (6V)  
5V DC via USB 2.0/3.0 |
| **Minimum PC System Requirements to run Vitalograph Reports** |  
Processor: Pentium, Celeron CPU, 1.73 GHz or better  
Operating Systems: Windows® 7 or higher  
Memory: 128 MB of RAM, 256 MB recommended  
Hard Disk: 40MB for the Vitalograph Reports application  
280MB for the .NET framework  
Display: A display supporting a resolution of 1280 x 800 pixels, higher recommended.  
Other:  
• Installation of the .NET Framework 3.5  
• CD-ROM drive  
• USB Port (to connect device)  
• Install the application as System Administrator and provide full read/write access rights to the folder and sub-folders where the application has been installed, for all applicable users.  
• Internet Explorer 8.0 or above required |
Notes:
• Class II device when powered by USB. Otherwise is internally powered device.
• ME system is created when the device is connect to a supporting PC via USB. This system made from the following parts: Supporting PC, USB Cable and micro Device.
• The BVF, flowhead and device body are type BF applied part. An applied part is a part of the equipment, that in normal use necessarily comes into physical contact with the subject for equipment or system to perform its function.

14. Contraindications, Warnings, Precautions and Adverse Reactions

1. No modification of this ME equipment or ME system is allowed, except for connecting and disconnecting of the USB interface. Any unauthorised changes to the Vitalograph micro device may compromise product safety and/or data and as such Vitalograph cannot be held responsible and the device will no longer be supported.

2. The Vitalograph micro is not designed as a sterile device. Always follow the safety guidelines given by the manufacturer of cleaning and disinfectant chemicals.

3. For the device to be used as intended, there is no requirement to clean any supporting computer. If cleaning is required to remove any visible soiling, this should be done as per the computer manufacturer’s instructions.

4. Vitalograph intends that a new Bacterial Viral Filter (BVF™) be used for every subject to prevent cross contamination. Using a new BVF provides a significant level of protection of the subject, the device and the user against cross contamination during spirometry manoeuvres. A BVF is for single use only.

5. Spirometry is a valuable tool that provides important information to clinicians which is used together with other physical findings, symptoms, and history to reach a diagnosis (ATS/ERS 2019).

6. When using the Vitalograph micro with a remote flowhead adaptor ensure that the flowhead connecting tube is not pinched or trapped as spirometry results may appear to be
inverted.
7. Take care not to block the mouthpiece with tongue or teeth during testing. A ‘spitting’ action or cough will give false readings.
8. Subject fatigue may occur during spirometry testing depending on the subject’s characteristics e.g. age, health status. For safety reasons, testing should be preferably done in the sitting position, using a chair with arms and without wheels. Subject may also take a break between tests. A subject fatigue warning will appear after 8 manoeuvres and the maximum number of allowed manoeuvres in one session is 20.
9. All values displayed are expressed as BTPS values
10. Time zero is determined using the back-extrapolated method, from the steepest part of the curve.
11. After fitting a New Flowhead to the Vitalograph micro it is necessary to set-up the flowhead with the device. The procedure is outlined in section 3.6.
12. Do not expose the micro to liquids, with the exception of 70% Isopropyl wipes for cleaning as detailed in Section 5 Cleaning & Hygiene.
13. The micro should not be used in the presence of flammable liquids or gases, dust, sand or any other chemical substances.
14. All spirometry standards recommend checking the accuracy of lung function measuring devices daily with a 3-L syringe to validate that the instrument is measuring accurately. The Vitalograph micro should never be outside accuracy limits. Accuracy should be checked after cleaning or disassembling the spirometer for any reason, after adjusting calibration or if the flowhead or device has been dropped.
15. Service and repairs should be carried out only by the manufacturer or by Service Agents specifically approved by Vitalograph.
16. Maintenance must not be performed while the device is in use by a subject.
17. The device contains uses 4 x 1.5V non-rechargeable IEC60086 certified AAA alkaline batteries.
18. Use of accessories, parts and cables other than those specified or provided by Vitalograph for this equipment is not
recommended.
19. Non-medical equipment must be kept outside the subject environment i.e. any area in which intentional or unintentional contact between the subject and parts of the system, or some other persons touching part of the system, can occur.
20. Including the BVF, the subject may contact any part of the device during a spirometry session. There are no adverse effects if the subject comes into contact with any part of the micro device.
21. The AAA batteries should be removed, if the device is intended to be stored, without use, for an extended period of time.
22. Reprocessing of single use devices is not permitted.
23. Non-ME equipment used with the device, should comply with its relevant IEC or ISO standard and also supplied via a suitably approved PSU.
24. The operator should not contact to patient while simultaneously contacting any of the following: Batteries, components within battery compartment and USB connector (whilst the USB cable is connected to the supporting PC) or supporting PC’s connectors

15. CE Notice
Marking by the symbol \( \bigcirc \) indicates compliance of the Vitalograph Model 6300 micro to the Medical Devices Directive of the European Community.
The Vitalograph Model 6300 micro is intended for use in a variety of professional healthcare environments, e.g. primary care, hospital wards and occupational health centres, except for near active high frequency surgical equipment and the RF shielded room of an ME system for magnetic resonance imaging, where the intensity of electromagnetic disturbance is high. The customer or the user of the Vitalograph micro should assure that it is not used in such an environment.
The Vitalograph Model 6300 micro has been tested in accordance with:


### EN 60601-1-2:2015 - Emissions tests

<table>
<thead>
<tr>
<th>Emissions test</th>
<th>Compliance</th>
<th>Electromagnetic environment - guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions CISPR 11</td>
<td>Group 1</td>
<td>The Model 6300 micro 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.</td>
</tr>
<tr>
<td>RF emissions CISPR 11</td>
<td>Class B</td>
<td>The Model 6300 micro is suitable for use in all establishments, including domestic establishments and those connected to the public mains network (e.g. at home and doctor’s offices in residential areas)</td>
</tr>
</tbody>
</table>

### EN 60601-1-2:2015 - Immunity tests

<table>
<thead>
<tr>
<th>Immunity test</th>
<th>Test level</th>
<th>Compliance level reached</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic discharge (ESD) EN 61000-4-2</td>
<td>Contact: ±8kv</td>
<td>Contact: ±8kv</td>
</tr>
<tr>
<td></td>
<td>Air: ±15kv</td>
<td>Air: ±15kv</td>
</tr>
<tr>
<td>Power frequency (60 Hz) magnetic field EN 61000-4-8</td>
<td>30A/m</td>
<td>30A/m</td>
</tr>
<tr>
<td>Radiated RF EN 61000-4-3</td>
<td>3 V/m 80MHz to 2700MHz</td>
<td>3 V/m 80MHz to 2700 MHz</td>
</tr>
</tbody>
</table>
Medical electrical equipment needs special precautions regarding EMC and needs to be installed and put into service according to the EMC information provided.

16. FDA Notice
Caution: Federal Law restricts this device to sale by, or on the order of a physician.

17. EU Declaration of Conformity
Product: Vitalograph micro™, model 6300

Vitalograph hereby ensures and declares that the above product associated with these instructions for use, is designed and manufactured in accordance with the following QMS regulations and standards:

• European Medical Devices Directive {MDD} 93/42/EEC, as amended.
  This device is classified as IIa per Annex IX of the MDD also meets the provisions of the Essential Requirements, Annex I, via compliance with Annex II of the Medical Devices Directive as per Article 11, section 3a, excluding point 4 of Annex II.

• Canadian Medical Device Regulation {SOR/98-282}
• FDA Quality System Regulation {QSR} 21 CFR 820.
• EN ISO 13485 Medical devices. Quality management systems. Requirements for regulatory purposes.

Certifying Body: British Standards Institute {BSI}.
BSI Notified Body #: 2797
Certificate Nos. CE 00772, CE 85553, MD 82182.

Signed on behalf of Vitalograph (Ireland) Ltd.

Frank Keane.
CEO, Vitalograph Ltd.
18. Guarantee

Subject to the conditions listed below, Vitalograph Ltd. and its associated companies, (hereinafter called the Company) guarantee to repair or at its option replace any component thereof, which, in the opinion of the Company is faulty or below standard as a result of inferior workmanship or materials.

The conditions of this Guarantee are:

1. This Guarantee shall only apply to hardware defects which are notified to the Company or to its accredited distributor within 1 year of the date of purchase of the equipment, unless otherwise agreed in writing by the Company.
2. Software (meaning computer software, or user installable modules) is guaranteed for 90 days from the date of purchase.
3. The Company warrants that the software when correctly used in conjunction with the hardware will perform in the manner described in the Company’s literature and user manuals. The Company undertakes to rectify at no expense to the customer any software failure notified within the period stated above, provided that the failure can be recreated and the software has been installed and used in accordance with the user manual. Notwithstanding this clause, the software is not warranted to be free of errors.
4. This Guarantee does not cover any faults caused by accident, misuse, neglect, tampering with the equipment, use of consumable items or parts not approved by the Company, or any attempt at adjustment or repair other than by personnel accredited by the Company, nor does it cover reinstatement of any configuration changes caused by the installation of any software.
5. If a defect occurs please contact the supplier from it was purchased for advice. The Company does not authorize any person to create for it any other obligation or liability in connection with Vitalograph equipment.
6. This Guarantee is not transferable and no person, firm or company has any authority to vary the terms or conditions of this guarantee.
7. To the maximum extent permitted by law, the Company does not accept liability for any consequential damages arising out of the use of, or inability to use any Vitalograph® equipment.
8. This Guarantee is offered as an additional benefit to the Consumer’s statutory rights and does not affect these rights in any way.