Resting Metabolic Rate Test
Metabolism

Metabolism can be understood as the conversion by the human body between food and accumulated fat into energy. The energy is used by the body to maintain constant temperature, to move and to make all the organ function. Measure of metabolism is: calories (cal).

Total Metabolic Rate

The total metabolic rate are the total calories that the human body needs in order to actuate the daily functional activities.

Resting Metabolic Rate (RMR)

Resting Metabolic Rate represents the calories that the vital organs need to properly operate at rest (heart, brain, lungs, liver, kidneys etc.). RMR represents between 60% and 75% of the human’s total metabolism.

Importance to measure RMR

A knowledge of the RMR is very helpful in order to understand the nutritional needs and to properly manage it.

Measure of the rest metabolic rate with indirect calorimetry

Energy expenditure can be measured directly by putting a person in a calorimeter and measuring the amount of heat produced by the body mass. This is expensive and very impractical in the clinical setting. Energy expenditure can be measured indirectly with a metabolic cart by analysis of respired gases (usually expired) to derive volume of air passing through the lungs, the amount of oxygen extracted from it (i.e., oxygen uptake VO\(_2\)) and the amount of carbon dioxide, as a by-product of metabolism, expelled to atmosphere (CO\(_2\) output – VCO\(_2\)). With these measurements the resting energy expenditure (RMR) and respiratory quotient (RQ) can be calculated. The RQ represents the ratio of carbon dioxide exhaled to the amount of oxygen consumed by the individual. RQ is useful in interpreting the results of the RMR. The abbreviated Weir equation is probably the most common calculation of RMR.

*Abbreviated Weir equation:*

\[
\text{RMR} = [3.9 \ (\text{VO}2) + 1.1 \ (\text{VCO}2)] \ 1.44
\]

How to perform a RMR test

For best results, when having a REE done, there are certain conditions that need to be controlled and others that just require documenting at the time of the test. During the test the individual is interfaced with a metabolic measurement system by means of a facemask. A mouthpiece with a nose clip is also sometimes used, but it may create overly stressful conditions to a subject (patient).

Important considerations or conditions to improve the RMR measurement:

- No food for at least 12 hours and no smoke for at least 2 hours before the test.
- Maintain quiet surroundings when the test is in progress and normal temperature. The individual should not move arms or legs during the test.
- Medications taken should be noted, such as stimulants or depressants.
- The first 5 minutes of acquisition should be discarded by the computation of RMR.
- Steady state should be achieved, which would be identified clinically by the following criteria: 5 minute period when average minute VO\(_2\) and VCO\(_2\) changes by less than 10%, average RQ changes by less than 5%
- Stable interpretable measurements should be obtained in a 15 to 20 minute test.
- Renal failure patients requiring hemodialysis should not be tested during dialysis therapy.
Recommendations

Resting metabolic rate test using the face mask

1. Since the ventilation is very low (normally <10 litres/min), the turbine calibration has to be performed with very slow manoeuvres (each complete manoeuvre in about 10-15 seconds), to obtain the best accuracy.
2. Use the following correction for the dead space (VD):
   - 50 ml for the small mask
   - 60 ml for the medium mask
   - 70 ml for the large mask

Resting metabolic rate test using the canopy option

1. Verify, before and during the test, that the FeCO$_2$ falls into the range 0.5%-0.8% and adjust the flow rate of the pump. If the FeCO$_2$ is too low, increase the flow, if it is too high decrease the flow. In fact, if the FeCO$_2$ is too low the measurement could be not reliable, while an high FeCO$_2$ could be dangerous for the patient.
2. In order to perform a correct ERGO calibration and to obtain more reliable data from the test, it is recommended to use a calibration cylinder with the following concentrations: 1% CO$_2$, 20% O$_2$, balance N$_2$. If you use this cylinder, please remember to modify the reference values, as explained in the chapter Calibration.
Performing a test using the face mask

Calibrations

Before the test, it is necessary to perform an ergo calibration (see Calibration chapter) and it is advisable to perform also a turbine calibration (see Recommendations in this chapter).

How to prepare a patient

The patient interfaces with the equipment by means of a face mask, like in the stress exercise. The mask has to be tight to the face, in order to avoid any air leakage.

Start the test

1. Enter in the ergometry program
2. Select a patient or add a new one (File/Patients...)
3. Select Start test from Test menu
4. Enter the patient’s data and select the RMR mode (1st picture).
5. Press Other Data... and enter the dead space value (50ml Small mask, 60ml Medium mask and 70ml Large mask). It is possible to enter the Ureic Nitrogen value NU (2nd picture).
6. Confirm and start the test by pressing OK.

Selecting RMR the system set automatically the following options:

- Data acquisition with a 30 seconds average
- RMR protocol, which is:
  - 5 minutes discarded;
  - 10 minutes with data acquisition, of which the software will make an average at the end of the test;
  - automatic end of the test after the 16th minute.
- Selection of the RMR workspace (windows placement);

The test is fully automatic, the software will stop it and save the data at the end of the 16th minute.

The real time view is as shown in the following picture:
Viewing the test

At the end of the test, it will be opened automatically a window with the test results.

At the end of the test, or if it is selected View/RMR, the main results are shown:

- The average time interval (default: 10 minutes)
- Average values of VO₂, VCO₂, R, RMR, RF, VE, HR, FAT% and CHO% and predicted values if available.
- Body Mass Index (BMI) and interpretation
- Graph of the energetic expenditure for all the data acquisition interval, highlighting the selected average interval.

In order to verify the goodness of the test, check that the ventilation and respiratory frequency are similar to the predicted ones (12 breaths/min for the respiratory frequency and 6 litres/min for the ventilation), and the heart rate is the rest heart rate of the patient.

Nota: The percentage of used Proteins (PRO%) is calculated assuming 12 grams of Ureic Nitrogen in 24 hours. You can modify this value selecting View/Information… -> Modify…
How to modify the average interval

If the average interval (automatically identified by the software) is not satisfying, for example because the patient was speaking in the first minutes, it is possible to modify the interval of the average.

Right-click and select **Edit RMR…** It is possible to move the start and the end lines.

To move the start line, left-click on the exact time in which you want to start the calculations, for the end line, right-click.

Print

The print of the current window generates a report similar to the one in the following page.
**Chapter 8 - Resting Metabolic Rate Test - 101**

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http://www.cosmed.it; E-mail: info@cosmed.it

Last name: BOND  First name: JAMES

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Rest</th>
<th>Pred.</th>
<th>% Pred.</th>
<th>BMI Statements</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time (mm:ss)</td>
<td>09:30</td>
<td>10:00</td>
<td>95.0%</td>
<td>BMI &lt;18.5</td>
</tr>
<tr>
<td>RMR (Kcal/day)</td>
<td>1981.1</td>
<td>1731.9</td>
<td>114.3%</td>
<td>Normal 18.5-24.9</td>
</tr>
<tr>
<td>R (→)</td>
<td>0.82</td>
<td>0.85</td>
<td>96.8%</td>
<td>Overweight 25.0-29.9</td>
</tr>
<tr>
<td>VO2 (ml/min)</td>
<td>286</td>
<td>266</td>
<td>107.7%</td>
<td>Obesity class I 30.0-34.9</td>
</tr>
<tr>
<td>VCO2 (m/l/min)</td>
<td>235</td>
<td>226</td>
<td>104.2%</td>
<td>Obesity class II 35.0-39.9</td>
</tr>
<tr>
<td>Rf (b/min)</td>
<td>13.7</td>
<td>12.0</td>
<td>114.2%</td>
<td>Obesity class III &lt;=40</td>
</tr>
<tr>
<td>VE (l/min)</td>
<td>9.0</td>
<td>6.0</td>
<td>150.4%</td>
<td></td>
</tr>
<tr>
<td>HR (bpm)</td>
<td>72</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FAT% (%)</td>
<td>59.5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CHO% (%)</td>
<td>40.9</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BMI (Kg/m²)</td>
<td>23.9</td>
<td></td>
<td>Normal</td>
<td></td>
</tr>
</tbody>
</table>

---

**Graph**

---

20/02/2003 15:24
Performing a test using the canopy option

The principle of a ventilated bubblehood system is that a stream of air is forced to pass across the face of a subject and mixes with the air which is collected by a transparent hood, placed over the subject’s head. A measurement system, knowing the flow rate, calculates the oxygen consumption and the CO$_2$ production and, starting from these values, the energy expenditure.

Calibrations

Before the test, it is necessary to perform an ergo calibration (see Calibration chapter) and it is advisable to perform also a turbine calibration (see Recommendations in this chapter).

How to prepare the canopy and the patient

Replacement of the power plug
If the power plug does not fit into the mains socket, replace it with the one in the packaging.
In order to replace the plug:
1. Extract the plug from the battery charger
2. Insert the proper plug in the battery charger.

Connecting the Canopy
1. Connect the Canopy unit to the mains by means of the medical grade AC/DC adapter provided.
2. Insert the bubblehood adapter into the bubblehood from the outside and fix it screwing the ring from the inside, being careful to insert it in the proper hole, as shown in the following picture.
3. Connect the bubblehood to the wrinkled tube, interposing a bacterial filter.
4. Connect the wrinkled tube to the unit through the Flow in connector.
5. Connect the optoelectronic reader of the Quark to the Flow out connector of the Canopy unit through the spirometry adapter.
6. Fix the vail to the bubblehood through the velcro strips.
How to prepare the patient

1. Switch on the Canopy unit. If there are no problems, the red led on the front panel of the unit flashes for few seconds and the alarm beeps. If the led does not flash and/or the alarm does not beep, the test cannot be performed, because the backup battery is exhausted or there is no backup battery.

2. When the green led turns on, the test can start. If the green led does not turn on, the red led flashes and the alarm beeps, the test cannot be performed because the pump does not work or the mains does not power the system.

3. After these checks, put the patient in a supine position.

4. Place the bubblehood with the vail on the patient’s head. The tube has to be placed near the patient’s mouth.

Performing the test

1. Enter in the ergometry program

2. Select a patient or add a new one (File/Patients...)

3. Select Start test from Test menu.

4. Enter the patient’s data and select the Canopy mode.

5. Confirm and start the test by pressing OK.

6. In the first part of the test the flow rate of the pump has to be adjusted by means of the Flow adjustment handle on the front panel of the Canopy unit, in order to measure an FeCO$_2$ between 0.5% and 0.8%. FeCO$_2$ values can be read on the right side of the PC monitor.
7. When the FeCO₂ remains within the acceptability range, press F₂ to start the data acquisition. Verify, also during the test, that the measured FeCO₂ is within the 0.5%-0.8% range. Otherwise, adjust it by means of the Flow adjustment handle.

**Warning:** If the green led turns off during the test, the red led flashes and the alarm beeps, abort the test, because the pump does not work or the mains does not power the system. In the last case, the pump works only because of the backup battery.

The test is fully automatic, the software will stop it and save the data at the end.

**Viewing the test**

At the end of the test, it will be opened automatically a window with the test results. At the end of the test, or if it is selected View/RMR, the main results are shown:

- The average time interval (default: 10 minutes)
- Average values of VO₂, VCO₂, R, RMR, VE, HR, FAT% and CHO% and predicted values if available.
- Body Mass Index (BMI) and interpretation
- Graph of the energetic expenditure for all the data acquisition interval, highlighting the selected average interval.

In order to verify the goodness of the test, check that the FeO₂ and FeCO₂ values are within the acceptability ranges (20.2%-20.8% and 0.5%-0.8% respectively), and the heart rate is the rest heart rate of the patient.

**How to modify the average interval**

If the average interval (automatically identified by the software) is not satisfying, for example because the patient was speaking in the first minutes, it is possible to modify the interval of the average.

Right-click and select Edit RMR…. It is possible to move the start and the end lines.

To move the start line, left-click on the exact time in which you want to start the calculations, for the end line, right-click.

**Print**

The print of the current window generates a report similar to the one of the RMR test using the face mask.

---

**Note:** The percentage of used Proteins (PRO%) is calculated assuming 12 grams of Ureic Nitrogen in 24 hours. You can modify this value selecting View/Information… -> Modify…
Sub-maximal Exercise Testing
Introduction

Several physiological responses to exercise are used to evaluate cardiorespiratory fitness, including oxygen consumption, heart rate, and blood pressure. Measuring these variables during exercise, particularly maximum exercise, increase the chance of detecting any coronary artery disease or pulmonary disease.

Unfortunately, maximum exercise tests are impractical because they are expensive, require extensive clinical supervision, and subject individuals to levels of physical stress that may be unnecessary depending on the objectives of the test. Consequently, maximal testing is reserved for clinical assessments, athletic evaluation, and research.

A sub-maximal exercise test costs less and carries a lower risk for the individual. Although less sensitive and specific for detecting disease or estimating maximal oxygen consumption, correctly performed sub-maximal tests can provide a valid estimate of cardiorespiratory fitness.

Pre-test screening

Pre-test health screening is essential for risk stratification and for determining the type of test that should be performed and the need for an exercise test prior to exercise training. A thorough pretest health screening includes the following:

- Complete medical history
- Medical contraindications to exercise
- Symptoms suggesting cardiac or pulmonary disease
- Angina or other forms of discomfort at rest or during exercise
- Unusual shortness of breath at rest or during exercise
- Dizziness or light-headedness
- Orthopaedic complications that may prevent adequate effort or compromise the validity of test results
- Other unusual signs or symptoms that may preclude testing
- Risk factors for coronary heart disease
- History of major cardiopulmonary events
- Current medications
- Activity patterns
- Nutritional habits
- Reading and signing an informed consent form
Sub-maximal exercise testing

Heart rate varies linearly with VO₂ to the point of maximum exertion; thus, VO₂max may be estimated using the relation between heart rate and VO₂ without subjecting the individual to maximum levels of physical stress. During sub-maximal exercise testing, predetermined workloads are used to elicit a steady state of exertion (plateau of heart rate and VO₂). The steady-state heart rate at each work level is displayed graphically and extrapolated to the VO₂ at the age-predicted maximal heart rate (HR = 220-age). A variety of protocols for different exercise modalities (i.e., treadmill, stationary cycle, and step increments) can be used as long as the VO₂ requirements of each selected workload can be estimated with accuracy.

The objectives of cardiorespiratory fitness assessments in the apparently healthy population are as follows:

- Determine the level of cardiorespiratory fitness and establish fitness program goals and objectives.
- Develop a safe, effective exercise prescription for the improvement of cardiorespiratory fitness.
- Document improvements in cardiorespiratory fitness as a result of exercise training or other interventions.
- Motivate individuals to initiate an exercise program or comply with an established program.
- Provide information concerning health status.

A few assumptions regarding testing are necessary to ensure the highest degree of accuracy when using sub-maximal exercise testing to estimate VO₂max:

- Selected workloads are reproducible. A steady-state heart rate is obtained during each stage of the test. Usually, workload durations of 3 minutes or more are used to ensure steady state.
- The maximal heart rate for a given age is uniform (HR = 220-age).
- Heart rate and VO₂ have a linear relation over a wide range of values; thus, the slope of HR/VO₂ regression can be extrapolated to an assumed maximum heart rate.
- Mechanical efficiency (i.e., VO₂ at a given work rate) is consistent.

Although if done correctly, sub-maximal exercise tests provide valuable information concerning cardiorespiratory fitness, they have extremely limited diagnostic capabilities and should not be used as a replacement for clinical exercise tests or other clinical treatment or management modalities. Health care professionals should avoid detailed interpretation beyond the scope of the information obtained.

Considerations with sub-maximal exercise testing

Considerations for selection of protocol and equipment include any physical or clinical limitations that may preclude certain types of exercise (i.e., age, weight, arthritis, orthopaedic complications, individual comfort, level of fitness, type of exercise training that will be performed, and individual preference).

For example, some individuals may perform better on a non-weight-bearing modality (cycle versus treadmill), while others may not have the required range of motion in the hip or knee to pedal and may perform better walking. Deconditioned, weak, or elderly persons may have to start the test at a low work level and increase the workload in small increments. Also, field tests may not be appropriate for those who require strict supervision during testing, who do not understand the concept of pacing, or who cannot be expected to put forth a good effort. More consistent results may be obtained by testing in a controlled environment such as a laboratory setting. Creativity when selecting protocols may allow adaptations of commonly used protocols to accommodate athletes competing in specific sports. Regardless of the type of exercise and protocol selected, the same type of exercise and protocol should be used for repeat testing if between-test comparisons are important.
Staffing

Staff members should be able to do the following:
1. Establish rapport with the subject and make him or her feel comfortable.
2. Recognize normal acute and chronic responses to exercise.
3. Recognize abnormal signs and symptoms during exercise.
4. Provide basic life support measures competently.
5. Adhere to established procedures and protocols.
6. Clearly explain test results to the individual.

Test termination

Sub-maximal tests should be terminated according to ACSM or other accepted guidelines (see table in the following). In the event of an abnormal response, the test should be terminated, the medical director of the facility and the individual’s primary care physician notified, and all specified follow-up procedures performed. In the event of mechanical or electrical failure that may compromise the accuracy of the test results or monitoring capabilities, the test should be terminated until the problem is corrected.

<table>
<thead>
<tr>
<th>General Indications for Stopping an Exercise Test in Apparently Healthy Adults</th>
</tr>
</thead>
<tbody>
<tr>
<td>Onset of angina or angina-like symptoms</td>
</tr>
<tr>
<td>Significant drop (20 mmHg) in systolic blood pressure or a failure of the systolic blood pressure to rise with an increase in exercise intensity</td>
</tr>
<tr>
<td>Excessive rise in blood pressure: systolic pressure &gt;260 mmHg or diastolic pressure &gt;115 mmHg</td>
</tr>
<tr>
<td>Signs of poor perfusion: tight-headedness, confusion, ataxia, pallor, cyanosis, nausea, or cold and clammy skin</td>
</tr>
<tr>
<td>Failure of heart rate to increase with increased exercise intensity</td>
</tr>
<tr>
<td>Noticeable change in heart rhythm</td>
</tr>
<tr>
<td>Subject requests to stop</td>
</tr>
<tr>
<td>Physical or verbal manifestations of severe fatigue</td>
</tr>
<tr>
<td>Failure of the testing equipment</td>
</tr>
<tr>
<td>Assuming that testing is non-diagnostic and is being performed without direct physician involvement or electrocardiographic monitoring.</td>
</tr>
</tbody>
</table>
Considerations for accuracy

The ability to obtain valid and reproducible results is essential to ensure that any differences between pre-treatment and post-treatment test results are due to exercise training rather than variations in testing procedures. Some inconsistencies that are inherent may increase variability:

- Sub-maximal heart rate is influenced by time of day, eating, smoking, and familiarization with test procedures.
- Prediction equations for estimating VO$_{2\text{max}}$ may overestimate trained individuals and underestimate untrained individuals.
- The efficiency of motion during walking, running, and cycling varies.
- Cardiac output and VO$_2$ have a test-retest variability of 3-4%.

Psychological factors, such as pre-test anxiety, may influence the heart rate, especially at rates below 120 beats per minute and at low workloads. It is not unusual for the heart rate and/or blood pressure to be higher at rest than during the initial stages of exercise in these cases. Having the subject repeat the first test may improve reliability, particularly if the subject has never previously performed such a test.

Factors that can cause variation in the heart rate response to testing:

- Dehydration
- Prolonged heavy exercise prior to testing
- Environmental conditions (e.g., heat, humidity, ventilation)
- Fever
- Use of alcohol, tobacco, or caffeine 2 to 3 hours prior to testing

Because of these inherent inconsistencies, standard procedures for each test must be strictly followed to ensure the greatest accuracy and reproducibility possible:

- Standard testing protocol
- The same testing modality and protocol for repeat testing
- A constant pedal speed throughout cycle ergometry testing
- Cycle seat height properly adjusted, recorded, and standard for each test
- The time of day for repeat testing consistent
- All data collection procedures standardized and consistent
- Test conditions standard
- Subjects free of infection and in normal sinus rhythm
- Prior to the test, no intense or prolonged exercise for 24 hours, smoking for 2-3 hours, caffeine for 3 hours, or heavy meal for 3 hours
- Room temperature 18-20°C (64-68°F) with air movement provided
Performing the test

In this chapter it is supposed that the user is able to:

- perform an exercise test
- create exercise protocols
- view, edit and print tests

If this is not the case, please read the Exercise testing chapter.

To perform a sub-maximal test, follow these instructions:

1. Create a proper protocol (procedural guidelines for several sub-maximal testing protocols are provided in [ACSM’s Guidelines for Exercise Testing and Prescription, 6th Edition Philadelphia: Williams&Wilkins, 2000:22-29]).

2. Start an exercise test.

3. Perform the test as it were a maximal exercise test, ending it when the heart rate reaches the 85% of the Hrmax, or it happens an event listed in the section Test termination.

4. Display a VO2/Kg vs. HR plot

5. Right-click on the graph and select VO2 submax from the pop-up menu.

An example of testing protocol

An example of protocol is reported here. The YMCA cycle ergometry protocol is defined as follows.

1st step: workload 150 kgm/min

2nd step: if the HR at the end of the 1st step is:  
- <80  750
- 80-89  600
- 90-100  450
- >100  300

3rd step: if the HR at the end of the 2nd step is:  
- <80  900
- 80-89  750
- 90-100  600
- >100  450

4th step: if the HR at the end of the 3rd step is:  
- <80  1050
- 80-89  900
- 90-100  750
- >100  600

If the predicted HR max (calculated as 220-age) is not suitable for the patient tested, it is possible to edit the HR max value from the View/Information page.
Spirometry
Setting spirometry options

The software allows to configure some options selecting Configure from the Option menu.

Spirometry

Automatic Interpretation

K4 $b^2$ has the function of interpreting each test performed by a patient visualising an automatic diagnosis. The algorithm has been calculated basing on “Lung Function Testing: selection of reference values and interpretative strategies, A.R.R.D. 144/1991:1202-1218”.

The automatic diagnosis is calculated at the end of the FVC Test if:

- the automatic diagnosis option is enabled.
- the patient’s anthropometric data allow the calculation of the LLN (Lower Limit of Normal range).
- at least one FVC test has been performed.

To enable/disable the automatic diagnosis:

1. Click on Enable Automatic Interpretation checkbox to enable or disable the calculation and the visualisation of the automatic interpretation.
2. Select the LLN (Lower Limit of Normal Range) criteria among the ATS (LLN=Pred-0.674*SD), ERS (LLN=Pred-1.647*SD) or 80%Pred (LLN=Pred*0.8) specifications.

Quality control

K4 $b^2$ allows a quality test control. The calculation has been carried out referring to “Spirometry in the Lung Health Study: Methods and Quality Control, A.R.R.D. 1991; 143:1215-1223”. The messages concerning the quality control are shown at the end of the test.

To enable/disable the quality control, click on Enable Quality Control checkbox.
Parameters manager

The program allows to calculate a huge number of parameters; it is advisable, in order to simplify the analysis of the results, to view, to print and to sort the desired parameters only. Select the menu item Options/Parameters...

View
Move the parameters to view into the Selected parameters list.

Print
Move the parameters to print into the Selected parameters list.

Sort
Drag the parameter up or down with the mouse.

Customise
Add, modify and delete custom parameters.

If it is necessary to restore the default parameters press the button in the left corner of the window to initialise the parameters database.

Predicted values manager

The program contains a preset of predicted equations, but the user is allowed to customise its own predicted sets. Select Predicteds... from Options menu.

The window is divided into two forms: Predicteds set and Formula definition.

Predicteds set
This form allows the user to manage the set of predicteds. The following information define a set:
Name: identifies the set and cannot be duplicated;
Description: free field;
Age: the adult predicted age starts since this age.

To enter a new set of predicted values click on the **New** button. The field Name must be filled and must be unique. To stop without saving click on the **Cancel** button. To save the set, click on the **Save** button.

To delete a set of predicted values click on the **Delete** button. If a set is deleted, also the associated formulae are deleted.

It is possible to generate a new set of predicted values with the same attributes and the same formulae of the selected one. To do this click on the **Copy** button and specify a new Name.

To import a set of predicted values click on the **Import** button and select a file of Predicted files type.

To export a set of predicted values click on the **Export** button.

In the list **Set current predicted** choose the current predicted values for printing and viewing.

**Set the current predicted**

K4 $b^2$ allows to calculate the predicted values according to the following configurable sets:

<table>
<thead>
<tr>
<th>Adult</th>
<th>Paediatric</th>
</tr>
</thead>
<tbody>
<tr>
<td>ERS 93</td>
<td>Zapletal</td>
</tr>
<tr>
<td>Knudson83</td>
<td>Knudson83</td>
</tr>
<tr>
<td>ITS white</td>
<td>ITS white</td>
</tr>
<tr>
<td>ITS black</td>
<td>ITS black</td>
</tr>
<tr>
<td>LAM</td>
<td>LAM</td>
</tr>
<tr>
<td>MC Barcellona</td>
<td>MC Barcellona</td>
</tr>
<tr>
<td>Nhanes III</td>
<td>Nhanes III</td>
</tr>
</tbody>
</table>

Select the desired choice in the group **Predicted**.

**Formula definition**

This form allows the user to manage the formulae associated to a set of predicted values.

Select the set of predicted values from the list **Predicted** set.

To insert a new parameter click on the **New** button.

The parameter formulae can be:
calculated according to the predicteds in the list Use the predicteds formulae;
customised by the user with the option ...or the customised formulae.
The Delete button deletes the selected parameter.
The Copy button stores the selected parameter in memory.
The Paste button inserts a new parameter from the one copied. If the name is not
unique, the user is asked whether to specify a new name or to replace the existing
parameter.

Page set-up

Select Page Setup... from the File menu.

Header
All the printouts carried out by the program are preceded by 3 rows of customisable header (usually they contain the name
and the address of the Hospital using the spirometer).

Data
Patient and visit information are printed below the header. These data are reported on 3 columns and 5 rows. the user may
configure the disposition, change and eventually cancel the fields, as he prefers.

Margins
Configures the print margins from the borders of the paper.
The unit of measure is decided in Units of measurements.

Footer
Configures information at the bottom of the page.

Printed file name
Defines the automatic name to be assigned to the pdf file, if
the report will be printed in this format.

In the example it has been set to create a filename composed by
(date of the test) followed by <last name> and <first name>. 
Spirometry tests

Once completed the phases of the introduction of the patient’s data and the visit data, it is possible to carry out the spirometric tests.

K4 b² allows to perform the following tests:

<table>
<thead>
<tr>
<th>Key</th>
<th>Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>FVC pre</td>
<td>Forced Vital Capacity</td>
</tr>
<tr>
<td>FVC post</td>
<td>Forced Vital Capacity after bronchial stimulation</td>
</tr>
<tr>
<td>SVC</td>
<td>Slow Vital Capacity</td>
</tr>
<tr>
<td>MVV</td>
<td>Maximum Voluntary Ventilation</td>
</tr>
</tbody>
</table>

Before performing any test make sure that:

1. K4 b² is properly connected to your PC and the selected serial port (COM1, COM2) corresponds to the one effectively use.
2. The name shown on the status bar corresponds to the patient who is to carrying out the tests.
3. The today’s visit card exists.

Note: Read carefully the contraindications in Chapter 1.
Forced Vital Capacity (pre)

FVC is a reference test to verify obstructive (airflow limitations) and restrictive disorders (lung volume limitations). To achieve good test results it is fundamental a good manoeuvre (quality control messages, real time plots …)

The main parameters measured during FVC tests are:

- **FVC** Forced Vital Capacity
- **FEV1** Forced Expiratory Volume in 1 second
- **FEV1/FVC%** FEV1 as a percentage of FVC
- **PEF** Peak Expiratory Flow
- **FEF25-75%** Forced mid-Expiratory Flow

The two representative plots are the Flow/Volume and Volume/Time loops.

By comparing FVC, FEV1 and FEV1/FVC% values the software allows an automatic interpretation concerning the levels of obstructive and/or restrictive disorders.

**Recommendations**

- The flowmeter has to be disconnected from the breathing valve
- The patient should wear the nose clips
- The turbine has been recently calibrated (ATS recommends a daily calibration)
- The paper mouthpiece or the antibacterial filter is properly connected to the flowmeter through the corresponding adapter

For hygienic reasons, we strongly recommend the use of a bacterial filter.

If a kid must perform the test it is recommended to enable the encouragement function which shows exactly the manoeuvre of the FVC test.

**Perform a FVC (pre) test**

1. Select **Forced Vital Capacity pre** from the Test menu and wait for the green led is prompted on the right side of the screen.
2. Explain the manoeuvre to the patient and press the F2 key.
3. Wait some seconds and perform the test.
4. After having performed the test, press F3 or wait for the automatic end (5 seconds without flow), so that the software displays the F/V and V/t graphs, the main parameters, and the predicted values.
5. In order to visualise the F/V and V/t graph and the main parameters press the following buttons:
   - view Flow Volume graph
   - view Volume Time graph
   - view data of the test
6. Press Alt+F3 to stop the acquisition discarding the results.
7. Repeat the test until it is correctly performed (ATS recommends 3 times).
8. Press Exit to visualise the test list carried out during current session together with the results of the main parameters.
9. Select the test that you want to save (the Best Test according to the ATS criteria is highlighted as default) and press OK.

**Test encouragement**

During FVC manoeuvre you might experience some lack of collaboration with kids or with other patients. In this case you may find a good help in using the encouragement software tool.
Perform the FVC test with the encouragement

1. Select **Encouragement** from **View** menu.

2. Perform the test as explained in the previous paragraph.
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Slow Vital Capacity

Important test for assessing COPD (chronic obstructive pulmonary disease) patients affected by this disease might present a the Slow Vital Capacity could be higher than the Forced one (FVC).

The main parameters measured during SVC tests are:

- **EVC** - Expiratory Slow Vital Capacity
- **IVC** - Inspiratory Slow Vital Capacity
- **ERV** - Expiratory Reserve Volume
- **IRV** - Inspiratory Reserve Volume

If the inspiratory/expiratory maximal manoeuvre is preceded by a some breaths at tidal volume the software allows to measure the Respiratory Pattern, represented by the following parameters:

- **VE** - Ventilation per minute
- **Vt** - Tidal volume
- **Rf** - Respiratory frequency
- **Ttot** - Breath time
- **Ti/Ttot** - Inspiratory time/Ttot
- **Vt/Ti** - Vt/Ti

**Perform a SVC test**

1. Select **Slow Vital Capacity** from the **Test** menu and wait for the green led is prompted on the right side of the screen.
2. Press **F2** and instruct the Patient to breath normally until the message “carry out...” is prompted; then ask to perform a Slow Vital Capacity (deep inhalation, maximal slow expiration and deep inhalation again).
3. Press **F3** or wait for automatic interruption (5 seconds without flow) in order to visualise the V/t graph together with the main parameters compared to the predicted values.
4. To visualise the V/t graph and the main parameters press the follow buttons:
   - view Volume Time graph
   - view data of the test
5. Press **Alt+F3** to stop the acquisition discarding the results.
6. Repeat the test until it is correctly performed (ATS recommends 3 times).
7. Press **Exit** to visualise the test list carried out during current session together with the results of the main parameters.
8. Select the test that you want to save (the Best Test according to the ATS criteria is highlighted by default) and press **OK**.

The reference for the ERV calculation is displayed on the V/T graph.
Test for assessing the maximum ventilatory capacity. In the past, it was commonly performed during routine PF tests, however its clinical use declined over the years. Today MVV test is most commonly performed as part of the exercise tolerance tests, where it is used as an index of maximum ventilatory capacity. Test consists in breathing in and out deeply and rapidly for 12, 15 seconds. The expired volume during this short period is then extrapolated.

The most important measured parameter is the following:

**MVV** Maximum Voluntary Ventilation

**Perform a MVV test**

1. Select **Maximum Voluntary Ventilation** from the test menu and wait for the green led is prompted on the right side of the screen.
2. Press **F2** and make the Patient breath as much deeply and rapidly as possible for at least 12 seconds.
3. Press **F3** or wait for automatic interruption (5 seconds without flow) in order to visualise the V/t graph together with the main parameters compared to the predicted values.
4. To visualise the V/t graph and the main parameters press the follow buttons:
   - view Volume Time graph
   - view data of the test
5. Press **Alt+F3** to stop the acquisition discarding the results.
6. Repeat the test until it is correctly performed (ATS recommends 3 times).
7. Press **Exit** to visualise the test list carried out during current session together with the results of the main parameters.
8. Select the test that you want to save (the Best Test according to the ATS criteria is highlighted as default) and press **OK**.
Bronchial Provocation Test

Bronchodilator test

Bronchodilators are administered routinely in the b² laboratory to determine whether airflow obstruction is reversible. Bronchodilators increase airway calibre by relaxing airway smooth muscle.

The test consists of comparing results between the reference FVC (FVC PRE) and the FVC POST performed after the administration of the drug. Increasing value of 13-15% of FEV₁, respect to the basal value (FVC Pre) is considered as a reversible condition.

Main parameters are the following:

- **DFEV₁%pre**: Change of FEV₁ as a percentage of test PRE
- **DFVC%pre**: Change of FVC as a percentage of test PRE
- **DPEF%pre**: Change of PEF as a percentage of test PRE

Some authors states that the above mentioned parameters are too dependent from the FVC Pre, hence latest reference (ERS93, [A comparison of six different ways of expressing the bronchodilating response in asthma and COPD; reproducibility and dependence of pre bronchodilator FEV₁: E. Dompeling, C.P. van Schayck et Al; ERJ 1992, 5, 975-981]) recommend the following parameters:

- **DFEV₁%pred**: Change of FVC as a percentage of predicted value
- **DFEV₁%poss**: Change of FEV₁ as a percentage of “possible value”

Methacholine and Histamine Bronchial provocation Tests

The most common indication for performing methacholine and histamine bronchial challenges is to diagnose hyperresponsive airways. Some patients demonstrate normal baseline pulmonary function despite complaints of “tightness” wheezing, cough, and a little or not response to bronchoconstrictor. Other patients demonstrate spirometric improvement after use of bronchoconstrictor have diurnal variation in peak flows. In this groups aerosolised bronchial challenges are used to confirm a diagnosis of Asthma.

We can summarise the use of the test as follows:

1. Diagnose asthma
2. Confirm a diagnosis of asthma
3. Document the severity of hyperresponsivness
4. Follow changes in hyperresponsivness

When patients with hyperresponsive airways inhale certain pharmacologic agents (i.e. Methacholine or histamine) the airways respond by constricting.

Test consists of executing repeated FVC following the pharmacologic agents inhalation according to an established protocol. The fall of the FEV₁ parameter is used to calculate the bronchial hyperresponsivness. The most important parameter is the PD20 that is amount of drug (mg/ml) that causes a reduction of 20% of the FEV₁ respect the basal value (without drug).

Main parameters are:

- **P₁₀**: Dose that causes a 10% fall of FEV₁.
- **P₁₅**: Dose that causes a 15% fall of FEV₁.
- **P₂₀**: Dose that causes a 20% fall of FEV₁.

The representative plot is the Dose/response curve, showing the percentage variation of FEV₁ versus the Drug dose in logarithmic scale.

The program assumes as the baseline test the best FVC pre carried out during the today’s visit. You can change the reference pre test editing the Post test.

The name of the drug, its quantity and its unit of measurement, can be typed immediately before any FVC post manoeuvre (manual protocol) or can be stored in a database of bronchoprovocation (File/Bronchial Provocation protocols Database…).
Perform the test

(During 1st step only) select **Protocol...** from the **Test** menu and choose the name of the bronchoprovocation protocol that you are going to use (**manual protocol** if you want to type the information about the agent before any manoeuvre)

1. Select **FVC post** from the **Test** menu.
2. Select an existing protocol or click on “manual protocol”, and wait the green leds turned on.
3. Press **F2**, or the button by side, to start the test.
4. Press **F3**, or the button by side, to achieve the test.
5. In order to visualise the V/t graph and the main parameters press the follow buttons:
   - view Flow Volume graph
   - view data of the test
   - view bronchial provocation response
6. Press **Alt+F3** to stop the acquisition discarding the results.
7. Repeat the test until it is correctly performed (ATS recommends 3 times).
8. Press **Exit** to visualise the test list carried out during current session together with the results of the main parameters.
9. Select the test that you want to save (the Best Test according to the ATS criteria is highlighted as default) and press **OK**.

**Bronchial Provocation protocols Database**

The response to a bronchoprovocator is usually assessed in terms of change in the FEV1, vital capacity or airways resistance on the basis of serial measurements (FVC manoeuvres) in which the results of the initial test constitute the reference values. The international literature proposes several standardised protocols in order to address the methodological issues of the various available techniques.

The possibility to store a bronchoprovocation protocol in a database is useful to simplify and automate the sequence of operations that the Physician need to execute during the bronchoprovocation tests.

The typical sequence of activities to carry out a bronchoprovocation test are:
1. Typing and storing a bronchoprovocation protocol in the database (usually only once).
2. Selection of protocol among the list of the ones already present in the database before carrying out the FVC post tests (the selection of “manual protocol” allows to execute the test fully manually).
3. Performing the Post tests.

**Enter a new Bronchial provocation protocol in the archive**

1. Select **Bronchoprov. protocols database** from the **File** menu.
2. Type the Protocol name, the Bronchoprovocator name and the unit of measurement in the proper input fields.
3. If the bronchoprovocator has a cumulative effect select the cumulative check button.
4. Enter the quantities for each step and press the button **.**
Viewing results

All the visualisation functions refer to the test carried out by the Current Patient, whose name is indicated on the left-side of the status bar.

To view tests results:
1. Select the Patients from the File menu
2. Select the patient corresponding to the test you want to view.
3. Select in the list box of the tests up to 5 tests of the kind (FVC, VC/IVC, or MVV) and press OK.

To switch between graph and or data use the following buttons on the toolbar:

- view Flow Volume graph (F5)
- view Volume Time graph (F6)
- view data of the test (F7)
- view bronchial provocation response.

If you need more than one visualisation meantime use the New Window function from the Window menu.

If you need to display a list of visits:
- Select Visits list... from the File menu.
- Type the name of the Company and/or the time interval desired or simply confirm for the complete list.

Tests of the current patient

If a current patient has been selected you can quickly view his tests selecting Test current patient... from the View menu.

Delete a test
1. Select Patients from the File menu or press the button by side.
2. Select the test that you want to eliminate from the list of the tests referred to the Current Patient and press Delete.
Printing results

You can print out in three different ways:

- printing the Report
- printing the Active Window
- printing a series of reports

Printing Reports

To print a report of the current visit, select Print report... from File menu. The software will choose automatically the best performed test.

The standard Report is composed by 1, 2 or 3 pages depending if you wish to printout the FVC data and the graphs together on the first page or if you wish to printout the bronchoprovocation response.

- Selecting the option One page (no ATS) the report will contain, on one page, the F/V and V/t graphs of the best test, overlapped on the FVC Post, the patient data, the notes, the diagnosis and the test results.
- Otherwise the report will contain two pages, the first with the patient data, the graphs and the diagnosis, and the second one with the measured parameters, according to the ATS recommendations.
- The 3rd page will contain the bronchoprovocation response.

Select the desired options:
- FVC graph Prints the F/V and V/t curves for the best FVC test.
- One page (no ATS) Prints data and graphs on the first page.
- Response Prints the bronchoprovocator response.
- Preview Views a report preview on the screen.

Printing the active window

This printout function is only enabled when the active window (title bar highlighted) is one of the following objects:

- Any kind of Graph.
- Numeric data
- List of visit

To print the active window

1. Ensure that the active window is one of the preceding objects.
2. Select Print Active window from File menu.

Printing a series of reports

Sometimes it is useful to printout automatically a series of reports (all tests carried out with the employees, all tests carried out in the today’s session).

To print out proceed as follows:

1. Select Visit List from the File menu
2. Set the criteria of the visits to be added in the list (from, to,...)
3. Select Print Report from the File menu.

Electronic reports (*.pdf)

If an Adobe PDF writer “Printer Driver” is installed and set as the default printer, it is possible to store the printout report automatically in any location of the HD or eventually LAN paths according to a customizable filename format.

It is possible to define the created filename format selecting File/Page Set up... (see Page set-up).
Export data

With this function you can export the test data in 4 different formats:

- *.txt (ASCII)
- *.xls (Microsoft Excel)
- *.wk1 (Lotus 123)
- *.xpo (Cosmed)

Export a test

1. Select **Export tests** from the **File** menu.
2. Select the test to export from the list box and press **OK**.
3. Type the name and the format of the file in the dialog **Save as**. If the ASCII format is selected, the Text button in the dialog box Save as allows you to configure the separators for character based files.
   
   With the *.xpo Cosmed file format it is possible to import data from another K4 b® archive. Press **OK** to confirm.
4. Select the folder for the export and type the file name. Press **OK** to confirm. A status bar will show the file creation.
External devices
GPS

GPS initialisation

The GPS operates on information gathered from satellites. To gather this information, take your GPS outside and find a large, open area that has a clear view of the sky (a nearby park would work fine). The GPS needs to receive at least three strong satellite signals to find your location.

At the first power on the GPS needs to be initialized; the initialisation is a fundamental procedure for obtaining accurate and reliable data and should be performed on a large area where the sky is fully "visible".

After the initial self test is complete, the GPS will begin the process of satellite acquisition and tracking. The acquisition process is fully automatic and, under normal circumstances, will take approximately 45 seconds to achieve a position fix (15 seconds if ephemeris data is known).

Like all GPS receivers, COSMED GPS utilizes initial data such as last stored position, date and time as well as satellite orbital data to achieve maximum acquisition performance. If significant inaccuracy exists in the initial data, or if the orbital data is obsolete, it may take 5.0 minutes to achieve a navigation solution. The GPS Autolocate™ feature is capable of automatically determining a navigation solution without intervention from the user. This procedure may be required if one of the following situations occurs:

1) Transportation over distances further than 1500 kilometers.
2) Failure of the internal memory battery without system standby power.
3) Stored date/time off by more than 30 minutes.

The GPS will automatically update satellite orbital data as it operates. The intelligence of the GPS combined with its hardware capability allows these data to be collected and stored without intervention from the host system.

Initialize the GPS

1) If the receiver is not operated for a period of six (6) months or more, the unit will "search the sky" in order to collect satellite orbital information. This process is fully automatic and, under normal circumstances, will take 3-4 minutes to achieve a navigation solution.
2) If the memory backup battery of the GPS fails, the receiver will search the sky as described above. Should the memory battery discharge, the unit needs to be powered on for several hours to insure a sufficient recharge to maintain several months of clock operation and memory storage.
3) If the initial data is significantly inaccurate, the receiver perform an operation known as AutoLocate™. This procedure is fully automatic and, under normal circumstances, will require 1.5 minutes to calculate a navigation solution.

During the acquisition process a message "acquiring satellites..." is prompted on the display of the Portable unit.

The AutoLocate™ function can be manually forced selecting GPS AutoLocate from the Calibration menu, in order to obtain the best accuracy.

Fixing the antenna to the subject

The GPS Receiver has to be positioned onto the harness of the K4b² according the following pictures, paying attention to keep the receiver in a position so that the sky will be always "visible" during the test.

Some applications such as cycle racing and rowing may require different positioning of the antenna.
Operating sequence

Test with GPS module can be carried out with K4 b² system in Holter Data Record or Telemetry Data Transmission mode only. In addition to the Operating sequence of this mode you must carry out the following operation.

Run a test with GPS

1. Connect the receiver antenna to the Portable Unit plugging phone jack into the RS232 port at the bottom of the PU.

2. Select Settings then External device and press Enter.

3. Enable the GPS option by moving the “*” sign on GPS and press Enter to confirm settings.

4. Check the GPS module functionality choosing Calibration then GPS Control and press Enter. Display will show latitude and longitude.

5. With the use of Up and Down key verify that the displayed altitude value (Alt) is different from zero. In case displayed altitude value is fixed on zero, please be sure that the antenna receiver is well plugged in, the “sky” is visible and wait until the Altitude value is shown.
Monitoring GPS parameters in real time

To monitor in real time GPS parameters during Telemetry Mode Transmission or as soon as test has been stored or downloaded, go to the PC software and select Parameters to view/Test execution... (real time) or Parameters to view/Test visualization... (after download) from the Options menu.

Select the following parameters:

- **Velocity**  
  GPS Vel (m/sec)
- **Distance**  
  GPS Dist (meters, incremented during exercise phases only)
- **Latitude**  
  Lat (DD°MM.MMM’ N/S)
- **Longitude**  
  Long (DD° MM.MMM’ E/W)
- **Altitude**  
  Alt (meters)

Only when test has been stored or downloaded you can verify the **Graphical path** (automatically drawn on a scaled X/Y plane oriented to North) selecting on the PC software **Visualization** and **GPS track**.

**Note:** Distance is automatically calculated only during the "exercise" phases.
Pulse Oximeter (option)

The oximeter option is useful to monitor \( \text{SpO}_2 \) value during the test. Test with this option can be carried out with K4b\(^2\) system in Holter Data Record or Telemetry Data Transmission mode only.

In addition to the Operating sequence of this mode you must carry out the following operation:

**Operating Sequence**

1. Connect the Oximeter module to the Portable Unit plugging phone jack into the RS232 port at the bottom of the PU
2. Go to the K4b\(^2\) control panel and select Oximeter like External device connected by choosing **Settings** than **External device** and press **Enter**
3. Enable the Oximeter option by moving the “*” sign on **Oximeter** and press **Enter** to confirm settings.
4. Positioning Finger or Ear Clip on the patient and fix well the cable with Velcro stripes on the harness to minimize motion artifact

To monitoring in real time \( \text{SpO}_2 \) value during Telemetry Mode Transmission or as soon as test has been stored or downloaded, go to the PC software, select **Parameters to view/Test execution**... (real time) or **Parameters to view/Test visualization**... (after download) from the **Options** menu and select \( \text{SpO}_2 \) parameter.
System maintenance
System maintenance

All service operations which are not specified in this user manual should be performed by qualified personnel in accordance with the service handbook (to be required to the manufacturer).

Rubber mouthpieces, face masks, breathing valve and the other parts are not shipped sterile. They should be disinfected before using according to the following instructions.

All materials used in the construction of the K4 b^2 are non toxic and pose no safety risk to the patient or operator.

Prior to the device cleaning, disinfection and inspection it is necessary to switch off the device and to disconnect adapters from the supply mains.

In order to guarantee the highest accuracy of measurements we recommend you to disinfect the turbine periodically.

Use disposable anti-bacterial filters or disinfect each part in contact with the patient before each test.

Cleaning and disinfection

Cleaning and disinfecting instructions are of fundamental importance to control infections and assure patient safety. In fact aspiration of residue, particles and contaminated agents are life-threatening.

In this handbook we strongly recommend you to follow the rules worked out by ATS and ERS (see: "Lung Volume Equipment and Infection Control" – ERS/ATS WORKSHOP REPORT SERIES, European Respiratory Journal 1997; 10: 1928 – 1932), which are summarised as follows:

- Accessible internal as well as external surfaces of equipment exposed to expired gas should be washed and disinfected prior to testing of subsequent patients.
- Liquid disinfection can be used if the equipment is well cleaned first (no droplets of saliva/sputum remain).
- Disposable gloves should be worn when handling mouthpieces, when cleaning equipment exposed to saliva or sputum and especially when drawing blood.
- Laboratory staff should wash hands prior to testing of each patient.
- Adopt particular precautions when testing patients with recognised high-risk communicable diseases (e.g. tuberculosis, multidrug-resistant staphylococcus). In these cases, the clinical need for such testing should justify the risks.

During the disinfection:
- do not use alcohol or other liquids containing gluteraldehyde on the exterior surfaces of the equipment. Actually they can damage polycarbonates plastics and may produce unhealthy substances.
- do not use abrasive powders or glass cleaners containing alcohol or ammonia on the plexiglas components of the equipment
- do not steam autoclave any parts of the equipment unless it is clearly specified.
- do not immerse the optoelectronic reader.

Preparing the disinfecting solution

The following recommendations are retrieved from:


As disinfecting solution it is suggested:

- Sodium hypochlorite 0.5% (5000 ppm) prepared fresh for use within 24 hours.
- Sodium hypochlorite 1% (10000 ppm) prepared fresh for use within 30 days.

The first solution can be easily prepared by adding 1 part household bleach (sodium hypochlorite 5.25%) to 9 parts water, the second one by adding 1 part household bleach to 4 parts water.
Cleaning the turbine flowmeter

It is necessary to disinfect periodically the turbine for sanitary measures or/and for the correct device function.

The disinfecting procedure is easy and may be effected every time the user needs, keeping attention to some precautions:

1. Take out the turbine.
2. Dip it in a disinfectant solution (non alcoholic based) for about 20 minutes.
3. Rinse the turbine in a vessel, filled of clean water, shaking gently to remove the disinfectant (do not clean the turbine by putting it under running water!).
4. Let it dry to air.
5. After cleaning the turbine, check if the turbine propeller rotates freely even with a low speed air flow.
6. Connect the turbine to the reader.

Precautions during the cleaning of the turbine

1. Do not expose the turbine to high heat and do not put it under running water.
2. Do not ever dip the optoelectronic reader in any kind of solution, the liquid infiltration would damage the internal circuit.
3. Do not use alcoholic solutions to clean the turbine.

Suggested disinfection solutions

Helipur H Plus   Braun Melsungen AG
Gigasept FF     Schulke & Mayr GmbH
Dismozon pur    Bode Chemie GmbH
TETA-S          Fresenius AG
CIDEKX          Johnson & Johnson

Masks cleaning and disinfection

The face masks should be cleaned and sterilised after each test.

Disassembling the different parts of the mask

1. Remove the valves from their place.
2. Remove the adapter for the optoelectronic reader.

Cleaning the mask

1. Clean the mask with hot water and a soap solution to remove the impurities.
2. Rinse the mask with energy in running hot water.

Warning: Do not use synthetic or petroleum-based products for the masks cleaning.
Disinfecting the mask

It’s possible disinfecting the mask following these procedures:

- **Standard autoclaving method**
  - Rapid cycles of autoclave lasting 10 minutes at 132°C (270°F)
  - Heavy cycles of autoclave lasting 30 minutes at 121°C (250°F)
  - Pre vacuum cycles of autoclave lasting 30 minutes at 121°C (250°F)

- **Hetilene oxide method (ETO)**
  - The hetilene oxide doesn’t deteriorate the face masks. Sterilisation by this method is not advised unless sufficient data is available regarding the time required for complete out-gassing of residual ETO. If you use this method, follow carefully the instruction provided by the maker of the sterilising product.

- **Pasteurisation**
  - The disinfecting with hot water is a sterilising method that may be used with the silicone masks.

Permapure maintenance

- Do not bend, squash or deform it.
- Do not keep it in open air, if not used, especially in crowded or smoky places.
- If saliva is entered in the tube, replace it immediately, because it lost its functions.
- Periodically grease the o-ring on the connector in order to simplify the flowmeter connection.
- Replace it every 100 test / 6 month.

Inspections

The equipment requires easy inspections to be carried out in order to assure a proper electrical and mechanical safety level in the years.

These inspections are highly recommended after a rough use of the equipment or after a period of storage in unfavourable environmental conditions.

Referring to the electrical safety, it is important to check the conditions of insulation materials of cables, plugs and any other visible part by means of simple inspection, when the equipment is switched off and adapters (or electrical feeders) are disconnected from the supply mains.

Mechanical parts to check are: the turbine and breathing circuits.

Follow these instructions:

- extract the turbine from the optoelectronic reader;
- verify, by inspection, that the turbine axis fits correctly its seats and the blade is strongly fastened on the axis itself (it can be useful to shake slightly the turbine in order to note any anomalous movement).

Check if there are any torn or broken components in the breathing circuits: remember that they can create safety risk to patients during tests.

Replace the fuses

The fuses can be replaced easily in the following way:

1. Open the power supply cover using a screwdriver as shown in the picture.

2. Extract the fuse holder as shown in the picture
3. Replace the damaged fuse(s).

**Note:** Be careful to use proper fuses: 
A 680 023 500 (Time lag fuses 5x20 250V T500mA)
Warranty and limitation of liability

COSMED provides a one (1) year limited warranty from the date of the original sale of COSMED products. All COSMED products are guaranteed to be free from defect upon shipment. COSMED’s liability for products covered by this warranty is limited exclusively to replacement, repair, or issuance of a credit for the cost of a defective product, at the sole discretion of COSMED. COSMED shall not be liable under the foregoing warranty unless (i) COSMED is promptly notified in writing by Buyer upon discovery of defect; (ii) the defective product is returned to COSMED, transportation charges prepaid by Buyer, (iii) the defective product is received by COSMED no later than four weeks after the last day of the one (1) year limited warranty period; and (iv) COSMED’s examination of the defective product establishes, to COSMED’s exclusive satisfaction, that such defect was not caused by misuse, neglect, improper installation, unauthorised repair or alteration, or accident. If the product is manufactured by a third-party, COSMED shall make available for the Buyer’s benefit only those warranties which COSMED has received from the third-party manufacturer(s). COSMED hereby specifically disclaims any and all warranties and/or liabilities arising from defect(s) and/or damage(s) to and/or caused by products manufactured by third-party manufacturers. Buyer must obtain written authorisation from COSMED prior to the repair or alteration of COSMED products(s). Failure of Buyer to obtain such written authorisation shall void this warranty.

COSMED hereby specifically disclaims any and all other warranties of any kind, whether express or implied, in fact or by law, including, but without limitation, any and all warranties of merchantability and/or fitness for a particular purpose.

COSMED shall not be liable for special, indirect and/or consequential damages, nor for damages of any kind arising from the use of any COSMED’s products, whether said products are used alone or in combination with other products or substances. Determination of the suitability of any of COSMED’s product(s) furnished hereunder for the use contemplated by Buyer is the sole risk and responsibility of Buyer, and COSMED has no responsibility in connection therewith. Buyer assumes all risks and liabilities for loss, damage or injury to persons or property of Buyer or others arising out of the use or possession of COSMED’s products.

The limited warranty as herein above set forth shall not be enlarged, diminished, modified or affected by, and no obligation or liability shall arise or grow out of, the renderings of technical advice or service by COSMED, its agents or employees in connection with Buyer’s order or use of the product(s) furnished hereunder.

Return goods policy for warranty or non warranty repair

Goods shipped to COSMED for repair are subject to the following conditions:

1. Goods may only be returned after your receipt of a Service Return Number (SRN) from COSMED S.r.l.
2. Place your SRN report and Packing List outside the package.
3. Goods returned must be shipped with freight and insurance charges prepaid. Collect shipments will not be accepted.
4. The following list of goods are not eligible for return unless proven defective.
   - Special order items
   - Expendable products
   - Goods held over 30 days from COSMED’s invoice date.
   - Used goods not in original shipping containers.
   - Goods which have been altered or abused in any way.
5. The following parts are not covered by warranty:
   - consumables
   - fragile glass or plastic parts
   - rechargeable batteries
   - damages at the
   - damages due to use of the device not conforming to the indication reported in this manual
Repair Service Policy

Goods returned to seller for Non-Warranty repair will be subject to conditions 1, 2, 3, 4. The returned goods need to re-enter COSMED together with the customs documents (Pro-forma Invoice and Customs Paper) as requested by the Italian law.

- The shipment has to be qualified as a Temporary Export.
- All the goods returned to COSMED without the customs papers will not be accepted.

For European Community members:
Pro-Forma invoice complete with:

- Number
- Description of the goods
- Quantity
- Serial Number
- Value in €
- Number of parcel
- Gross weight
- Net weight
- Reason for resent (i.e. Resent for repair)

In case you should send the system for repair please contact the nearest service centre or contact COSMED at the following address:

**COSMED S.r.l.**
Via dei Piani di Monte Savello 37
P.O. Box 3
00040 Pavona di Albano - Rome, Italy
tel. +39 (06) 9315492
fax +39 (06) 9314580
E-mail: customersupport@cosmed.it

For USA customers only please contact:

**COSMED USA Inc**
2758 North Paulina
Chicago IL 60614 USA
Phone: +1 (773) 528-8113
Fax: +1 (773) 528-8116
e-mail: usa.sales@cosmed.it

To ensure that you receive efficient technical assistance, please specify as precisely as possible the nature of the problem as it is specified on the assistance information form. We advise you to save the original packaging. You may need it in case to ship the unit to a technical assistance centre.
Dear Customer,

we inform you that your personal data are gathered and will be used by Cosmed Srl in conformity with the requirements of the Italian privacy law (Decreto Legislativo 196/2003). We believe it is important for you to know how we treat your personal data.

**Personal data treatment and purposes**

We request and process your personal data:

a. to place an order, register a product, request a service, answer a survey, enter a contest, correspond with us (all of the above, in the following: “service”) and, if necessary, to supply the Competent Authorities with the required information;
b. in order to define your commercial profile;
c. in order to use your commercial profile for own marketing and advertising purposes;
d. for accounting purposes, including e-mailing of commercial invoices;
e. for providing your information to selected business partners (also abroad), in order to supply the service;

**How your personal data are treated**

Your personal data will be stored in electronic format, and protected at the best from destruction, loss (even accidental), not authorized accesses, not allowed treatment or use not in conformity with the purposes above listed.

**The consent is optional, but...**

If you deny the consent, we regret we cannot supply the service.

**Holder of the treatment**

The holder of the treatment is Cosmed Srl, Via dei Piani di Monte Savello 37, Pavona di Albano Laziale (RM). The responsible of the personal data treatment is indicated in the documentation stored by Cosmed Srl itself.

**Customer rights**

In accordance with art.7 of the Law, you can:

a. obtain confirmation of the existence of your personal data and their communication in intelligible form;
b. obtain:
   • updating, correction or integration of your data;
   • deletion or transformation in anonymous form of your personal data;
c. deny your consent to the treatment of your personal data;

These rights can be exercised directly requesting in writing to the holder of the treatment.
Converting factors configuration

You can edit the parameters shown in Control Panel by selecting *Control Panel* from the *Calibration* menu in the calibration program, then pressing the button by side.

You might configure the following options:

- **Name:** identify the parameter
- **Unit of meas.:** unit of measurement
- **Base line and Gain:** factors used to convert the acquired raw data (mV) into the final format according to \( Y = (mV - BL) \times Gain \). The value entered for gain must be multiplied by 1000 (for Gain=1, enter 1000).
- **Precision:** the number of decimals shown as 0
Calculations references

**VO₂ and VCO₂**


"Nutritional Assessment in Critical Care, A Training Handbook": Donald C. Zavala

**Anaerobic threshold (modified V-Slope)**

The break-point or intercept of the two slopes can be selected by a computer program that defines the VO₂ above which VCO₂ increases faster than VO₂, without hyperventilation.

During an incremental exercise above the Lactate Threshold, the net increase in lactic acid production results in an acceleration of the rate of increase in VCO₂ relative to VO₂. When these variables are plotted against each other (squared graph without recovery points), the relationship is composed of two apparently linear components, the lower of which has a slope of slightly less than 1.0, whereas the upper component has a slope steeper than 1.0. The intercept of these two slopes is the LT or AT point measured by gas exchange.

The increase in VCO₂ in excess of that derived from aerobic metabolism must be generated from the buffering of lactic acid. This is an obligatory gas exchange phenomenon seen in all subjects who exercise to work levels above their LT. This technique is referred to as the V-Slope method.


**O₂ kinetics**


In the following picture it is shown how the O₂ debit and deficit values are computed.
ATS recommendations


**ATS recommendations**

- **Volume range:** 8 l (BTPS)
- **Flow range:** ±14 l/sec
- **Volume accuracy:** ±3% or < 50 ml
- **Flow accuracy:** ±5% or < 200 ml/sec
- **Flowmeter resistance:** <1.5 cmH2O da 0 a 14 l/sec

**Reproducibility:** the 2 largest of 3 acceptable FEV1 and FVC values should be within 5% or 150 ml.

**The end of test:** no change in volume for 1 second with at least 6 seconds of collected volume.

**Accumulation time:** the maximum time allowed for volume accumulation during the VC manoeuvre should be at least 30 seconds and at least 15 seconds during the FVC.

The spirometer should be store at least 8 FVC manoeuvres.

FEV1 should be calculated by using the “back extrapolation” method to detect the start of the test, extrapolated volume must not be higher then 5% FVC or 150 ml.

The graphic resolution of the printed report must be as in the following:

- **Volume:** 10 mm/l
- **Flow:** 5 mm/l/sec
- **Time:** 20 mm/sec
- **F/V ratio:** 2:1

The total number of error (FVC e FEV1 >±3.5%, FEF25-75% >5.5%) during the measurement of the 24 standard waveforms must be lower than 4.
Predicted values

ERS93


KNUDSON 83

ITS

LAM

Multiéncrino de Barcelona

Nhanes III

Pneumobil (Brazil)
Valores extraídos do Programa Pneumobil/Brasil para a Tese de Doutoramento do Dr. Carlos Alberto de Castro Pereira. (Boehringer).

Gutierrez (Chile)
Gutierrez et Al. Reference values for Chile population

Knudson, Morris and Bass


Pereira (Brazil)
Pereira CAC; Barreto SP; Simões JG; Pereira FWL; Gerstler JG; Nakatani J. Valores de Referência para Espirometria em uma amostra da população brasileira adulta. Jornal de Pneumologia 1992; 18: 10-22.

Mallozi MC. Valores de referência para espirometria em crianças e adolescentes, calculados a partir de uma amostra da cidade de São Paulo. Valores finais publicados em : Pereira CAC; Lemle A; Algraniti E; Jansen JM; Valença LM; Nery LE; Mallozi M; Gerbasi M; Dias RM; Zim W. I Consenso Brasileiro sobre Espirometria. Jornal de Pneumologia 1996; 22:105-164.


Neder JA; Andreoni S; Castelo-Filho A; Nery LE. Reference values for lung function tests. I. Static Volumes. Brazilian Journal Medical and Biological Research 1999; 32:703-17.


DLCO


**Single Breath Oxygen Test**


**Rint**


**Mip/Mep**

Leo F. Black, Robert E. Hyatt: Maximal Respiratory Pressures: Normal Values and Relationship to Age and Sex, American Review of Respiratory Disease, Volume 99, 1969


### Automatic diagnosis (algorithm)


- LLN=Pred-0.674*SD (ATS, 50° percentile)
- LLN=Pred-1.647*SD (ERS, 95° percentile)
- LLN=Pred*0.8 (80% Pred)

<table>
<thead>
<tr>
<th>Message interpretation</th>
<th>Criterion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal Spirometry</td>
<td>FVC and FEV1/FVC &gt; LLN</td>
</tr>
<tr>
<td>Obstructive abnormality (it may be physiological)</td>
<td>% Pred FEV1 &gt;= 100</td>
</tr>
<tr>
<td>Obstructive abnormality: mild</td>
<td>% Pred FEV1 &lt; 100 and &gt;= 70</td>
</tr>
<tr>
<td>Obstructive abnormality: moderate</td>
<td>% Pred FEV1 &lt; 70 and &gt;= 60</td>
</tr>
<tr>
<td>Obstructive abnormality: moderately severe</td>
<td>% Pred FEV1 &lt; 60 and &gt;= 50</td>
</tr>
<tr>
<td>Obstructive abnormality: severe</td>
<td>% Pred FEV1 &lt; 50 and &gt;= 34</td>
</tr>
<tr>
<td>Obstructive abnormality: very severe</td>
<td>% Pred FEV1 &lt; 34</td>
</tr>
<tr>
<td>Restrictive abnormality: mild</td>
<td>FVC &lt; LLN and % Pred FVC &gt;= 70</td>
</tr>
<tr>
<td>Restrictive abnormality: moderate</td>
<td>% Pred FVC &lt; 70 and &gt;= 60</td>
</tr>
<tr>
<td>Restrictive abnormality: moderately severe</td>
<td>% Pred FVC &lt; 60 and &gt;= 50</td>
</tr>
<tr>
<td>Restrictive abnormality: severe</td>
<td>% Pred FVC &lt; 50 and &gt;= 34</td>
</tr>
<tr>
<td>Restrictive abnormality: very severe</td>
<td>% Pred FVC &lt; 34</td>
</tr>
</tbody>
</table>

### Quality Control Messages

Reference: Spirometry in the Lung Health Study: Methods and Quality Control, ARRD 1991; 143:1215-1223.

<table>
<thead>
<tr>
<th>Message</th>
<th>Criterion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start faster</td>
<td>VEXT &gt;5% of the FVC and &gt;150ml</td>
</tr>
<tr>
<td>Blast out harder</td>
<td>PEFT &gt;120 msec</td>
</tr>
<tr>
<td>Avoid coughing</td>
<td>50% drop in the flow in first second</td>
</tr>
<tr>
<td>Blow out longer</td>
<td>FET100% &lt;6 sec.</td>
</tr>
<tr>
<td>Task</td>
<td>Criteria</td>
</tr>
<tr>
<td>-------------------------------------</td>
<td>-----------------------------------------------</td>
</tr>
<tr>
<td>Blow out more air</td>
<td>flow &gt;0.2 l/s within 20 ml of FVC</td>
</tr>
<tr>
<td>Blow out harder</td>
<td>dPEF &lt; 10%</td>
</tr>
<tr>
<td>Take a deeper breath</td>
<td>dFVC &lt; 200 ml and 5% best FVC</td>
</tr>
<tr>
<td>Blow out faster</td>
<td>dFEV1 &lt; 200 ml and 5% FEV1</td>
</tr>
<tr>
<td>That was a good test</td>
<td>No errors</td>
</tr>
<tr>
<td>FVC reproducible</td>
<td>diff. 2 max FVC within 0.2 l</td>
</tr>
<tr>
<td>FEV1 reproducible</td>
<td>diff. 2 max FEV1 within 0.2 l</td>
</tr>
<tr>
<td>PEF reproducible</td>
<td>diff. 2 max PEF within 10%</td>
</tr>
<tr>
<td>MVV time too short</td>
<td>MVV time less than 12 sec</td>
</tr>
</tbody>
</table>
References

Gas Exchange References
[“Measurement and analysis of gas exchange during exercise using a programmable calculator”; Sue, Hansen, Blais, Wasserman, JAP, 49(3), 1980:456-461]
[“Principles of exercise testing and interpretation, 2nd edition”; Wasserman et Al, 1994]

Indirect calorimetry
[“Nutritional Assessment in Critical Care, A Training Handbook”: Donald C. Zavala]

Spirometry
Lung function”, J.E. Cotes, Blackwell scientific publications

Sub-maximal testing