

Review Station

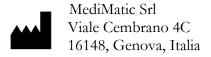
User's Guide

Rev. 10.12



Viale Cembrano 4C, 16148 Genova, ITALIA Tel: +39 010 3071634 Fax: +39 010 3074548 http://www.medimatic.com, info@medimatic.com







USER'S GUIDE¹

Basic UDI-DI 805647730COMPACSRH

 UDI-DI
 08056477300008

 Document Emission Date:
 March 21st, 2025

SW Revision: 10.12.x Manufacturer: MediMatic S.r.l.

Document Revision: 2

It is forbidden to copy this document – All rights reserved

© Copyright MediMatic S.r.l. 2004-2025

¹ All trademarks, names and logos in this document belong to their legitimate owners.

SUMMARY

User's Guide	
1. GENERAL INFORMATION	9
1.1. HOW TO USE THIS GUIDE	9
1.2. TEXT ICON	9
1.3. USING THE MOUSE	10
2. INTRODUCTION	
2.1. INDICATIONS FOR USE	12
2.2. INSTALLATION, DECOMISSIONING AND DISPOSAL	
LICENSE - COPYRIGHT - WARRANTY	16
3. LAUNCHING THE APPLICATION	24
3.1. START-UP PROCEDURE	
3.2. LOGIN	
3.3. SHUTDOWN PROCEDURE	
3.4. CHANGING PASSWORD	28
3.5. IDLE TIME PROTECTION	29
4. APPLICATION LAYOUT	30
5. QUICK START	
5.1. QUICK ACCESS TO MAIN FUNCTIONS	
6. REVIEW STATION	
6.1. OPENING FILES	
6.1.1 FILE FORMATS	36
6.1.2 OPENING A FILE FROM REMOVABLE DRIVES	36
6.1.3 OPENING A FILE FROM LOCAL HARD DRIVES 6.1.4 OPENING A PATIENT/STUDY FROM THE ARCHIVE	38
6.2. VIEWING IMAGES AND REPORTS	
6.2.1 THUMBNAILS	
6.2.2 IMAGE LAYOUT	42
6.2.3 FULL SCREEN	
6.2.4 ADDITIONAL INFORMATION	
6.2.5 INFORMATION OVERLAY	47
6.2.6 VIEWING REPORTS	
6.2.7 CLOSING ITEMS	
6.2.8 MULTI-MONITOR	51
6.3. PLAYING LOOPS	
6.3.1 HOW TO PLAY 6.3.2 SPEED CONTROL AND TRIMMING	52
6.3.2 SPEED CONTROL AND TRIMMING6.3.3 MULTI LOOP CONTROL	
6.3.4 STACKING/UNSTACKING	
6.4. FILTERING IMAGES	58
641 COLOR EUTER	58

6.4.2 DYNAMIC FILTER	59
6.5. TRANSFORMING IMAGES	_60
6.5.1 MIRROR AND, ROTATION	60
6.5.2 INTERPOLATED ZOOM	61
6.5.3 PAN	61
6.5.4 LENS	62
6.5.6 ROI SELECTION	62
6.5.7 MASKING	
6.6. STRESS ECHO	65
6.7. PRINTING OR EXPORTING TO MULTI-MEDIA FORMATS	67
6.7.1 LOOP EXPORT	
6.7.2 FRAME EXPORT	68
6.7.3 PRINTING IMAGES	
6.8. EXPORTING TO DICOM FOLDERS OR MEDIA	71
7. PROPERTIES SETTINGS	74
8. THE ARCHIVE	76
8.1. QUERY & RETRIEVE	76
8.1.1 NORMAL QUERY	76
8.1.2 QUERY RESULTS, COLORS AND INFORMATION ICONS	
8.1.3 ADVANCED QUERY	
8.1.4 RETRIEVAL AND DISPLAY 8.1.5 WEB COMPRESSION	80
8.2. ASSIGNING STUDIES	
8.3. ORDER MANAGEMENT	
8.4. PATIENT AND STUDY URGENCY STATUS	
8.5. ADDING OR EDITING NEW ITEMS	
8.5.2 ADDING OR EDITING FATIENTS	07
8.5.3 ADDING OR EDITING SERIES	
8.6. MERGING AND MOVING ITEMS	
8.6.1 MERGING ITEMS 8.6.2 MOVING ITEMS	91
8.6.2 MOVING ITEMS	
8.6.3 SPLITTING ITEMS	92
8.7. ARCHIVING PATIENT IMAGES AND LOOPS	93
8.7.1 EDITING INFORMATION	95
8.7.2 ANONYMIZING IMAGES	
8.7.3 TRIMMING IMAGES	96
8.8. ATTACHMENTS	98
8.9. MEMO	
8.10. PERMISSIONS	
9. KEY IMAGE NOTES	
9.1. REVIEWING A KEY IMAGE NOTE	
9.2. CREATING A NEW KEY IMAGE NOTE	107

9.3. EDITING A KEY IMAGE NOTE	107
9.4. OTHER OPERATIONS	109
9.5. OPENING A KEY IMAGE NOTE	110
9.6. SEARCHING FOR A KEY IMAGE NOTE	110
10. MANAGING EVIDENCE AND REPORTS	
10.1. CREATING A NEW REPORT	
10.2. IMPORTING EXISTING REPORTS	
10.3. EDITING A REPORT	
10.4. OPENING REPORT DOCUMENTS	
10.5. REPORT REVISION HISTORY	
11. ECHO ANALYSIS	
11.1. ACTIVATING OFF-LINE ECHO ANALYSIS	
11.2. OFF-LINE MEASUREMENTS	
11.3. IMPORTED MEASUREMENTS 11.3.1 IMAGE CALIBRATION	
11.3.2 PERFORMING MEASUREMENTS	126
11.3.3 CALCULATIONS AND FORMULAS	130
11.4. MEASUREMENTS MANAGEMENT	
11.5. CUSTOM SETTINGS	133
12. ANGIOGRAPHIC ANALYSIS	135
12.1. ACTIVATING QCA II	135
12.2. ACTIVATING LVA	135
12.3. CALIBRATION	136
12.3.1 DEFAULT CALIBRATION	137
12.3.2 CALIBRATION BASED ON A KNOWN DISTANCE	138
12.3.3 CATHETER BASED CALIBRATION	
12.4. CORONARY ANALYSIS	141
12.4.1 THE RESULTS WINDOW 12.4.2 ANALYSIS AND STENT	142
12.5. VENTRICLE ANALYSIS	
13. IVUS ANALYSIS	
13.1. ACTIVATING IVUS ANALYSIS	
13.2. LONGITUDINAL RECONSTRUCTION (ILD)	
13.3. OFF-LINE MEASUREMENTS	
13.3.1 IMAGE CALIBRATION	155
13.3.2 LONGITUDINAL RECONSTRUCTION CALIBRATION	157
13.3.3 PERFORMING MEASUREMENTS	158
13.3.3 PERFORMING MEASUREMENTS 13.3.4 CALCULATIONS AND FORMULAS	160
13.4. MEASUREMENTS MANAGEMENT	161

13.5. CUSTOM SETTINGS	163
14. GENERIC MEASUREMENTS	164
14.1. ACTIVATING OFF-LINE GENERIC MEASUREMENTS	164
14.2. OFF-LINE MEASUREMENTS 14.2.1 IMAGE CALIBRATION 14.2.2 PERFORMING MEASUREMENTS 14.2.3 CALCULATIONS AND FORMULAS	166 169
14.3. MEASUREMENTS MANAGEMENT	
14.4. CUSTOM SETTINGS	
15. PHILIPS QLAB	
15.1. ACTIVATING PHILIPS QLAB	
15.2. LAUNCHING PHILIPS QLAB	
16. INVIA CORRIDOR 4DM	
16.1. ACTIVATING INVIA CORRIDOR 4DM	
16.2. LAUNCHING INVIA CORRIDOR 4DM	
17. GE ECHOPAC PLUGIN	
17.1. ACTIVATING GE ECHOPAC PLUGIN	
17.2. LAUNCHING GE ECHOPAC PLUGIN	
18. TOMTEC ANALYSIS	
18.1. ACTIVATING TOMTEC ANALYSIS	182
18.2. LAUNCHING TOMTEC ANALYSIS	183
19. MEDIS ANALYSIS	184
19.1. ACTIVATING MEDIS ANALYSIS	184
19.2. LAUNCHING MEDIS ANALYSIS	185
20. EPSILON ECHO INSIGHT ANALYSIS	186
20.1. ACTIVATING EPSILON ANALYSIS	186
20.2. LAUNCHING ECHO INSIGHT ANALYSIS	187
21. CIRCLE CVI42 ANALYSIS	188
21.1. ACTIVATING CVI42 ANALYSIS	188
21.2. LAUNCHING CVI42 ANALYSIS	189
22. ECG MANAGEMENT	190
22.1. REVIEW OF DICOM ECG	190
22.2. ACTIVATING ECG ANALYSIS AND REPORTING	192
22.3. REPORTING	193
22.4. COMPARING	194
22.5. OFF-LINE MEASUREMENTS	195

22.5.1 ECG TRACE CALIBRATION	
22.5.2 PERFORMING MEASUREMENTS	197 198
22.6. MEASUREMENTS MANAGEMENT	
22.7. CUSTOM SETTINGS	
23. REPORTING	
23.1. ACTIVATING COMPACS REPORTING	
23.2. CREATING OR OPENING A REPORT	
23.3. PREVIEWING AND PRINTING THE REPORT	
23.4. ADVANCED OPERATIONS	
23.4.1 ADDING IMAGES	
23.4.2 ADDING ATTACHMENTS	210
23.4.3 ALL NORMAL STATEMENTS	
23.4.4 IMPORTING PREVIOUS REPORTS	210
23.4.5 RECOMPUTING VALUES	211
23.4.6 MEASUREMENT DETAILS	211
23.5. CONFIRMING OR DISCARDING CHANGES	
23.6. REVISION HISTORY	
24. CONTEXT SYNCHRONIZATION	
25. APPENDIX A	
25.1. POST MARKETING SURVEILLANCE	
25.2. Hazard Notification Form	
26. APPENDIX B	
26.1. ADDITIONAL ARCHIVES	
26.2. OTHER OPTIONS	
27. APPENDIX C	
27.1. ECHO ANALYSIS ACCURACY AND PRECISION	
28. APPENDIX D	224
28.1. ANGIOGRAPHIC ANALYSIS ACCURACY AND PRECISION	224
28.2. KNOWN DISTANCE BASED CALIBRATION	224
28.3. CATHETER BASED CALIBRATION	225
28.4. CORONARY ANALYSIS	225
28.5. VENTRICULAR VOLUME	225
28.6. BODY SURFACE	226
28.7. REGIONAL SHORTENING	226
29. APPENDIX E	227
29.1. IVUS ANALYSIS ACCURACY AND PRECISION	227

30. APPENDIX F	228
30.1. GENERIC MEASUREMENT'S ACCURACY AND PRECISION	228

1. GENERAL INFORMATION

1.1. HOW TO USE THIS GUIDE

Use this guide for both training and reference, regardless whether you are a new user, or a user who is already familiar with the system.

Symbols, called icons, show up throughout the guide to assist you in quickly finding cautions, suggestions, urgent information, reminders, and custom settings. You will find an explanation of these icons in this chapter.

1.2. TEXT ICON

Text written next to this caution sign indicates the most important information, safety precautions, or warnings.

The stop sign indicates information that you should stop and read before continuing.

This icon is a reminder to do something. It may be information you read about in a previous chapter. This icon may also indicate a suggestion or an idea, which may make it easier for you to use the software.

1.3. USING THE MOUSE

In this chapter are described the main functions of your mouse to quickly accomplish common functions and speed up your work. It teaches you mouse behavior, which is common to all WindowsTM OS.



If you are used to working with Windows you can jump to the next section.

• Left Mouse Button

This button is primarily used to select/deselect items on the desktop (for example an icon), to move items on the desktop and to open items or start applications.

To select an item on the desktop

- 1. Place the cursor over the item to be selected.
- 2. Click once and quickly release the left mouse button (called left-click). The item is now selected.

To deselect an item on the desktop

- 1. Position the cursor away from the selected item.
- 2. Click the left mouse button to deselect the item.

To move an item on the desktop

- 1. Point to the item.
- 2. Click and hold down the left mouse button and drag the item to another location. Release the left mouse button at this point and this will release the item in the new location.

To open an item on the desktop

- 1.) Double-click on the item with the left mouse button.
- 2.) If the item is connected to an application, e.g. ComPACS Report Manager, this application will be started. Other items like word processing files or spreadsheets will be opened with the tool they were created with.

• Right Mouse Button

This button is primarily used to activate a context specific task menu associated with each item on the desktop.

To complete common tasks on an item on the desktop

- 1. Right-click on any item on the screen.
- 2. A task menu appears, listing common tasks that you may want to carry out on this item. Click a task from this new menu to activate it. In this case, it is irrelevant which mouse button you click with.

• Middle Mouse Button

This button (if available) provides application specific functionality. Refer to the manuals of the applications you use for detailed information.

2. INTRODUCTION

The present user's guide refers to the ComPACS Review Station software and integrated modules and components.

2.1. INDICATIONS FOR USE

The device is a software product intended to be used by qualified medical professionals, after proper installation on an appropriate hardware and middleware platform, for retrieving, archiving, transmitting, viewing, analyzing and, reporting medical image and loops.

The software is designed to be installed by qualified IT professionals on an adequate and compatible hardware platform.



CAUTIONS AND WARNINGS

ComPACS can be used for diagnostic purposes only according to the European 93/42/EEC, 2007/47/EC Medical Device Directive and USA FDA 510k regulations. Diagnostic use under other regulations is expressly forbidden. **Restrictions under some regulations may also apply to specific product components.**

ComPACS and integrated modules and components may be subject to additional limitations. Please review the related certificates to determine which modules can be used for diagnostic purposes according to the above regulations.

ComPACS and integrated modules and components are under the license, terms and warranty in this document. Please review the license and warranty conditions carefully before using the device. Notice that the use of this device indicates your acceptance of this license agreement and warranty.

ComPACS and integrated modules and components system requirements are indicated in the following section. If the requirements are not met, the device may not function correctly.

The present document must be read carefully before using ComPACS and integrated modules and components. In particular it is important to note the following warnings and advices.

This software should not be used without prior training and a thorough understanding of the operating instructions.

No other software must be installed in the review station other than compliant software for the released components. Please contact ComPACS support if you want to install additional software. The manufacturer or its representative shall not be responsible for problems attributable to unauthorized changes, additions or deletions to the software or hardware device, or unauthorized installation of third-party software.

Application usage on system configurations different to those explicitly indicated is prohibited. The manufacturer or its representative shall not be responsible for any and all consequences, including incorrect application behavior, performance, data loss, data corruption, and/or incorrect computations.

Workstation or viewer deployment and/or usage based on virtual desktop or remote desktop technologies is not supported. The usage of such technologies may alter image quality displayed and lead to incorrect diagnosis.

The information contained in this guide is intended only for the operation of the device. It does not include information such as how to perform medical imaging procedures on the medical devices used to obtain the DICOM images, on any medical procedures or on other medical devices.

Do not remove or change disks from the disk drive during disk read and write operations. This may cause incomplete storage of data files, resulting in data loss.

Do not disconnect the system from the network during its use. This may cause incomplete storage of data files, resulting in data loss.

If system power is lost or the network disconnected during a patient study the content of the stored patient file may be incomplete.

Before saving or reviewing a patient study file, always double-check the patient name to be sure that it corresponds to the correct one.

When a patient study is deleted, the files are permanently lost. Be sure to copy the patient study to another storage media for archival before deleting it.

When lossy compression is selected some information will be lost. The higher the compression ratio, then greater will be the data loss.

System administrators are responsible to ensure that the patient data is backed up daily to prevent the risk of permanent data loss. Furthermore, a daily backup of the SQL Server database also needs to be scheduled. Please refer to the System Administrator Manual for more detailed information.

System administrators are responsible to ensure that the systems used are adequately protected against virus and malware. Please refer to the System Administrator Manual for more detailed information.

System administrators are responsible to ensure that the decommissioning and disposal instructions are carefully followed when the software is decommissioned in order to avoid the disclosure of confidential information.

2.2. INSTALLATION, DECOMISSIONING AND DISPOSAL

The software is designed to be installed by qualified IT professionals on an adequate and compatible hardware platform.

Please refer to the ComPACS Installation Manual for detailed installation, decommissioning and disposal instructions.

Please refer to the ComPACS System Administrator's Manual for detailed configuration instructions.

In order to display the Electronic Instructions For Use it is necessary to have Adobe® Acrobat Reader installed on your system.

2.2.1 SYSTEM REQUIREMENTS

ComPACS Workstation and ComPACS Web Workstation

- Intel/AMD Quad Core processor or higher
- 16 GB RAM or higher (32 GB recommended)
- 80 GB disk or above
- Gigabit NIC
- (minimum) Graphics Card with 256MB, Dual DP/HDMI, Dual 19" flat panel LCD color monitors Resolution: 1280*1024 @ 60Hz, Brightness: 250cd/m2, Dot Pitch: 0.27mm Response Time: 8 ms
- (recommended) Graphics Card with at least 1 GB dedicated memory, Dual DP/HDMI, Dual 23" flat panel LED color monitors Resolution: 1920*1080 @ 60Hz, Brightness: 300cd/ m2, Dot Pitch: 0.27mm, Response Time: 8 ms, Contrast: 1.000:1 (typical), 2.000.000:1 (dynamic/max)
- Windows 10 Pro / Enterprise, 11 Pro / Enterprise
- Web Browser: Edge version 79 or above, Chrome version 54 or above, Firefox version 49 or above, Opera version 41 or above.
- Keyboard, Mouse

Note. Deployment based on Citrix® Virtual Desktops is not formally supported though it has been known to work. See further warnings below.

ComPACS Web Viewer

- Intel/AMD Dual Core equivalent processor or above
- 8 GB RAM or higher (16 GB recommended)
- 5 GB disk or above
- 100Mbps or higher NIC
- (minimum) Graphics Card with 256MB and Single DP/HDMI, Single 19" flat panel LCD color monitors Resolution: 1024*768 @ 60Hz, Brightness: 250cd/m2, Response Time: 8 ms
- (recommended) Graphics Card with at least 1 GB dedicated memory, Single DP/HDMI, Single 23" flat panel LED color monitors Resolution: 1920*1080 @ 60Hz, Brightness: 300cd/ m2, Dot Pitch: 0.27mm, Response Time: 8 ms, Contrast: 1.000:1 (typical), 2.000.000:1 (dynamic/max)
- Windows 10 Pro / Enterprise, 11 Pro / Enterprise
- Web Browser: Edge version 79 or above, Chrome version 54 or above, Firefox version 49 or above, Opera version 41 or above.
- Keyboard, Mouse

Note. Deployment based on Citrix® Virtual Desktops is not formally supported though it has been known to work. See further warnings below.

Third Party Components

When ComPACS is integrated with third party components (for example Philips QLAB) some additional requirements may apply. Please refer to the third-party component documentation for additional requirements specifications.

LICENSE - COPYRIGHT - WARRANTY

ComPACS Review Station and integrated modules and components

MediMatic S.r.l., Copyright © 2004-2025. All rights reserved.

NOTICE TO USER: PLEASE READ THIS LICENSE AGREEMENT CAREFULLY. BY COPYING, INSTALLING OR USING ALL OR ANY PORTION OF THE SOFTWARE YOU ACCEPT ALL THE TERMS AND CONDITIONS OF THIS AGREEMENT, INCLUDING, IN PARTICULAR THE LIMITATIONS ON: USE, TRANSFERABILITY, WARRANTY AND LIABILITY.

YOU AGREE THAT THIS LICENSE AGREEMENT IS LIKE ANY WRITTEN NEGOTIATED AGREEMENT SIGNED BY YOU. THIS LICENSE AGREEMENT IS ENFORCEABLE AGAINST YOU AND ANY LEGAL ENTITY THAT OBTAINED THE SOFTWARE AND ON WHOSE BEHALF IT IS USED: FOR EXAMPLE, IF APPLICABLE, YOUR EMPLOYER.

IF YOU DO NOT AGREE TO THE TERMS OF THIS LICENSE AGREEMENT, DO NOT USE THE SOFTWARE. PLEASE CONTACT MEDIMATIC OR YOUR RESELLER FOR TERMS OF AND LIMITATIONS ON RETURNING THE SOFTWARE FOR A REFUND.

BY USING THE SOFTWARE YOU ACKNOWLEDGE THAT YOU HAVE READ THIS LICENSE AGREEMENT, UNDERSTAND IT, AND AGREE TO BE BOUND BY ITS TERMS.

LICENSE AGREEMENT

This License Agreement constitutes the entire, final, and exclusive Agreement and supersedes any and all other prior or contemporaneous oral representations or agreements between the parties relating to the subject matter hereof. Please note however that you may have another written agreement directly with MediMatic and undersigned by both parties that supplements or supersedes all or portions of this agreement.

No amendments or modifications of this Agreement may be made except in writing signed by both parties. This Agreement is severable, and if any term or provision hereof is determined to be invalid or unenforceable, such determination shall not in any way affect the validity or enforceability of the remaining provisions hereof and any such term or provision shall be reformed, if reasonably possible, only to the extent necessary to make it enforceable.

Software License

MediMatic Srl and its suppliers own all intellectual property in the Software. The Software is licensed, not sold. MediMatic Srl permits you to copy, download, install, use, or otherwise benefit from the functionality or intellectual property of the Software only in accordance with the terms of this agreement. Use of some third-party materials and services included in or accessed through the Software may be subject to other terms and conditions typically found in a separate license agreement, terms of use or "Read Me" file located within or near such materials and services.

As long as you comply with the terms of this agreement, MediMatic Srl grants you a non-exclusive, non-transferable license to use the Software in the manner and for the purposes described in the Documentation, as further set forth below.

You may install and use one copy of the Software on one (1) computer for each valid license in your possession. If the valid license in your possession grants you concurrent user access then you may install a copy of the Software on more computers belonging to you or your organization up to the limit indicated by such license. The total number of concurrent users at any time may not exceed the number indicated by your valid license.

You may make a one (1) or two (2) backup copies of the Software, provided your backup copies are not installed or used for other than archival purposes.

Intellectual Property Ownership

The Software and any authorized copies that you make are the intellectual property of and are owned by MediMatic S.r.l. and its suppliers of third-party components, if any. The structure, organization and code of the Software are the valuable trade secrets and confidential information of MediMatic and such suppliers.

MediMatic, and its third party component MediMatic S.r.l. if any, retain all right, title, and interest in and to the Software, the Documentation, and any and all services performed in connection therewith, and any other content, products, documentation, software, materials, or other technology related to the Software including, but not limited to, any and all modifications, enhancements, updates, and configurations, whether written, printed, electronic, or in source code or any other format, and any patent, copyright, trade secret, trademark, service mark, or other intellectual property rights in any of the foregoing (the "MediMatic Property").

MediMatic grants only those rights and licenses expressly provided for herein and it does not hereby transfer any title or ownership interest in the MediMatic Property, nor in any third-party intellectual property, nor in any modifications, enhancements, or derivative works thereof. You shall not interfere with or otherwise challenge MediMatic's rights in the MediMatic Property, nor remove or alter, nor allow to be removed or altered, any notice, symbol, or legend of any trademark, copyright notice, or other proprietary rights appearing in or on any of the MediMatic Property. You hereby assign to MediMatic all rights you may have in any and all suggestions, concepts, improvements, or other enhancements communicated by you to MediMatic that are incorporated into the Software.

You shall keep the Software and the Documentation free and clear from any claims, liens, or encumbrances. You shall not: (i) modify, translate, reverse engineer, decompile, disassemble, create derivative works of, or otherwise attempt to derive any source code of the Software; (ii) alter or copy, or permit a third party to alter or copy, any part of the Software, except for a reasonable number of back-up copies of the Software for safety or archival purposes only; (iii) use the Software to provide service bureau, time sharing, access through a public computer bulletin board or "shareware" distribution process, or other services to third parties; nor (iv) sublicense, distribute, sell, assign, transfer, lease, rent, disclose, demonstrate, or allow access to the Software or the Documentation to any person except your agents or employees, and then only to the extent necessary to operate the Software as licensed hereunder.

Retention of Title

Title to the software licenses not fully paid by you although delivered shall be retained by MediMatic until the receipt of full payment.

User Responsibilities

When using the Software, you shall be solely responsible for administering access privileges and maintaining the security of all user names, passwords, and other Confidential Information (as defined below) relating to your use of the Software. You shall promptly: a) report to MediMatic any errors or irregularities in connection with the Software and provide copies of any and all programs, reports, data files, and other materials requested by MediMatic; and b) provide support sufficient for MediMatic to attempt to duplicate such failure; and c) provide sufficient test time within which MediMatic shall use commercially reasonable efforts to correct such failure.

You understand that Licensed Material may include third party software or other intellectual property and agrees that you shall treat such third-party material as you do the Licensed Material. You shall assume full responsibility for all risk associated with your use of the Software, including, but not limited to, any financial obligation arising with respect to your patients or other third parties.

Restrictions

You may not copy the Software except as expressly stated in writing in the present agreement. Any permitted copy of the Software that you make must contain the same copyright and other proprietary notices that appear on or in the Software.

You may not modify, adapt or translate the Software. You may not reverse engineer, decompile, disassemble or otherwise attempt to discover the source code of the Software except to the extent you may be expressly permitted under applicable law to decompile only in order to achieve interoperability with the Software.

The Software may include various applications, utilities and components, may support multiple platforms and languages and may be provided to you on multiple media or in multiple copies. You are not required to use all component parts of the Software, but you may not unbundle the component parts of the Software for use on different Computers. You may not unbundle or repackage the Software for distribution, transfer or resale.

YOU MAY NOT RENT, LEASE, SELL, SUBLICENSE, ASSIGN OR TRANSFER YOUR RIGHTS IN THE SOFTWARE, OR AUTHORIZE ANY PORTION OF THE SOFTWARE TO BE COPIED ONTO ANOTHER INDIVIDUAL OR LEGAL ENTITY'S COMPUTER EXCEPT AS MAY BE PERMITTED HEREIN.

It is forbidden to transfer the rights to use this software without explicit written permission by MediMatic S.r.l.

Except as permitted by the present license, no part of this guide may be reproduced, stored in a retrieval system, or transmitted, in any form or by any means, electronic, mechanical, recording, or

otherwise, without the prior written permission of MediMatic Srl. Please note that the content in this guide is protected under copyright law even if it is not distributed with the related software.

The content of this guide is furnished for informational use only, is subject to change without notice, and should not be construed as a commitment by MediMatic Srl. MediMatic Srl assumes no responsibility or liability for any errors or inaccuracies that may appear in the informational content contained in this guide.

Updates

If the Software is an upgrade or update to a previous version of MediMatic software, you must possess a valid license to such previous version in order to use such upgrade or update. After you install such update or upgrade, you may continue to use any such previous version in accordance with its end-user license agreement only if (a) the upgrade or update and all previous versions are installed on the same device, (b) the previous versions or copies thereof are not transferred to another party or device unless all copies of the update or upgrade are also transferred to such party or device and (c) you acknowledge that any obligation MediMatic may have to support the previous version(s) may be ended upon the availability of the upgrade or update.

No other use of the previous version(s) is permitted after installation of an update or upgrade. Upgrades and updates may be licensed to you by MediMatic with additional or different terms.

Warranty

MediMatic warrants that it has the right to license the Software to you and that the Software, as delivered to you, will function substantially as described in the Documentation for a period of one (1) year (the "Warranty Period") following the date of Software purchase. MediMatic's sole responsibility under the warranty with respect to any material defects in the Software shall be its use of commercially reasonable efforts to correct such defect or to replace the Software within a reasonable period of time. If MediMatic is unable to so repair or replace the Software, MediMatic shall refund you the portion of the Software license fee that relates to the defective portion of the Software. MediMatic further warrants that all services provided by MediMatic shall be performed in a professional manner appropriate to industry standards. Notwithstanding the foregoing, MediMatic shall have no responsibility for, and this warranty shall be voided in the event of: (a) errors or defects caused by your neglect, misuse, or damage to the Software; (b) use of the Software with any technology not in accordance with MediMatic's written recommended configurations; or (c) any loss or damaging dissemination of data due to breach of security technology used by you in connection with the Software. MediMatic does not warrant that the operation of the Software will be uninterrupted or error-free. Except as expressly set forth in the Agreement, MediMatic makes no other warranty of any kind, either express or implied, including, but not limited to, the implied warranties of merchantability, fitness for a particular purpose, title, and non-infringement.

You understand and acknowledge that the Software involves the use of computer applications performing sophisticated operations in order to assist you in the practice of medicine, but that the Software is not a substitute for competent human intervention and judgment. You shall be solely responsible for the accuracy and adequacy of information and data furnished for processing and any use made by you or your employees of the output of the Software or any reliance thereon. Regardless of whether this Agreement or any remedy hereunder may be considered to have failed of its essential

purpose, in no event shall MediMatic's cumulative liability to you, your patients or other parties for any and all claims regarding the Agreement, regardless of the form of action, whether arising under contract, tort, warranty, or otherwise, exceed the amount paid by you to MediMatic: (a) for the software fee; (b) for the software support and maintenance fees paid during the twelve months prior to the date this cause of action first occurred; or (c) for that portion of the services that related to the specific liability. Notwithstanding the foregoing, under no circumstance shall MediMatic, or its third party suppliers of component software if any, have any liability to you or any third party for any consequential, incidental, indirect, special, or punitive damages or costs, including, but not limited to: loss of revenue, profits, services, data or equipment; interruption of business; costs of substitute goods or services: computer malfunction, viruses, or other digital hazard; unavailability or inaccuracy of any data in the software; any action or non-action taken by you in reliance upon the software; medical malpractice claims, and any and all other circumstances, regardless of whether MediMatic has been advised, knew, or should have known of the possibility thereof. You understand that the fees charged by MediMatic in this Agreement reflect the allocation of risks expressed by the limited warranties, the exclusive limited remedies for breach of those limited warranties, and the limitations of liability damages that are set forth in this Agreement. By using the Software you accept these terms and affirm that you understand that to change them would affect the economic bargain expressed in this Agreement.

MediMatic, at its own expense, shall defend, indemnify and hold you harmless against claims that the Licensed Material constitutes an infringement of a patent, copyright, or other proprietary rights of a third party (except for those claims for which you indemnify MediMatic, below), provided you permit MediMatic, in its sole discretion, to defend or settle the claims. If the Software is held to constitute an infringement of a copyright, patent, or other proprietary right of a third party, and your of such Software is enjoined, MediMatic shall, at its sole expense and option: (i) procure for you the right to continue using the Software free of any liability for infringement or violation; (ii) replace or modify the Software with equivalent, non-infringing Software; or (iii) upon your return of the infringing Software, provide you a refund of the amount you paid to MediMatic for the infringing item(s), less depreciation based on a five (5) year straight-line depreciation schedule. THIS PARAGRAPH STATES THE ENTIRE OBLIGATION OF MEDIMATIC FOR INFRINGEMENT OF TRADEMARKS, PATENTS, OR OTHER INTELLECTUAL PROPERTY RIGHTS.

You shall indemnify, defend, and hold harmless MediMatic, and its officers, directors, employees, agents, successors, and its third-party suppliers of software components ("MediMatic Indemnitees") for, from, and against any claim or action brought against MediMatic by or on behalf of any of your patients or by or on behalf of any other person or third party arising out of the acts or omissions of your employees or agents. Moreover, you shall indemnify, defend, and hold harmless the MediMatic Indemnitees for, from, and against any claim by any third party against MediMatic based on: (a) any error, defect, or inaccuracy in the Software or the Medical Data caused by you; (b) your modification of the Medical Data, provided that the foregoing claim or action would not have arisen but for such modification; or (c) the infringement by the Software of any proprietary or intellectual property rights of a third party resulting from your design, ideas, specifications, or alteration of the Software; or (d) use of the Software in combination with any non-MediMatic items except those specifically approved in writing by MediMatic.

The indemnification obligations of both Parties herein are contingent on the indemnitee providing to the indemnitor: (a) prompt written notice of existence of any claim; (b) full cooperation in the settlement or defense of such claim; (c) within ten (10) days of receiving such claim, all information

reasonably necessary to defend or settle such claim; (d) all information within indemnitee's control necessary for the indemnifying party to conduct a defense; and (e) sole control of the defense or settlement negotiations. The indemnitee may participate in the defense or settlement at its own expense.

The Parties hereby acknowledge that their personnel may access information that the other Party deems to be confidential, proprietary and of commercial value (the "Confidential Information"). As used herein, Confidential Information means: (a) any and all proprietary or confidential business information or data related to the disclosing Party, or such Party's operations, employees, services, patients, or customers, including any such information of which the receiving Party becomes aware as a result of its performance under this Agreement, and (b) Trade Secrets, whether in oral, written, or electronic form. With respect to MediMatic, the MediMatic Confidential Information shall include for example the Software and the Documentation, and any source code, object code, data structures, methods, algorithms, flowcharts, and other materials related to the Software. As used herein, "Trade Secrets" means information related to the business of the disclosing Party that derives actual or potential economic value from not being generally known to or readily ascertainable by other persons and that the disclosing Party uses reasonable efforts to maintain as secret. Each Party shall only disclose the Confidential Information to its employees or agents that have a strict need to know, and shall use at least a reasonable standard of care to prevent disclosure of the other Party's Confidential Information. Without limiting the foregoing, neither Party shall disclose any terms of this Agreement to any other entity without prior written consent. The receiving Party shall immediately notify the disclosing Party upon gaining knowledge of any disclosure, loss, or use of the disclosing Party's Confidential Information in violation of this Agreement. The limitations within this paragraph shall not apply to any information that: (i) was in the public domain at the time of disclosure or became a part of the public domain after disclosure through no fault of the receiving Party; (ii) was independently developed by the receiving Party; (iii) was provided to the receiving Party by a third party who had a lawful right to such information without a breach of duty owed to the disclosing Party; or (iv) is required to be disclosed by judicial or administrative process or by law or regulation, provided that the Party so required to disclose the Confidential Information shall notify the other Party and shall provide reasonably necessary information to enable the other Party to take action to protect such Party's interests. The covenants of confidentiality set forth herein shall survive the termination of this Agreement for a period of three (3) years.

Term, Termination and Disputes

This Agreement is effective until terminated according to the procedures specified below.

You may terminate this Agreement if MediMatic S.r.l. materially breaches any of the terms and conditions of this Agreement and fails to remedy that breach. Upon material breach of this Agreement by MediMatic S.r.l., you shall notify MediMatic S.r.l. of its claim in writing and MediMatic S.r.l. shall have ninety (90) days to remedy the breach. If the breach is not so remedied, you may terminate this Agreement. Upon such termination of this Agreement by you, you shall immediately return the Software, Documentation, and any other MediMatic Confidential Information to MediMatic, and permanently destroy any such materials that may be stored in any non-returnable form including, but not limited to, electronic, paper, fiche, etc. You shall nevertheless be responsible for all outstanding payments for products and services provided and expenses due up to the date of termination, excepting only payment for defective product components or services which are the proximate cause of the breach

MediMatic may terminate this Agreement if you materially breach any of the terms and conditions of this Agreement and fails to remedy that breach. Upon material breach of this Agreement by you, MediMatic shall notify you of its claim in writing and you shall have ninety (90) days to remedy the breach. If the breach is not so remedied, MediMatic may terminate this Agreement, and you shall immediately return the Software, Documentation, and any other MediMatic Confidential Information to MediMatic, and permanently destroy any such materials that may be stored in any non-returnable form, and shall immediately pay all unpaid amounts which would have been due MediMatic after the complete delivery of software licenses, services and support.

The parties hereto declare to be willing to settle, in a friendly way, any dispute arising about the validity, interpretation and/or performance of these terms. Prior to bringing any action in a court of law or to terminating this Agreement, the Parties agree that senior management representatives ("Representatives") from each Party will meet, in person or telephonically, within twenty (20) business days of a Party's written request for resolution of a dispute. The Representatives will attempt in good faith to resolve any disputed issues. No action or notice of termination shall be brought or shall be effective until 20 (twenty) business days after such meeting has taken place, or such later date to which the Parties mutually agree. Should the dispute not be resolved during this meeting or a later date agreed to by the Parties, then the aggrieved Party or Parties may bring notice of breach or termination.

All provisions of this Agreement relating to disclaimers of warranties, limitation of liability, remedies, damages, confidentiality, and any other terms that either expressly or by their nature should survive, shall survive any termination of this Agreement, and shall continue in full force and effect.

Assignability, Notice and Force Majeure

You may assign its rights and delegate its duties and obligations hereunder to an acquiring or successor entity upon written notice to MediMatic. MediMatic may assign its rights and delegate its duties and obligations hereunder to an acquiring or successor entity upon written notice to you or throughout communication on the MediMatic's web site.

Any notice hereunder shall be deemed to have been duly given only if submitted in writing and delivered by (i) certified mail, return receipt requested, (ii) facsimile, or (iii) reputable overnight courier service, to the address of the receiving party as set forth herein or such other address as such party may indicate. Notices shall be deemed delivered when received by the party being notified. The failure by any Party to exercise any right provided herein shall not be deemed a waiver or forfeiture of any such right.

Neither Party shall be liable for any default in its obligations under this Agreement due to natural disasters, acts of God, riots, war, epidemics, labor disputes, governmental restrictions, mechanical or electrical breakdown, interruption of utility services, shortages or delays in obtaining suitable materials, transportation difficulties, acts of subcontractors, or any other acts which are beyond either Party's reasonable control.

Independent Contractors

The Parties agree that each is performing its obligations hereunder as an independent contractor, and no joint venture, partnership, employment, or other relationship is being created by this Agreement.

Neither Party has any express or implied right, power, or authority to enter into any agreement or any commitment on behalf of the other.

Governing Law

Without limiting any other provision of this Agreement, each party shall be solely responsible for its own compliance with all applicable laws and regulations. This Agreement shall be construed in accordance with the substantive laws in force in Italy without respect to its conflicts of laws principles, and any action or proceeding related to this Agreement shall be brought only in the courts located in Genova, Italy which shall have exclusive jurisdiction over all disputes relating to this agreement. No legal action related to this Agreement, regardless of its form, may be brought by either party more than two (2) years after the cause of action first accrued.

VP8 CODEC

MediMatic DICOM Web Compression 1 is a non standard, lossy DICOM compression syntax that makes use of Google® VP8 Codec that is subject to the following licensing terms.

Copyright (c) 2010, Google Inc. All rights reserved.

THE VP8 CODEC SOFTWARE IS PROVIDED BY THE COPYRIGHT HOLDERS AND CONTRIBUTORS "AS IS" AND ANY EXPRESS OR IMPLIED WARRANTIES, INCLUDING, BUT NOT LIMITED TO, THE IMPLIED WARRANTIES OF MERCHANTABILITY AND FITNESS FOR A PARTICULAR PURPOSE ARE DISCLAIMED. IN NO EVENT SHALL THE COPYRIGHT HOLDER OR CONTRIBUTORS BE LIABLE FOR ANY DIRECT, INDIRECT, INCIDENTAL, SPECIAL, EXEMPLARY, OR CONSEQUENTIAL DAMAGES (INCLUDING, BUT NOT LIMITED TO, PROCUREMENT OF SUBSTITUTE GOODS OR SERVICES; LOSS OF USE, DATA, OR PROFITS; OR BUSINESS INTERRUPTION) HOWEVER CAUSED AND ON ANY THEORY OF LIABILITY, WHETHER IN CONTRACT, STRICT LIABILITY, OR TORT (INCLUDING NEGLIGENCE OR OTHERWISE) ARISING IN ANY WAY OUT OF THE USE OF THIS SOFTWARE, EVEN IF ADVISED OF THE POSSIBILITY OF SUCH DAMAGE.

3. LAUNCHING THE APPLICATION

3.1. START-UP PROCEDURE

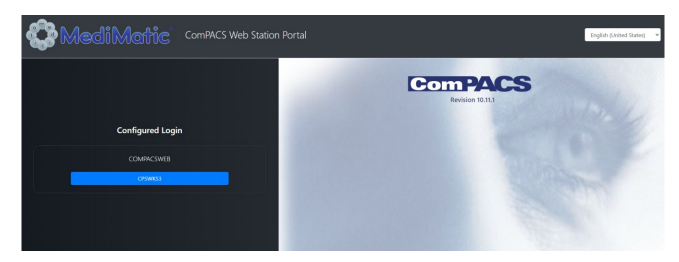
To start the ComPACS Review Station simply double-click on the icon that is on your desktop.



or connect to the ComPACS Web Server portal, generally: /webserver">https://servername>/webserver.

When the web server portal is opened, the landing page verifies if the software is installed and up to date.

The following page will then be displayed to allow the user to select the appropriate login configuration among the ones available and licensed to the connecting client.



If only one configuration is present, as in the example above, the browser will automatically prompt to launch the application. If more configurations are present, the user can select the appropriate one by pressing the related blue button.

If the software is not installed or it is not up to date, the web page will prompt the user to install or upgrade application.

If it is the first installation or if the local *ComPACS Installation Service* is not present, after the download operation, the user will have to execute the install package and follow the related instructions. Alternatively, the user will be prompted to launch the ComPACS *AppManager* to perform the application upgrade.

The review station installation or upgrade, though it is an intuitive and guided operation, is designed to be performed by qualified IT professionals.

After installation is completed successfully, it is mandatory to close and restart the web browser for the software to be successfully detected.

The present operator's guide for ComPACS plugged-in modules documents their use integrated into the ComPACS Review Station. In case such components are integrated into a third-party product please refer to such product operator's guide for specific and additional instructions.

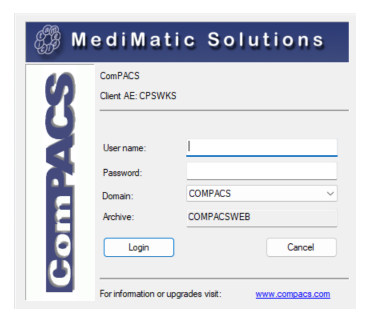
ComPACS Report Manager operates together with the Reporting and Off-line analysis modules. If none of these modules are activated, you will be prompted to activate at least one of them before proceeding.

3.2. LOGIN

Once launched, the software needs some time to configure and to load the application and the installed modules correctly. A splash window will be displayed during this operation.



Then a window will prompt you to login as follows.



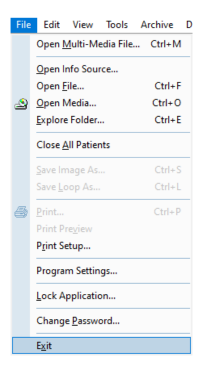
In this window you must enter your *User Name*, your *Password* (generally the *Microsoft® Active Directory /* Windows® login to which your user name belongs) and select the login *Context* or *Domain* from the available choices. The last parameter is generally set to a default value.

ComPACS does not force you to use an independent authentication mechanism but exploits the same user name and passwords used to log on the Windows® OS.

If Single Sign On is enabled for the default *Archive* configuration, the system will automatically proceed to authentication on the basis of the current Windows logged on user. In order to inhibit such functionality, keep the *Shift* key press while the application is launching.

3.3. SHUTDOWN PROCEDURE

In order to shutdown the application, you need to select the Exit item from the File menu.



Alternatively, you may also click on close button of the application primary window or windows:



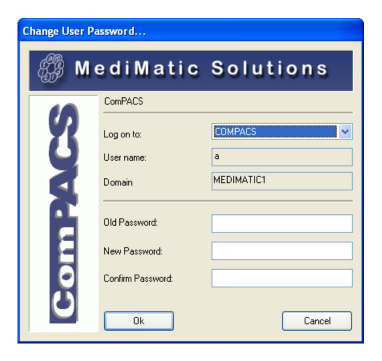
If there are unsaved items, you will be prompted to save them or discard changes.

3.4. CHANGING PASSWORD

In order to change the log-in password, you need to select the Change Password item from the File menu.



A dialog window will display the main log-in information of the Archive you are referring to (Archive Configuration, User Name and Domain) and prompt you to enter the current password (for safety purposes) and the new password. The new password also needs to be retyped for confirmation.

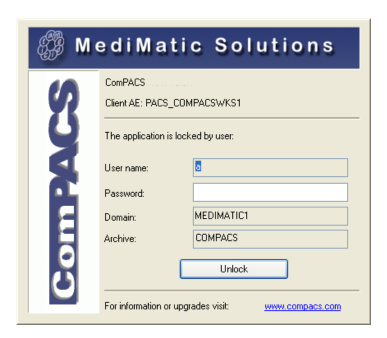


ComPACS exploits the same user name and passwords used to log on the Windows® Domain. This means that changing the password within ComPACS will also require changing the password to access the Windows® system.

The new password must follow the rules and restrictions (minimum length, complexity, etc..) defined by the related Windows® Domain.

3.5. IDLE TIME PROTECTION

In order to lock the application, you need to select the *Lock Application* item from the *File* menu. The following dialog window will prompt you to enter the logged in user credentials before re-accessing the application.



The application will also lock automatically after a period of user inactivity is reached on the workstation. This period of time is defined by the system. A second timeout can also be defined to automatically exit the application.

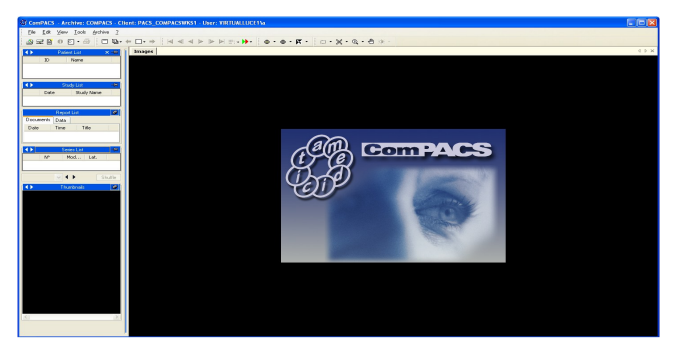


If an item such as a report is in edit mode the application will not exit, but will only lock.

4. APPLICATION LAYOUT

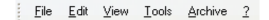
The appearance of some objects within the viewer may vary. It depends on the installed modules.

Once you have started the program, the main window will appear.



The following are the four key components to the main window.

1) The menu bar, on the top of the screen. On the menu bar you can open, save, print, edit, and close data. You can also retrieve information and set the general properties of the software;

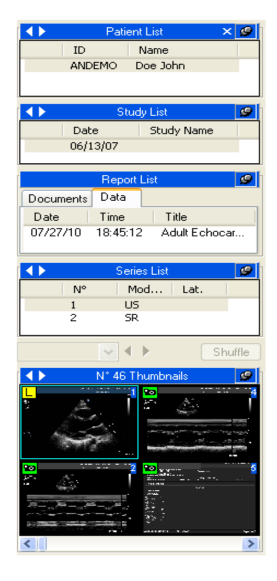


The menu bar can differ in relation to the optional modules installed. For example, the *Archive* menu is only present if a local or remote archive is connected. Additional menus or submenu items may also be present.

2) The tools bar just under the menu bar. On this bar there are various icons that represent tools (i.e. filters, zoom, etc.) that you can apply to the selected image. In addition, there are play commands, and buttons to enter the archive (if installed) or to open a media. Additional buttons to create reports may also be present if the related modules are installed.



3) The controls window on the left allows you to have direct control on a patient, a study, a study documents and reports, a series and on the images (that are displayed as thumbnails). Here you can retrieve the information (on patient, studies, documents and reports, series, images). You can archive, you can attach files, and write a memo if the related modules are installed.



4) Finally, the tabs window is the main window of ComPACS Review Station, where the images and the loops are displayed. If you are searching for a media the explore window appears in the tabs window, indicated as an additional tab at the top; the attachments, the memos, and the reports (if the related modules are installed) are all displayed in the tabs window.



5. QUICK START

ComPACS Review Station automatically recognizes DICOM media. To view the content of a media simply put it in the applicable media reader (for example CD/DVD drive or Magneto-Optical drive) of your computer.

Once the content of the media is interpreted, ComPACS automatically opens an explorer interface.

A *DICOMDIR* file has to be present in the root directory of the media otherwise the media is not recognizable by the software. The ComPACS review station can read some proprietary file formats as well, such as *DSR-TIFF* and *DEFF*.

If the media is not automatically detected it is possible to access and explore the media content by clicking the Open Media button on the toolbar.

A similar operation can be performed to access the central archive instead of the media content. To do so, click the ** Search Archive* button on the toolbar. Please refer to Archive sections for more detailed instructions.

At this point the list of the studies is presented, displayed with some associated information and image thumbnails. To select a patient select it and click on open, or simply double-click on it. The same operation can also be performed at the study, series, or image level.

The selected items will be then opened for review. The service window on the left will display the information about the selected patient, the study, the series and the images' thumbnails. The *Images* tab window will display the first image or loop selected. The picture below shows a sample screenshot.



If the viewer is configured to display the patient and study information header, such information will be displayed at the top of the "Images" tab.

If multiple studies for the same patient or multiple patients are displayed, then the header will also convey this information to the user.

5.1. QUICK ACCESS TO MAIN FUNCTIONS

If the DICOM items are image loops, they can be played by using the play commands on the tool bar.

- Forward play / Stop if re-clicked
- Backward play / Stop if re- clicked
- and together Sweeping play
- Next frame
- Previous frame

- Last frame
- First frame

Or by the smart keys on your keyboard.

- → Play Forward / Stop
- **Pg**↑ Previous Thumbnail
- **Pg**↓ Next Thumbnail
- ← Previous Frame
- → Next Frame
- ↑ First Frame
- ↓ Last Frame
- **F1** On-Line Help
- **F2** Toggle Normal/Full Screen view

By clicking the right mouse button when the cursor is in the tabs window, a quick access menu will appear to allow a rapid selection of some program function (i.e. zooming, display mode selection, information, etc.).

If the mouse wheel is configured for image navigation, by moving the mouse wheel when the cursor is over the film window it is possible to change the images displayed. Alternatively, the mouse wheel is used to select a frame within a loop.

6. REVIEW STATION

6.1. OPENING FILES

6.1.1 FILE FORMATS

Opening and closing a file set through the *DICOMDIR*, (the file that according to the DICOM 3.0 standard contains all the information on the images present in the media support), is typically performed automatically by the program that detects respectively, media insertion and removal.

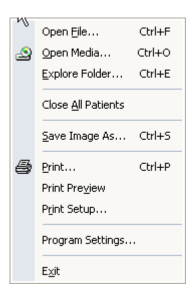
The *DICOMDIR* file has to be present in the root directory of the considered media. ComPACS viewer can read some proprietary file formats as well, such as *DSR-TIFF* and *DEFF*.

6.1.2 OPENING A FILE FROM REMOVABLE DRIVES

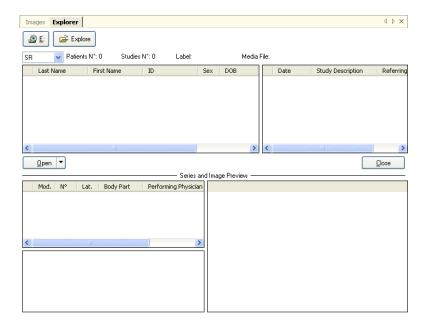
If your computer does not allow the detection of disk insertion and removal, or the *DICOMDIR* file is not in the root of the media, it is still possible to open a DICOM file set by clicking the explore icon



This can be done as well by clicking on the File menu then choosing Open Media (hotkey Ctrl+O)

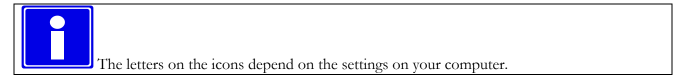


This operation opens in the tabs window, an explorer interface where you can explore the content of a removable drive, of the local hard drive, and of the Local Area Network.

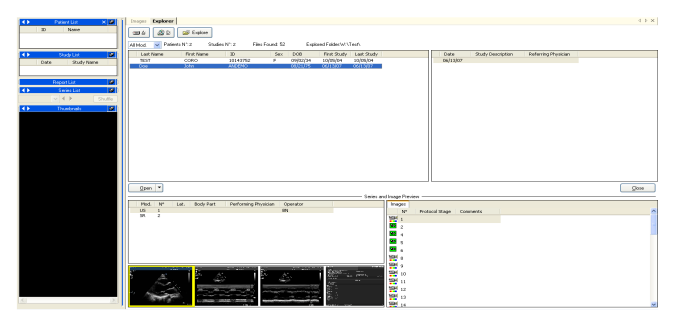


To explore the content of a removable media directly, click on one of the icons representing the removable media drives (CD, DVD, USB-keys etc.). For example:



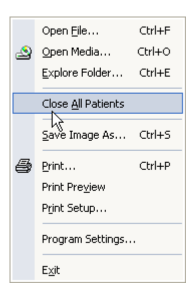


The software searches for the DICOMDIR and if found, it will automatically open the related file set.



A list of studies in the fileset is presented with some related information. To select a patient (or a study) simply double-click on it. If you wish to open just one image or loop, double-click on the related thumbnail and similarly, for single studies or series. Please note; double clicking the patient name will open and load all studies of this patient that are in the media you are querying. If the patient has had numerous studies, this will take some time.

The fileset is closed automatically when the medium is extracted. The fileset can also be closed by the *Close All Patients* command in the *File* menu.



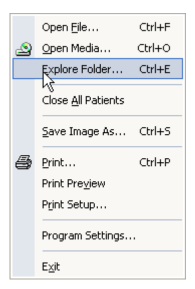
The same operation can also be performed from the patient list window by right-clicking and then selecting *Close Patient* or *Close all Patients*.

6.1.3 OPENING A FILE FROM LOCAL HARD DRIVES

To open a fileset present on a local Hard Disk, or on a network place, in the explore window, click the explore icon.



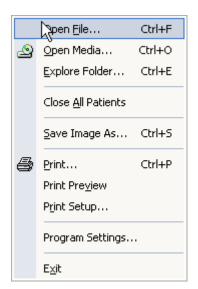
This can be done as well by choosing *File* menu, then *Explore Folder* (hotkey **Ctrl+E**).

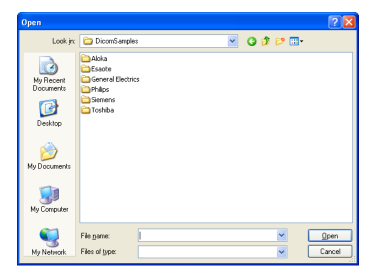


A dialog window appears that allows the selection of the desired folder present on a local disk, on a local device or on a network remote device



Another way to navigate through the folders is to select *Open File* (hotkey **Ctrl+F**) from the *File* menu and then select the specific file or *DICOMDIR* to open.





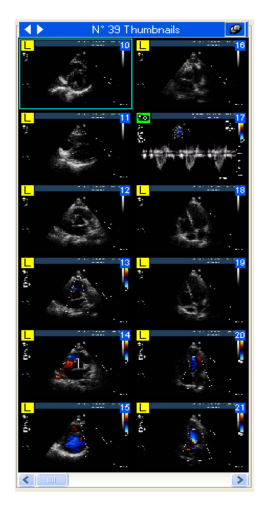
6.1.4 OPENING A PATIENT/STUDY FROM THE ARCHIVE

If a local or remote archive is connected it is possible to access the central archive instead of the media content. To do so, click the Search Archive button on the toolbar. Please refer to the Archive sections for more detailed instructions.

6.2. VIEWING IMAGES AND REPORTS

6.2.1 THUMBNAILS

Once you have loaded the image set to be analyzed, you can see it on the service window as thumbnails.



You can browse it with the mouse, or by using the Control keys, or by using the Numeric pad. Also you can use one of the two buttons created for this purpose.

next thumbnail (hotkey **PgUp**) and previous thumbnail (hotkey **PgDn**).

The thumbnails will display a small key image of the object they refer to if it represents an image or loop. Alternative, if it represents a different type of object such as a waveform or a report, it will display some key information such as the series modality, number and date/time.

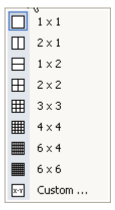
Moreover, the thumbnails can be associated to different symbols to convey additional information about the object they refer to:

- for objects that contain data in a lossy format (for example a lossy DICOM transfer syntax).

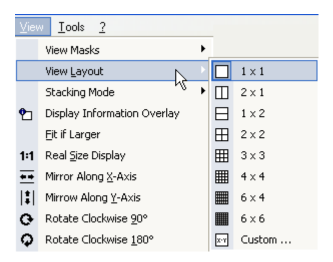
- for objects that represent single frame images.
- for objects that are not supported by your software version.
- for objects that are still being downloaded from the central archive or prepared for display. This symbol will disappear once the download operation has completed and the object is accessible.
- for thumbnails that refer to a series of objects (and not just a single object, image or loop). This is typical for objects that often need to be stacked such as those of MR and CT modality.

6.2.2 IMAGE LAYOUT

ComPACS Review Station allows viewing one or more images at the same time. This can be done by clicking the *layout* icon and then selecting the desired splitting of the screen from the *layout* menu



You can also do this by clicking the View menu and then selecting View Layout



You can customize the division of the screen (the maximum allowed value is 10x10). By double-clicking on a thumbnail, you will see on the screen the image selected and the following images until the

screen is full. You can move as you prefer, the images in the tabs window by simply dragging-and-dropping them.

Please note that it is also possible to specify stacking information for a loop composed of a series of independent single image files (typically for MR and CT series). Additionally you can specify the unstacking of a DICOM loop into independent window views to observe the different frames in the loop side by side. This operation will be addressed in more details in another chapter.

Below as an example, a 3x3 grid is shown; please note that changing the layout does not affect other functions of the software.



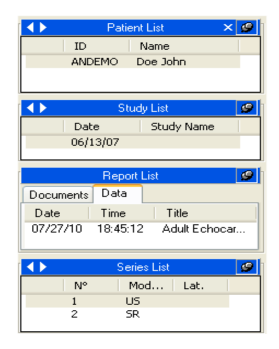
The default stacking mode is *Image Number Order*. Please revert to this mode if you experience some anomalous behavior in the image or loop display.

6.2.3 FULL SCREEN

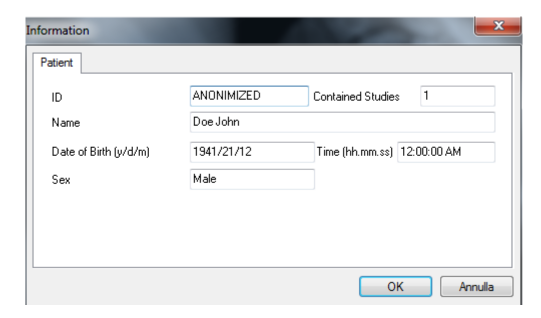
ComPACS Review Station allows the full screen mode. This option can be activated and deactivated using the button in the tool bar, or with the hotkey **F2**.

6.2.4 ADDITIONAL INFORMATION

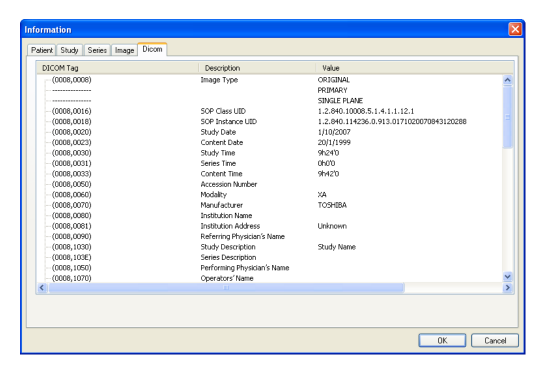
On the service window, together with the thumbnails, you can see three other windows showing additional information about the active items. You have *Patient list, Study list, Report list* and *Series list.* In other words on the service window you can retrieve all the data about a patient, a study, a report or a series that is open for reviewing.



Additional information is available by right-clicking on an item. Then click on the selection, *DICOM Information*. By doing so, you will open a new window (*Information Properties* window) displaying different information about the item selected.



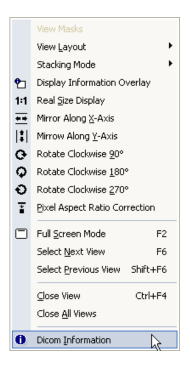
Patient Information



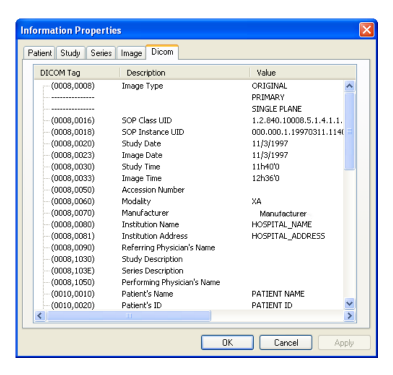
Extended DICOM Information

The tabs *Patient, Study, Series,* and *Image* have the same meaning of the service window menus. The *DICOM* tab contains all the information derived by the interpretation of the data contained in DICOM object itself.

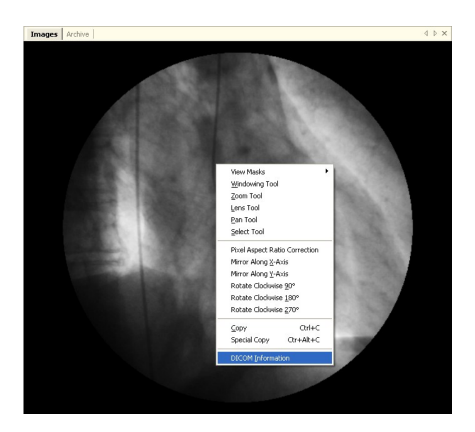
Another way to display this window is to click on the *View* menu and then on *DICOM Information*, or directly by clicking on the *Information* icon on the tool bar.



This procedure will open a similar new window with all the DICOM information and properties of each tab and subsequent field.



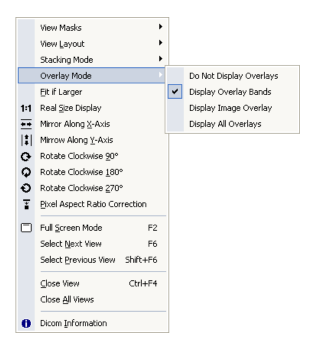
Alternatively, you may Right Click on the image in the tab window and select the DICOM Information item in the popup menu.



6.2.5 INFORMATION OVERLAY

It may be useful to hide or show the overlay information. That means showing information about the patient, study or image directly over the image itself. This can be done by clicking on the overlay icon or by clicking the *View* menu and then *Overlay Mode*.





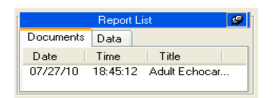
There are two main different types of overlays: a) image bands shown above the image to display main patient, study, b) overlay on the image to show more detailed and configuration information (see the settings chapter for more information).

If the Overlay Information is disabled, it will nevertheless remain on the screen for five seconds whenever different studies or different patients are open for review to warn the user of the change. Please be very careful when in this mode to avoid possible confusion between patients or studies.

A dynamic overlay will also be displayed when the cursor is positioned in the bottom part of the image. The overlay will display compression, series number, image number and frame information. For loops, it also contains a button to start or stop playback and a slider to select specific frames.

6.2.6 VIEWING REPORTS

The "Report List" service window on the left will display the report documents and structured data respectively in the *Documents* and *Data* tabs.



To display the report document simply click on the related item. A new tab will be opened on the right and it will display the selected document.

In order to display the report document it is necessary to have Adobe® Acrobat Reader installed on your system.

The structured data items in the *Data* tab can be analyzed only if the analysis or reporting modules are licensed and activated.

6.2.7 CLOSING ITEMS

To clear the main window while keeping the study open for review, click on *View* menu and then *Close View* (hotkey **Ctrl+F4**).



In case you have more than one image opened (i.e. layout modified) clicking on *Close View* will close only the selected image. To close all the images in the main window (without clearing the thumbnails) click on *Close All Views* or press the button on the tabs window.

Clicking the button on the *patient service window* will close all the opened patients for review and all the related studies. If only a single patient or study needs to be closed simply *right click* the appropriate service window and select the *Close xxx*... option.

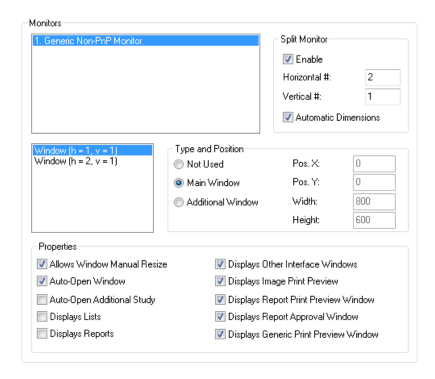
6.2.8 MULTI-MONITOR

The ComPACS Review Station allows using a multi-monitor configuration (if the appropriate module is installed). To access the multi-monitor options click on the *File* menu, then on *Program Settings*.



This opens a window with different tabs and subsequent fields. The tab *Multi-Monitor* allows the user to select the desired options for each monitor. The software settings can be modified to associate specific functions to each monitor.

- Not used.
- Main Window, i.e. Main software interface (containing the service windows).
- Additional Window, i.e. Additional image or report windows.



You can decide which roles are assigned to the additional monitors. Obviously one monitor must be the main window. The others can be selected as Additional Window. Once done, checking the desired options in the properties box assigns that function to that monitor. Remember that if a property is checked for a monitor, it should in general be not checked for the others.

The monitor primary properties are the following:

- <u>Auto-Open Window</u>. This property defines in which monitor the images and loops will be automatically displayed when a study is opened for review. If more than one monitor is selected, the images will be distributed across the selected monitors.
- <u>Auto-Open Additional Study</u>. This property defines in which monitor the images and loops will be automatically displayed when a second study is opened for comparison. If not set, all studies will be opened in the default monitors selected above. If more than one monitor is selected, the images for the studies to be compared will be distributed across the selected monitors.
- <u>Display Lists</u>. This property defines in which monitor the patient and study lists (for example the archive search windows) will be displayed.
- <u>Display Reports</u>. This property defines in which monitor the windows for creating and editing reports will be displayed.

All the remaining properties are self-explicatory.

Furthermore it is also possible to split monitors both horizontally and vertically, emulating a multimonitor configuration and application behavior even in single monitor systems. The position and size of the windows within the monitor can also be defined.

These settings will become operational only after the application is restarted.

Please note that multi-monitor options also depend on the video card installed on your computer. If no multi-monitor function is detected then the related settings will not be available.

6.3. PLAYING LOOPS

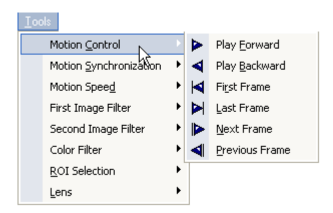
6.3.1 HOW TO PLAY

Once a DICOM file has been opened, ComPACS Review Station displays the image in the tabs window. If the tabs window is not large enough to host a full size DICOM image it will be automatically resized to fit in the window.

You have different ways to control the loop playing. You can use the play icon as follow

- Forward play / Stop if re-clicked
- Backward play / Stop if re-clicked
- and together Sweeping play
- Next frame
- **◄** Previous frame
- ► Last frame
- First frame

All these commands are also included in the Tools menu, selecting Motion Control

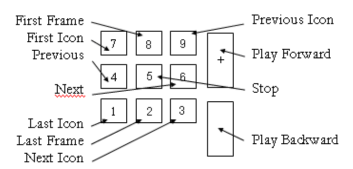


Also, a dynamic overlay will be displayed when the cursor is positioned in the bottom part of the image. For loops, this overlay contains a button to start or stop playback and a slider to select specific frames.

There is another way to control the flow of images. You can use the Control keys

- → Play Forward / Stop
- **Pg**↑ Previous Thumbnail
- **Pg**↓ Next Thumbnail
- ← Previous Frame
- → Next Frame
- † First Frame
- ↓ Last Frame

Finally the Numeric Pad may be used as follows:



By clicking the right mouse button when the cursor is in the tabs window, a quick access menu will appear to allow a rapid selection of some program function (i.e. zooming, display mode selection, information, etc.).

If the mouse wheel is configured for image navigation, by moving the mouse wheel when the cursor is over the film window it is possible to change the images displayed. Alternatively, the mouse wheel is used to select a frame within a loop.

6.3.2 SPEED CONTROL AND TRIMMING

Press the *Playback Settings* button to have access to a menu with which you can control the speed of playback and set different limits of the loop in terms of time or frame.

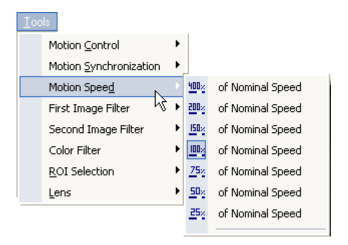


The speed can be changed between 400% and 25% of real time speed. The time and the frame limits can be modified and the progress bar shows the changes.

If Cardiac ECG R-Wave information is present in the DICOM file, a button for each beat is also displayed. By pressing one or more buttons, it is possible to limit the loop to play the selected beats.

The requested speed can only be obtained if sufficient computational resources are available to accomplish the operation. Otherwise a reduced playback speed will occur.

The same window is accessible by *Tools* menu and then clicking on *Motion Speed*.



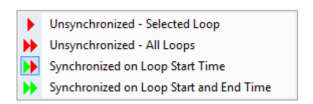
In this dialog, you may trim the loop to be displayed by selected the start and end portion and respectively clicking the and arrows. The markers in the bar will change accordingly.

These settings will become operational only after the application is restarted.

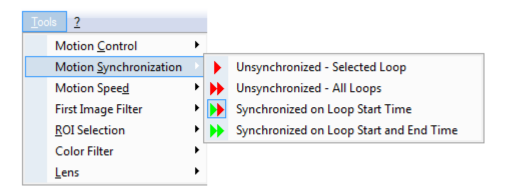
These settings are just temporary and will be lost the next time the loop is re-opened for display.

6.3.3 MULTI LOOP CONTROL

In case you are analyzing more than one loop (i.e. layout changed) you can easily control the synchronization of different loops. You can enter the *Synchronization* window by clicking the button. A drop down menu will be displayed.

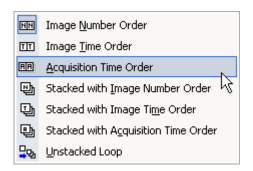


Most menu items have titles that are self-explanatory. This window allows, for example, to synchronize only on the start frame , or on the start and end frame etc. The same could be obtained by using the *Tools* menu and then choosing *Motion Synchronization*.



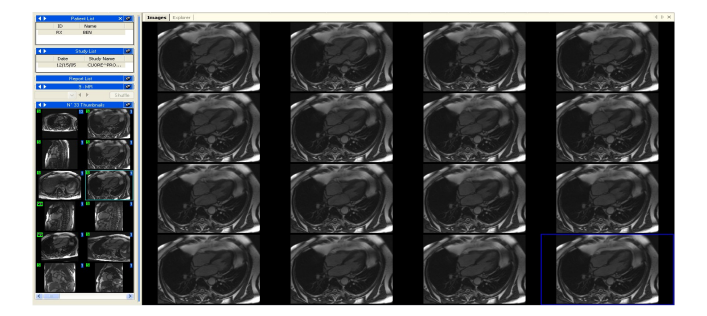
6.3.4 STACKING/UNSTACKING

Stacking means that you can build a loop with a series of single frames (for example a series of CT or MR images) or vice versa you can divide a loop in single images to analyze the video frame-by-frame. It is possible by accessing the stacking menu and using the



In this menu, you may decide what kind of sorting you need. The default behavior is to display images independently and sort by *Image Number Order*. This ordering information is generally generated by the modality device upon acquisition and conveys the order in which the images were effectively acquired. Other possible, but less often used, alternatives are to order using the *Image Time* (the time at which pixel data creation started) or *Acquisition Time* (the time at which the acquisition of the data resulting in this image started).

For example the next screenshot displays an MR series in which the different slices are displayed in a grid. Clearly it is important to note that before doing any display operation, you should set the desired layout (in the following case the layout grid is 4x4).



The system also allows a series of MR or CT images to be stacked to be displayed and processed as if it were a loop. In this case it is important to define how the images are going to be ordered in this loop, generally choosing the *Stacked with Image Number Order* option. Once this option is selected, all open operations that refer to a *series thumbnail* will be stacked into a loop.

In the opposite way the system also allows a loop to be unstacked into independent frames, shown side by side. To activate this option, select *Unstacked Loop* as the stacking mode. Once this option is selected, all open operations that refer to a loop will be unstacked into multiple image views.



Most of the icons presented take the look of the last choice done by the user.

6.4. FILTERING IMAGES

There are two different filtering options: a color filter and a dynamic filter.

6.4.1 COLOR FILTER

To add a color filter to the images click on the Color Filter button on the tool bar, and from the menu that appears, you can choose one of the following filters.



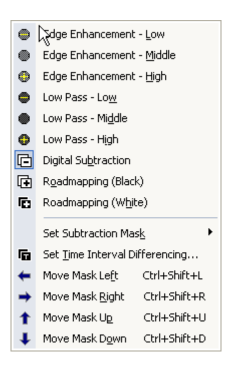
To access this window you can also click on the Tools menu and then Color Filter.

This filter works with each kind of analysis (single image, multi image, and loop). To remove the effects of the filter, just re-click the *Color Filter* button.

6.4.2 DYNAMIC FILTER

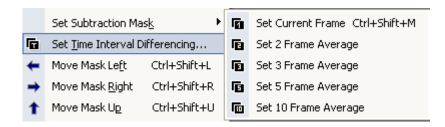
Another set of filters available are the *Dynamic Filters* or *Image Filters*. These filters are characterized by buttons such as or (The list of possible buttons is displayed in the menu below). By clicking the *Image Filter* button you open a window where several kinds of filter are available to use.

As you can see in the following figure, by selecting a filter you impose the look of the related button. This menu is also available by clicking on *Tools* menu and then on *First/Second Image Filter*.

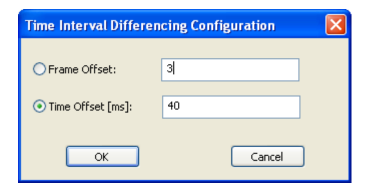


If the *Digital Subtraction* filter is chosen from the menu, the system automatically applies the subtraction masks as defined in the DICOM file. If no mask is defined, a set of additional operations is required. You then need to set a subtraction mask for a single frame with the hotkey **Ctrl+Shift+M** or choose one of the other options from *Set Subtraction Mask* menu, as shown below.

If the *Digital Subtraction* filter is chosen from the menu, a set of additional operations is required. You then need to set a subtraction mask for a single frame with the hotkey **Ctrl+Shift+M** or choose one of the other options from *Set Subtraction Mask* menu, as shown below.



There is another mask control for the Digital Subtraction filter. You can decide to use a dynamic mask by clicking *Set Time Interval Differencing* on the previous menu. This operation opens a window where you can decide the offset of the mask in terms of frame or time interval.



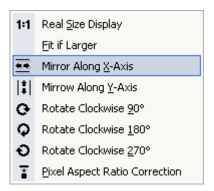
Once set, you can also move a mask along the image. This is useful in case where there are small movements which need to be adjusted. You can move a mask up (hotkey Ctrl+Shift+U), down (hotkey Ctrl+Shift+D), left (hotkey Ctrl+Shift+L) and right (hotkey Ctrl+Shift+R). The same options can be accessed from the *Image Filter* menu.

It is important to underline that the filtering implies a computationally intensive operation that reduces playback efficiency. For this reason we suggest avoiding its use during cine loop viewing in slow system configurations.

6.5. TRANSFORMING IMAGES

6.5.1 MIRROR AND, ROTATION

ComPACS Review Station allows rotation, mirroring, and zooming on the images and loops. To do this, click on the *Zoom/Transforms* button and obtain access to the following menu.



Here you have several options concerning rotation and mirroring. Please note that there's the possibility to correct the size of the displayed image to fit it to the screen, display it in real size or correct the pixel aspect ratio (square pixel transform). The same menu is available by *right clicking* on the tabs window.

6.5.2 INTERPOLATED ZOOM

For *Zooming*, click once on the button. The mouse pointer will now resemble the button itself. In this situation use the mouse wheel to obtain a zoom in/out effect. To return to the default settings, simply *double-click* on the view window.

The Zoom tool is also available by right-clicking on the image and then choosing the Zoom tool.

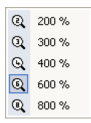
The quality of the display and speed of the zoom depends on the hardware installed. It is important to note that zooming on some platforms is a computationally intensive operation that reduces playback efficiency.

6.5.3 PAN

When the zoom is activated and the dimension of the image is bigger than the screen, you can move the screen along the image by selecting the *Pan* button, and then keeping the *left mouse button* pressed and moving the mouse on the view window. This function is available also by *right-clicking* on the image and then selecting *Pan Tool*.

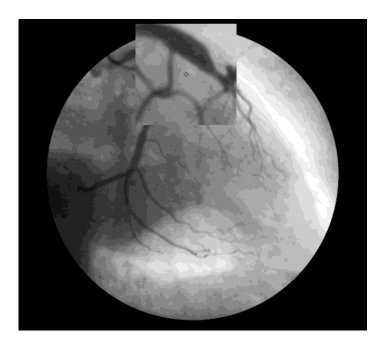
6.5.4 LENS

Another way to zoom the image is the *Lens* tool. This tool is identified by this button that gives access to a *Lens* menu.



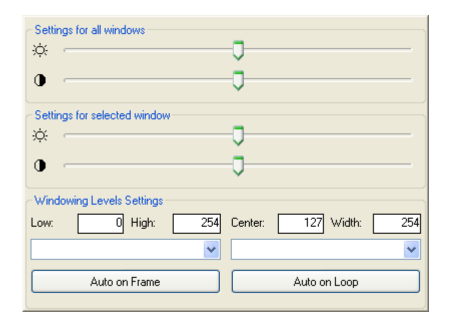
The same menu can be opened by clicking on *Tools* menu and then on *Lens*.

The appearance of the mouse pointer will be similar to a lens. Keeping the *left mouse button* pressed and then moving the mouse along the image creates a lens effect. Refer to the following page for an example.



6.5.5 BRIGHTNESS AND CONTRAST

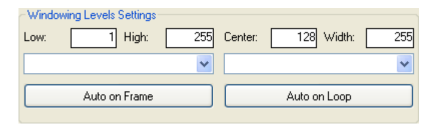
To change the brightness and contrast levels, use the *Brightness* button. It will open a window where you can change the brightness and contrast values for one image or for all the images.



Another way is to click on the button and then keep the left button pressed while over the image. Scrolling the mouse up and down will change the level of contrast; scrolling the mouse right-left will change the brightness level. To reset the original value, double-click on the image.

Another tool present on the above window is the *windowing* tool. A monitor is not capable of displaying more than 256 gray levels therefore if the selected image contains a greater number of gray levels, it is necessary to establish the value range that must be displayed and rescale it to a 256 levels range.

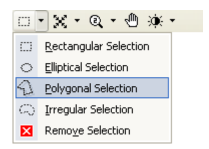
Normally, if the image requires a windowing operation, the possible window(s) are stored with the image in the DICOM file. These automatic settings, if present, are available in the combo box. By selecting an item in the drop down menu the related windowing levels will be applied.



If these settings are not present, it is possible to calculate them automatically by use of the *Auto* buttons or to set them manually in the edit boxes at the bottom of the control window. The boxes indicate the *Center* level of the window and the window *Width*.

6.5.6 ROI SELECTION

It is possible to select a ROI (Region of Interest) on the view window by using the drop down menu will list all the possible choices.



Once the option is activated, you may draw the ROI by using the mouse buttons. The left click adds a point and the right click removes it. The geometry of the ROI will depend on the option chosen.

These ROIs can then be used for all operations that can support them, for example printing or exporting to AVI or to Multi-Media image formats.

6.5.7 MASKING

You may hide or replace portions of the image or loop by applying a mask which will be overlaid onto the image. The opaque portion of the mask will replace the image content whereas the transparent ones will keep the underlying region visible.



Simply right click on the image and access the View Masks menu to manually select a mask or let the system automatically pick the best one out of the available choices. These masks can then be used for

all operations that can support them, for example printing or exporting to AVI or to Multi-Media image formats.

The available need to configured in application properties for each User Profile before they can be used in the application.

6.6. STRESS ECHO

In cases where ComPACS Lite Viewer recognizes a Stress Echo examination in the DICOM fields, it creates a new menu on the service window (above the thumbnails window).



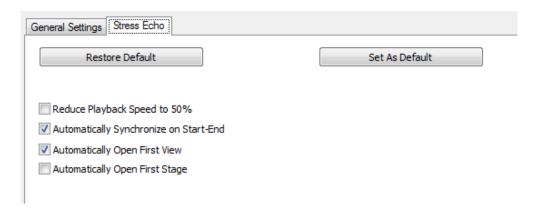
In this menu, all the different stages and views of the examination are available so it is very easy to choose a particular stage by clicking on the *Stress Echo* menu. You can also change stage or view by clicking the small arrows on the right of the menu.



Depending on user settings, ComPACS will automatically open the first view or the first stage. Furthermore the playback speed can be set to 50% and the synchronization to the start-end of the loops. The information about the view, stage and heart rate will be displayed as image overlays.

When the study is closed, the viewing preferences will return to their original value.

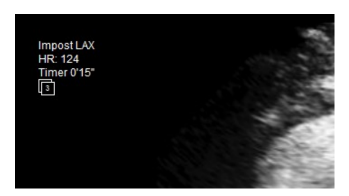
Clicking the *File* menu, then on *Settings* and selecting the *Stress Echo* tab, the user settings can be viewed and modified if necessary.



ComPACS Lite Viewer allows shuffling between a stage and a view by clicking the *shuffle* button.

This shuffling is possible with all views and stages recognized by the viewer in the DICOM header of the file. If such information is not stored in the DICOM file, the stress echo protocol will not be automatically detected.

If more than one acquisition has been performed for the same stage and view, a rectangle, containing the number of such acquisitions, will be displayed as an overlay to inform the user.



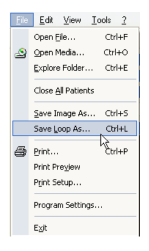
Clicking the left mouse button within the rectangle will display a menu with the list of such different acquisitions. Selecting an item in the list will update the display accordingly. The selection will persist until the study is closed.



6.7. PRINTING OR EXPORTING TO MULTI-MEDIA FORMATS

6.7.1 LOOP EXPORT

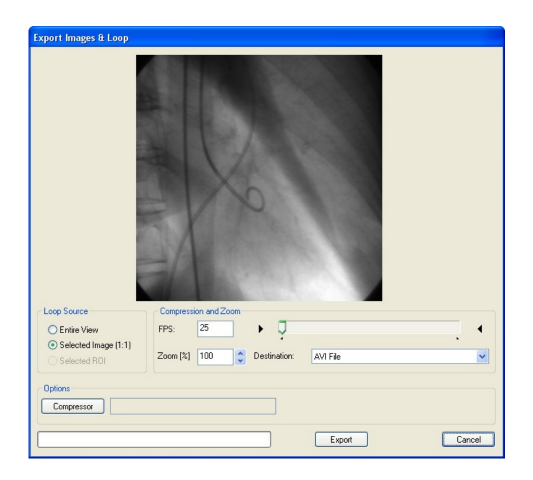
The ComPACS Review Station allows saving loops in AVI or MP4 format to embed them in *Microsoft® PowerPoint* presentations. To save a loop in AVI or MP4 format, click on *File* menu and then on *Save Loop As* (hotkey **Ctrl+L**)



This operation opens a new window where you can set some of the properties of the AVI or MP4 file:

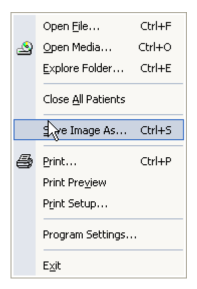
- Trimming, i.e. setting the start and end frames.
- Cropping, i.e. *Entire View*, *Selected Image*, or *Selected ROI*. (Please note that a ROI needs to be first selected in the view window for the *Selected ROI* option to be available.)
- Masking. For this option to be available it is necessary to first select and activate the mask in the view window. The mask will remain valid for all export operations.
- Zoom, expressed as a percentage of the original view.
- FPS Frames per Second.
- For AVI, the Compressor. This exploits the list of AVI CODECS installed on your Windows® OS.
- For MP4, the Quality represented as a percentage value.

Please make sure the CODEC used to generate the AVI file is also available on the system that is intended to open/review the file.



6.7.2 FRAME EXPORT

With the software it is also possible to export a single frame to a multi-media file. To do so, click the *File* menu, and then on *Save Image As* (hotkey **Ctrl+S**).



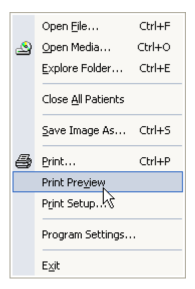
This operation opens a new window where you can set some of the properties of the output file:

- Cropping, i.e. Entire View, Selected Image or Selected ROI. Please note that a ROI needs to be first selected in the view window for the Selected ROI option to be available.
- Masking. For this option to be available it is necessary to first select and activate the mask in the view window. The mask will remain valid for all export operations.
- Zoom, expressed as a percentage of the original view.
- Output format (for example *TIFF*, *JPEG* or *BMP*).
- Compression Quality, where applicable.

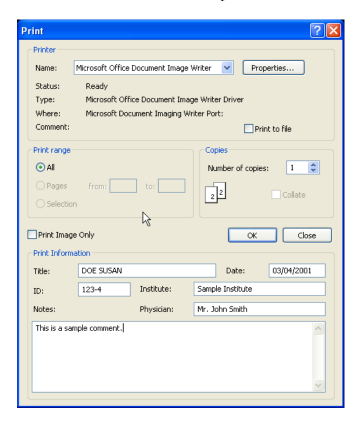
6.7.3 PRINTING IMAGES

Another way to export images is printing.

ComPACS Review Station allows printing images by clicking the *Print* button or by clicking on *File* menu and then *Print* (hotkey **Ctrl+P**).

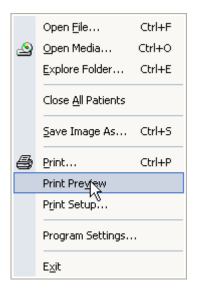


In that window you can hide, show or edit the details of a patient.

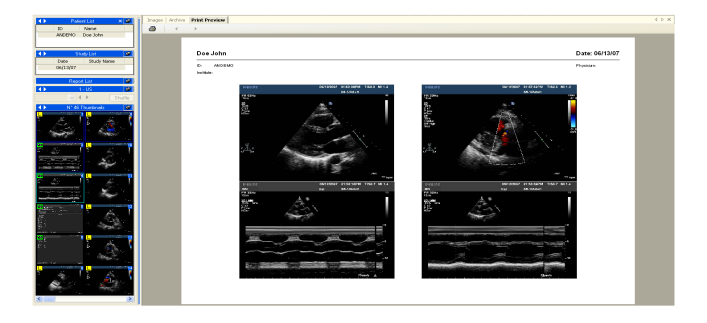


The software also allows setting some printer properties such as page type and margins by clicking on *File* menu and then on *Print Setup*.

Also, before printing it may be useful to have a preview. This is possible with ComPACS Review Station by clicking on *File* menu and then on *Print Preview*.



By doing this a new tab is opened displaying the page to be printed.

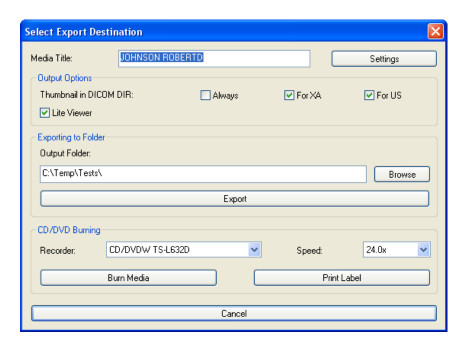


6.8. EXPORTING TO DICOM FOLDERS OR MEDIA

To export selected items to a folder or to removable media you can simply right-click on the image thumbnail, series, study or patient in the *service window* displays and select the *Export* function from the pop-up menu.



The process is practically identical to the archive process. Please refer to the section of this guide that addresses the archive process for a more detailed description. Here you need to click the *Export* button instead of the *Archive* button to activate the function. At this point an additional dialog is displayed, as shown below.

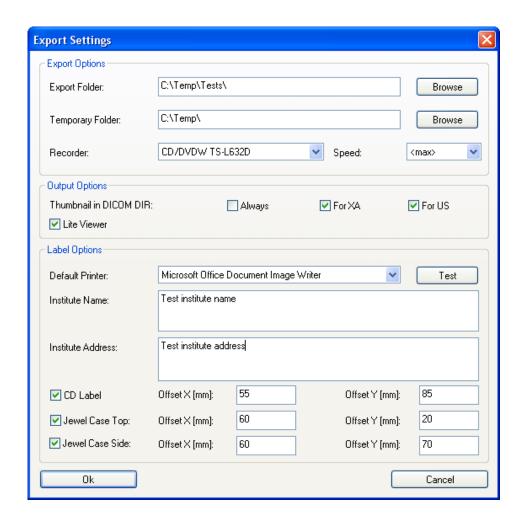


In this window you need to indicate the desired destination, such as a folder or a removable media, as well as some indications on the output format:

• If pre-built thumbnails need to be placed in the DICOMDIR (some viewers are not capable of managing DICOMDIRs with such information).

• If a lite viewer needs to be burnt together with the image set.

These settings can be defined as well as some additional information for label printing by clicking on the *Settings* button.

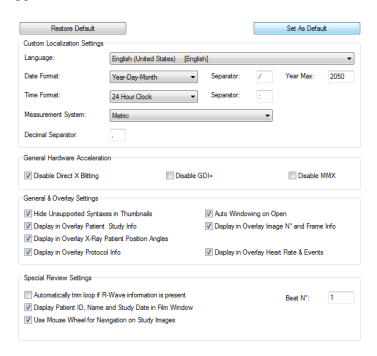


7. PROPERTIES SETTINGS

It is possible to set several general properties of the software by clicking on *File* menu and then on *Program Settings*.



A new dialog window appears.



Here you can set the general parameters to optimize the performance of ComPACS Review Station and to customize some functions. In particular you will notice the following tabs:

<u>General Settings</u> – On this tab you can select the localization preferences (language, measurement system, decimal separator, date & time displays), the hw/sw acceleration features, and the image overlay settings.

In this section it is also possible to automatically trim the loops to a specified number of beats, if the Cardiac ECG R-Wave information is present in the DICOM file. If the number of beats is not specified but the trimming option is enabled, the loop will be trimmed to play all the complete cardiac cycles.

It is also possible to always display the patient and study information in the film window and to define the application behavior when the mouse wheel is used over the film window.

<u>DICOM</u> – On this tab you can select specific DICOM conversion/interpretation options as well as checking if any specific proprietary plugins are installed.

<u>Windowing</u> – On this tab you can select the pre-defined windowing levels to be later selected. For easier access and management, these levels can be arranged in groups and indicated either as *center/width* levels or *low/high* levels depending on your preferences.

<u>Masking</u> – On this tab you can select the pre-defined masks to be applied to automatically anonymize images and loops. For easier access and management, these masks can be arranged in modalities and groups. Different DICOM values can be specified for each mask in order to allow the system to automatically select the best mask for a particular DICOM file.

<u>Multi-Monitor</u> – Addressed in the multi-monitor chapter.

<u>Stress Echo</u> – In this window it is possible to select the specific settings for the Stress Echo Module. Please refer to the Stress Echo chapter for further information.

Changing these properties requires in-depth knowledge of ComPACS Review Station and with the applications of your computer.



Additional tabs may be present in relation to the optional modules installed.

8. THE ARCHIVE

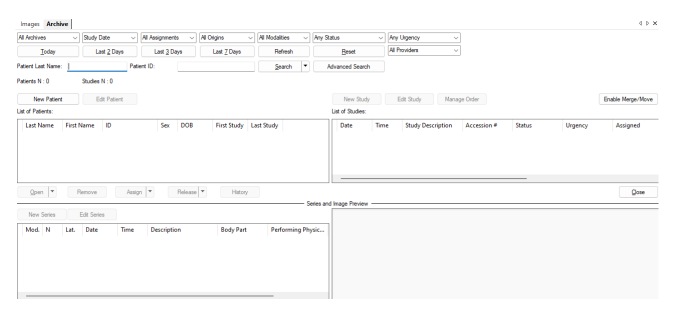
8.1. QUERY & RETRIEVE

8.1.1 NORMAL QUERY

To query the local or remote archive to retrieve data, click on the *Search Archive* button or select *Search Archive* from *Archive* menu.



The Archive interface is opened in the tabs window. In this interface, you can decide the querying option.



In the top portion of the window set the query parameters; the results are displayed in the window in a similar way to the one used to retrieve studies from media.

At the top of the window, by clicking the drop down menus, you can choose different search options and some result filtering information:

- Choosing which Archive to search or if all archives are to be searched simultaneously.
- Choosing the search date criteria. This determines if the queries based on study dates are to be interpreted as the date in which the study was performed, the date in which the study was stored into the archive or the date of the study last assignment.
- Filtering on the assignments (if a study is assigned to a user or a group).
- Filtering on the origins (the Application Entities that stored the object as defined and grouped using the *Web Tool*).
- Filtering on the providers (the entities that are responsible for performing the studies ordered, as defined and grouped using the *Web Tool*).
- Filtering on the modalities to query for.
- Filtering on the status of the study (created, in progress, etc.).
- Filtering on the urgency of the study.

Some filters or query options may depend on Server Side functionality and may not function correctly for all archives.

The top buttons allow the operator to process the query (Search button filtered on the Patient Last Name or Patient ID or predefined date queries such as Today, Last 2 Days, Last 3 Days or Last 7 Days).

The default behavior of the search buttons is to clear the result list and execute the query. Alternatively, by clicking the arrow to the right of the *Search* button, it is possible to search without resetting the results. In this case the new results will be added to the already listed ones.



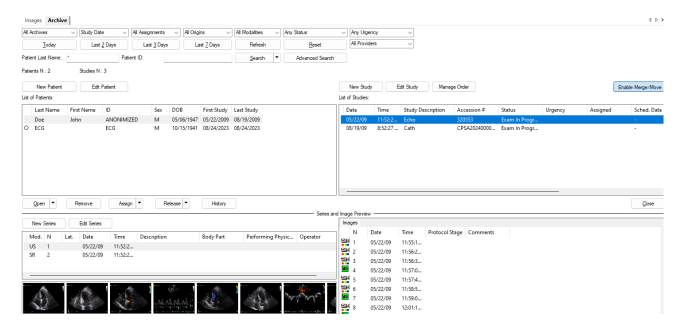
8.1.2 QUERY RESULTS, COLORS AND INFORMATION ICONS

The results are shown in the lists below.

The layout is arranged as follows:

- On the top left hand side are the patients matching the query.
- On the top right are the available studies for the selected patient.
- On the lower left are the available series for the selected study.

- On the lower right are the available object items for the selected series. These objects are organized in tabs according to the different types.
- On the bottom left, if available, a thumbnail display for the identified objects.



To the left of each item (patient, study, series or object) a number of different icons may be present, providing additional information on the status of the corresponding item:

- Attachments. This means that the item has attached one or more external documents such as, Microsoft Word files, Excel sheets, or PDF files.
- Notes. This means that the item has an associated note to inform users of important information.
- *Media*. This means that the item has been backed up on media. If the icon is only partially present it means that only a portion of the item (for example a subset of the images within a study) has been backed up on media.
- Deep Archive. This means that the item has been backed up to a deep archive. If the icon is only partially present it means that only a portion of the item (for example a subset of the images within a study) has been backed up to a deep archive.
- Near Line Data. This means that the item is not entirely present in the on-line archive and needs to be retrieved from a deep archive before being displayed. The retrieve and display time will consequently take a longer time.

At the patient level additional icons will also inform on:

• the urgency status for the studies of the patient the icons refers to. These icons $^{\circ}$, $^{\bullet}$, $^{\bullet}$, are explained in more detail in the PATIENT AND STUDY URGENCY STATUS section.

• the report status for the studies of the patient the icons refers to. These icons , , are explained in more detail in the PATIENT, STUDY AND REPORT STATUS section.

Furthermore the color of text for the query results may change to underline the status of the related item in the archive as follows:

- ONLINE. This means that the item is on line. The text color is standard black text.

- NEARLINE. This means that the item is present only in a deep archive and it needs to be retrieved before being displayed with a consequent longer display time. The text

color is green.

- OFFLINE. This means that the item is present only in backup media and can only be

retrieved by physically locating the media and inserting it into the review station.

The text color is brown.

- UNAVAILABLE. This means that the item files are either corrupt or have been removed from the

system before backup. These items can no longer be retrieved. The text color is

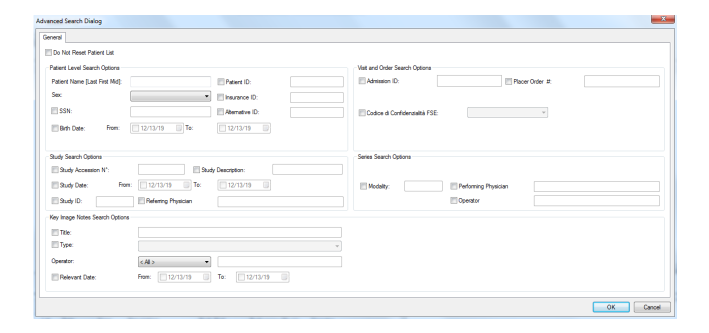
red.

8.1.3 ADVANCED QUERY

This module allows performing complex queries by posing constraints on patient, study, and series data.

In the Archive screen, by clicking on the Advanced Search button, you can further specify the parameters for the query.

As you can see in the following figure there are many parameters that allow you to make a more organized and optimized query.



Please note that the information inserted in the fields may be incomplete; it is enough to enter just the first characters. Wildcard characters (*?) are allowed.

The availability of simple or advanced queries depends on the features of the server. Whereas it is fully functional with the *ComPACS Server* you may experience some limitations with third party DICOM servers.

8.1.4 RETRIEVAL AND DISPLAY

To retrieve a selected item from the archive, click the *Open* button. The images will be automatically displayed when retrieved. The software automatically detects if the information is already present in the local archive or if it is necessary to open a communication channel with the remote server to retrieve the necessary information

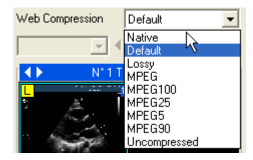
The system behavior when opening a patient has changed. Instead of opening all studies belonging to the patient, only the most recent study is opened. If you would like to open one or more studies, please first select the patient, then select the studies to be opened and finally click the *Open* button.

If you want to compare different studies or patient you can click on small arrow to the side of the *Open* button and select the related function.



8.1.5 WEB COMPRESSION

When retrieving images from a ComPACS Web Server it is possible to adjust the compression to optimize the download time in relation to the available bandwidth and requested image quality. A drop down menu allows the selection of the compression level applied.



The items in the list are those pre-configured by the system administrator. The *Native* value corresponds to the files as they were stored to the server, whereas *Default* corresponds to a default compression also pre-configured by the system administrator.

The selected compression level in the drop down list must be chosen before the images are retrieved from the archive window. To change the compression level of an already opened study you must first close it, change the compression level, and then re-open it.

8.2. ASSIGNING STUDIES

It is possible to Assign or Release a study by using the respective buttons after having selected a specific study.

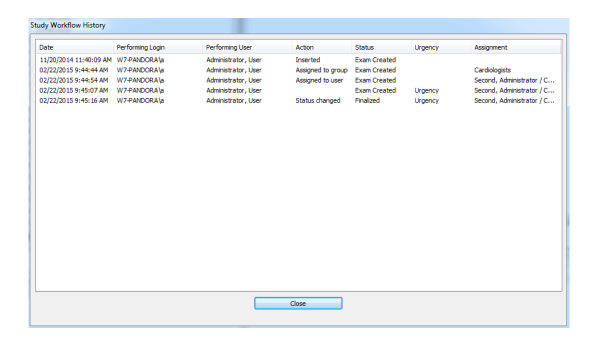


If you assign it to a different user or group, a dialog will prompt you for user selection. The assignment information will be immediately updated in the study panel.

Additionally when a study is assigned to a user identified as belonging to a *Technician* or *Physician* type, the report status will automatically progress during the different reporting phases.

The user and group assignments are independent. A study may be assigned both to a group and to a user who belongs to a different group.

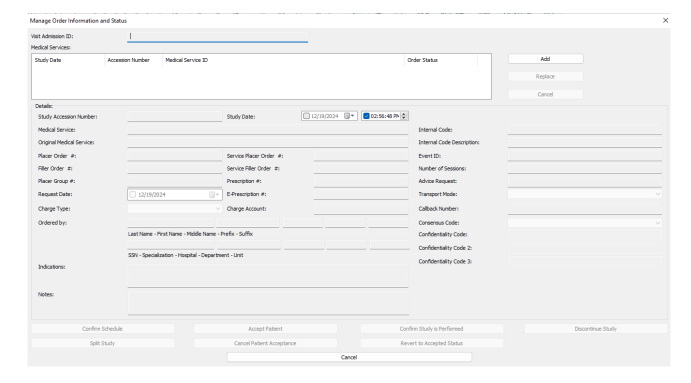
Clicking the button will display a window with the history of the study assignment and status changes. The window also displays the date and time in which the change occurred and the user performing the operation.



8.3. ORDER MANAGEMENT

You can manage the order for the selected study or modify the status by pressing the button. A dialog window will allow such operations.

Manage Order



In this window it is possible to manage the order status: confirm the schedule, accept the patient, confirm that the study has been performed, discontinue the study, split the study or revert previous operations.

In this window it is also possible to add, replace or remove "Medical Services" associated to the study. When adding one or more medical services, a new dialog window will be displayed, optionally allowing the indication of a prescription number to associate to the new service added or the generation of an electronic prescription.

You can also manually for a study status change by clicking on the status field in the list. A dialog window will allow such operations.

Unless your user account belongs to a profile with "Admin Management" rights, you may be subject to certain restrictions on the basis of the system configuration. It is possible that some stallates can be set only automatically by the system (for example using DICOM MPPS) and not manually by the user. Also it is possible that some states, once reached, cannot be manually modified. The "Change" button will be grayed out in such circumstances to prevent the change.

The above status change is also performed automatically by the ComPACS Worklist and/or HL7 Server when such components are installed.

Additionally when a study is assigned to a user identified belonging to a *Technician* or *Physician* type the report status will automatically progress during the different reporting phases.

The generation of an electronic prescription is subject to the interfacing of a suitable prescription module.

The status information will be displayed in the study panel list. Moreover at the patient level a set of symbols to the left of the patient name will warn the user if some reports are pending activity for the specified patient.

The possible icons with descending priority are the following:

- No icon. This means that at least one study for the patient is still Not Read.
- Tech Sheet Started. This means that at least one study for the patient is in the preliminary tech sheet reporting phase.
- - Tech Sheet Completed. This means that at least one study for the patient has a completed technician sheet which is pending reporting or report verification.
- Report Verified. This means that at least one study for the patient has a verified report which is pending finalization.
- Report Finalized. This means that all reports for the study have been reported and finalized.

8.4. PATIENT AND STUDY URGENCY STATUS

You can modify the urgency status of a study by clicking on the urgency field in the list. A dialog window will allow the new urgency status selection.



Unless your user account belongs to a profile with "Admin Management" rights, you may be subject to certain restrictions on the basis of the system configuration. It is possible that some states can be set only automatically by the system (for example using HL7) and not manually by the user. Also it is possible that some states, once reached, cannot be manually modified. The "Change" button will be grayed out in such circumstances to prevent the change.

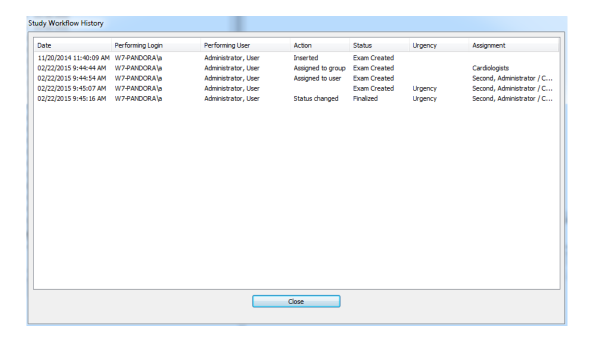
The above status change can be performed automatically by the HL7 Server when such components are installed.

The status information will be displayed in the study panel list. Moreover at the patient level a set of symbols to the left of the patient name will warn the user about the urgency of its related studies.

The possible icons with descending priority are the following:

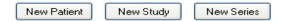
- - Emergency. This means that at least one study for the patient has an emergency status.
- - *Urgency*. This means that at least one study for the patient has an urgency status.
- - Low Urgency. This means that at least one study for the patient has a low urgency status.
- - No Urgency. This means that at least one study for the patient has a no urgency status.
 - No icon. This means that the urgency for all the patient studies has not been defined.

Clicking the History button will display a window with the history of the study urgency status changes. The window also displays the date and time in which the change occurred and the user performing the operation.



8.5. ADDING OR EDITING NEW ITEMS

To add a new item it is sufficient to click the New Patient, New Study or New Series buttons and enter the content in the displayed dialog windows:

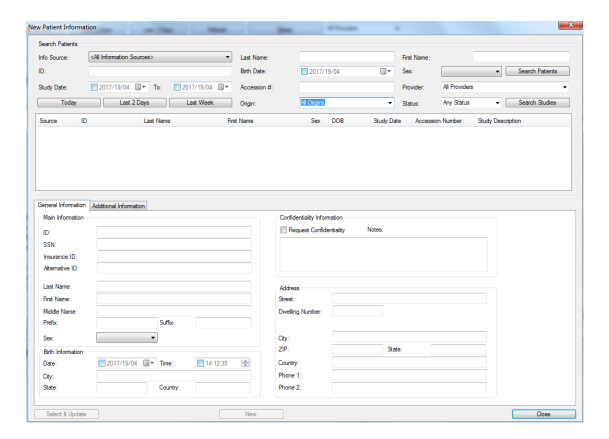


Alternatively a selected item can be edited by clicking the corresponding buttons:



8.5.1 ADDING OR EDITING PATIENTS

The information about the patient to be added is displayed in the following page:



In the top portion of the dialog window it is possible to import the patient from the available databases. By default the system will search all the available information sources but a specific one can also be selected using the drop down menu. The "Default Demographics Provider" is always present and it searches the ComPACS patient database. Other external sources may also be present.

The Search Patients button searches the database patient demographics and the Search Studies button search the study worklist.

The bottom portion of the dialog window displays the selected patient information. Such information may also be manually inserted if the patient cannot be found in the information sources.

The Select & Update button confirms the selection of the identified patient and updates the ComPACS database information in case it differs from the selected information source. The New button instead adds a new patient to the ComPACS database. Both buttons will complete the operations and the dialog will close.

If a study search has been performed, a new dialog to add a study to the patient will be opened. This dialog will be populated with the data imported from the information source.

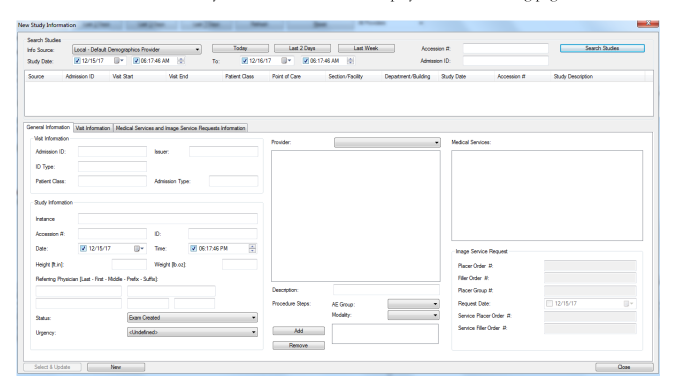
The window displayed for editing is very similar. In place of the *Select & Update* and *New* buttons, the *Reset*, *Update* and *Merge* buttons can be found.



These buttons respectively reset the patient information to the original values, update the database with the new inserted information or merge the patient with the patient selected from the list.

8.5.2 ADDING OR EDITING STUDIES

The information about the study to be added or edited is displayed in the following page:



In the top portion of the dialog window it is possible to import the study from the available databases. By default the system will search all the available information sources but a specific one can also be selected using the drop down menu. The "Default Demographics Provider" is always present and it searches the ComPACS study database for the selected patient. Other external sources may also be present.

The Search Studies button searches the database study worklist.

The bottom portion of the dialog window displays the selected study information. Such information may also be manually inserted if the study cannot be found in the information sources.

In the left portion of the lower panel, it is possible to view and edit the study information and, if the study is associated to a visit, the related visit information.

In the middle portion of the panel, it is possible to select the diagnostic provider and, accordingly, a pre-built list of study descriptions for the selected provider can be found for rapid selection. These descriptions are system wide settings to be configured by your system administrator.

Below the study descriptions you can find information about the procedure steps related to the selected study. For each procedure step it is possible to define the *Modality* and the group of Application Entities (AE Group) associated to the procedure.

In the right portion of panel it is possible to identify the medical services associated to the selected description and optionally set related information such as placer order number, placer group number or prescription number. If the common placer order number or placer group number are defined, an image service request order will automatically be generated in relation to such services.

The Visit Information tab allows the setting of more detailed visit information. The Medical Service and Image Service Requests Information tab allows the setting of more detailed order and medical service information.

The Select & Update button confirms the selection of the identified study and updates the ComPACS database information in case it differs from the selected information source. The New button instead adds a new study to the ComPACS database. Both buttons will complete the operations and the dialog will close.

The window displayed for editing is very similar. In place of the *Select & Update* and *New* buttons, the *Reset*, *Update* and *Merge* buttons can be found.

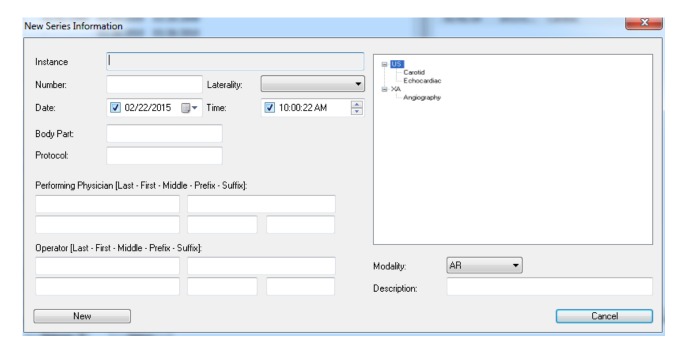


These buttons respectively reset the study information to the original values, update the database with the new inserted information or merge the study with the study selected from the list.

It is possible to modify the medical service information only when the study is first created. Once created, medical services need to be modified using the study acceptance window.

8.5.3 ADDING OR EDITING SERIES

The information about the series to be added is displayed in the following page:



In the panel on the right, a pre-built list of series descriptions for each modality can be found for rapid selection. These descriptions are system wide settings to be configured by your system administrator.

Click the *New* button to confirm the operation. The new series will be added.

The window displayed for editing is very similar. In place of the *New* button an *Update* button is present to confirm the new information. The confirmed changes will immediately affect the series and all its contained objects.

8.6. MERGING AND MOVING ITEMS

Merging and moving Objects, Series, Studies, or Patients can be easily and intuitively performed using mouse based drag and drop actions.

These operations carry out critical changes to the information in the ComPACS database and they are not reversible. To avoid their inadvertent activation, they must be first enabled by clicking the following button:



8.6.1 MERGING ITEMS

To merge patients, studies or series, simply select one or more such items in their related result window, drag them over the item in the same window to which they should be merged and them drop

the selection. A dialog box will warn you about the importance and non-reversibility of the operation which you will need to accept in order to proceed.

At the end of the requested operations the user is advised to de-select the *Enable Merge/Move* button to inhibit other operations.

8.6.2 MOVING ITEMS

To move studies, series, or objects simply select one or more such items in their related result window, drag them over the parent item to which they should be moved and them drop the selection. A dialog box will warn you about the importance and non-reversibility of the operation which you will need to accept in order to proceed.

For parent items, only the direct parent is intended. An object has a series as its parent, a series has a study, and a study has a patient.

At the end of the requested operations the user is advised to de-select the *Enable Merge/Move* button to inhibit other operations.

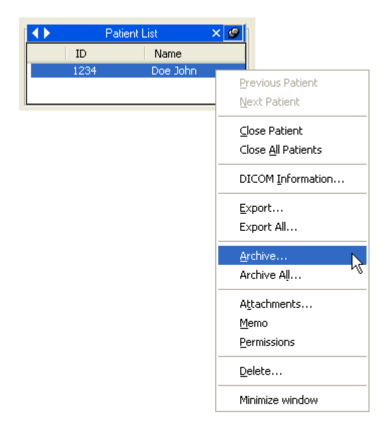
8.6.3 SPLITTING ITEMS

The splitting of the content of an item can be performed with new and move operations. For example to split the content of a study you would proceed as follows:

- Create a new study.
- Create new series of the same type of the ones already contained in the study to be split. For example if a study contains a US series, create a new additional US series.
- Select the objects that need to be separated into a different study and first move them to the new created series within the same study.
- Now complete the operation by moving the new series to the new study.

8.7. ARCHIVING PATIENT IMAGES AND LOOPS

To archive the images displayed or opened for review you can simply right-click on the image thumbnail, series, study, or patient in the *service window* displays and select the archive function from the pop-up menu.

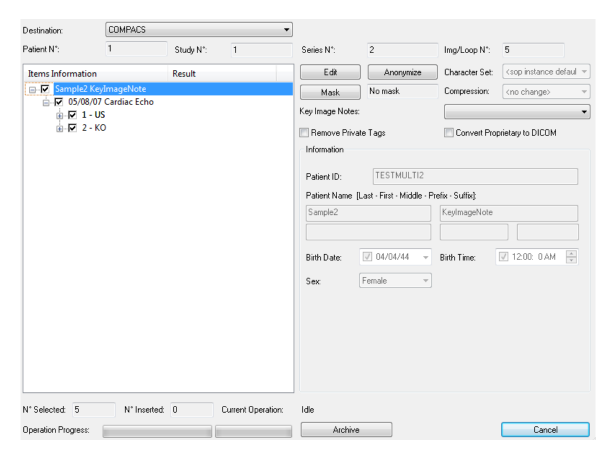


A dialog window with the list of selected image objects will be displayed. On top of the window you can choose the destination archive and see a brief summary on the items selected.

It is now possible to refine the selection by clicking on each check box to select/deselect the image or loop. You can also select/deselect a patient, a study, a series and an image.

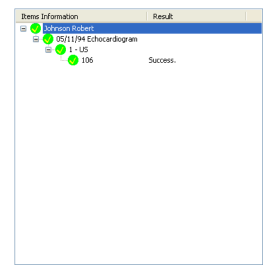
If the patient has associated Key Image Notes, it is also possible to limit the items exported to only the items referenced in the Key Image Note. The "Key Image Notes:" drop down window will display the available Key Image Notes.

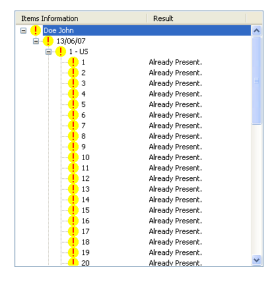
While navigating in this object hierarchy, it is possible to examine the related DICOM information in the panels on the right hand side of the window.



If you click the *Archive* button the selected images will be transferred to the archive. If a network archive has been selected the transfer will take longer. The slider bars indicate the progress of the transfer to the archive.

When the transfer is completed successfully a green icon will be displayed. A yellow icon indicates a warning, whereas a red icon indicates a failure. Please look at the result section at all levels for more information. In the example below to the left the archive was successful whereas on the one to the right a warning is returned because the patient was already present in the archive.



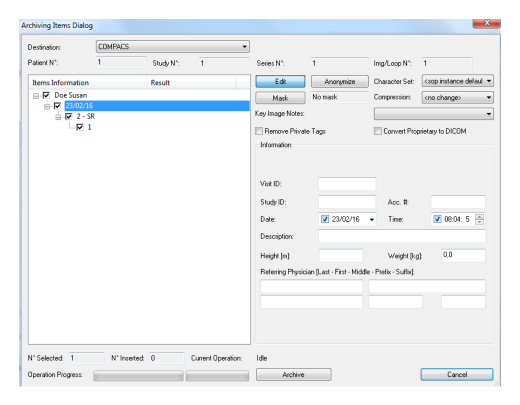


You can also archive all the data present in the control window. To do this, right-click on the *Patient List* of the control window and then select *Archive All*.

At the object level, the "already present" warning does not prevent the storage operation from occurring. In this scenario the already present object file is overwritten with the new object stored.

8.7.1 EDITING INFORMATION

During the archive process it is also possible to edit some of the DICOM information or even anonymize the items selected. To perform this operation, select an item and click the *Edit* or *Anonymize* button. The information panels will now be editable for you to set to the desired values.



During this operation it is possible to change the character set used or change the object/image compression. Changing the character set can be performed for the entire patient set (including all subseries) whereas changes in compression can be performed for different individual images.

It is also possible to remove any DICOM private tag from the selected objects or convert them to a standard DICOM format (for example Aloka Line Data).

By clicking again on the *Edit* button you can revert back to the original data values.

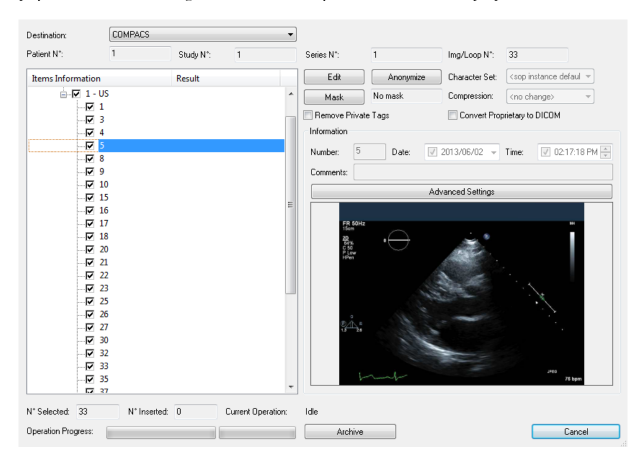
When re-archiving objects while editing the patient information, please ensure the same Visit Admission ID is not used across different patients. This identifier cannot be automatically coerced and the use of the same identifier in different patients would cause a conflict and a consequent storage failure.

8.7.2 ANONYMIZING IMAGES

In addition to data, ComPACS also allows for the anonymization of DICOM images and loops. The technique is identical to the one described in this guide for what concerns *Masks* usage. You can allow the system to automatically detect the masks or manually select it for each image or loop.

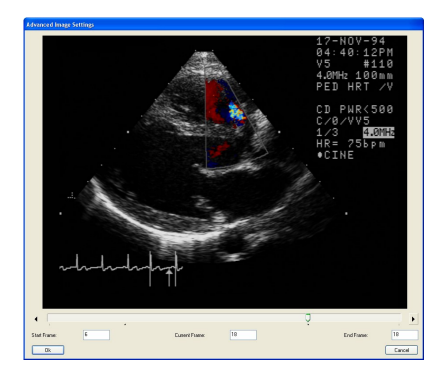
8.7.3 TRIMMING IMAGES

The archive dialog can be resized (by dragging the bottom right corner with the mouse). This causes the resize of the image and loop information page to better display the preview of the selected item. If the item is a loop, it will be displayed as playing at a fixed frame rate of 25 frames per second. This display is not intended for diagnostic review but only for item identification purposes.



ComPACS also allows you to trim the image loops during storage. To perform this operation you need to select a specific loop and then click the *Advanced Settings* button.

A new window will appear in which it is possible to review the loop and select the trimming points.



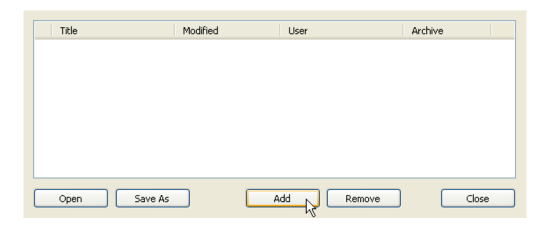
8.8. ATTACHMENTS

ComPACS Review Station, in addition to DICOM data, allows the management of non-DICOM information such as Adobe® PDF and Microsoft® Word documents as attachments.

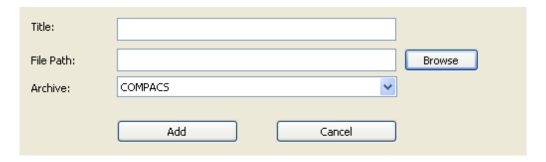
Attachments can be related to each level of the Patient/Study/Series/Object hierarchy. To attach, display or remove an attachment you simply need to *right-click* on an image thumbnail, a series, a study, or a patient name and then click on *Attachments*.



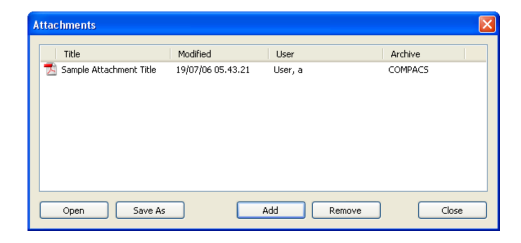
A new window is displayed in which all attachments for the selected item are listed.



Clicking the *Add* button allows you to add a new attachment.



After you have given the attachment a title (and selected the appropriate archive if more than one is configured) you can browse your computer and then attach any kind of file. Clicking on *Add* will effectively attach the selected file. If the file is associated to an icon, the related icon will be displayed to the left of the selected title as shown below.

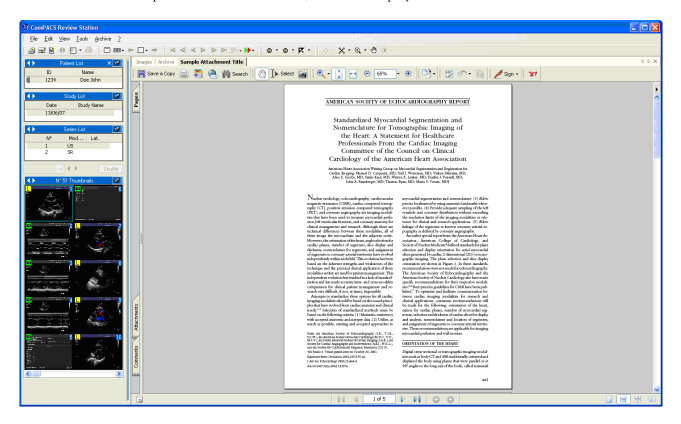


The attachment list also identifies the user that stored the attachment, the archive in which it has been stored and the last modification date.

It is also possible to manage the attachments by selecting them and then clicking on the button related to the requested operation: Open – to open the attachment within ComPACS, $Save\ As$ – to save it to an external file, Remove – to delete the attachment.

Please note that the operations allowed on attachments depend on your privileges to the selected item (Patient, Study, Series, or level). Therefore some or all of the operations may be grayed out because they may not be accessible.

If the attachment is opened within ComPACS, it will be displayed as a new tab window.



For the document embedding function to work, it is necessary that the selected document and related viewing application comply with ActiveX and Active Document technology. Examples are Adobe® PDF and Microsoft® Office files.

If an item contains one or more attachments, the user will be warned by a *paper clip* symbol that will be visible to the left of the related item as shown in the picture below. The same information is also displayed in the query window.



8.9. **MEMO**

A memo is a way to attach an important note to an item in the Patient/Study/Series/Object hierarchy.

To access the memo options, right-click on an image thumbnail, a series list, a study or on a patient and then select *Memo*.



The operation will open a *memo* window.



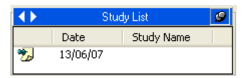
The window, in addition to the main memo text, will also show the *Archive* and the item's main information (for example patient name or study date) to which the *Memo* belongs, the user that created or last modified the memo and the modification date.

To edit the memo text simply type directly into the memo box. By clicking the X in the upper right corner you will confirm your selection and close the window.

To *Remove* the memo you can clear the text and close the window. Alternatively, this and other functions are available by clicking on the small *memo* icon on the top-left of the window accesses.

Please note that the operations allowed on memos depends on your privileges on the selected item (Patient, Study, Series or level). Therefore some or all of the operations may be grayed out because they are not accessible.

If an item contains a *Memo*, the user will be warned by a special symbol visible to the left of the related item as shown in the picture below. The same information is also displayed in the query window.

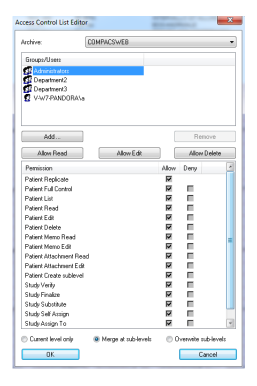


8.10. PERMISSIONS

ComPACS allows advanced and granular permissions to be associated to each item (Patients, Studies, Series, Objects, or even Attachments and Memos). To open the permission menu, right-click on an image thumbnail, series, study, or on patient name and then select *Permissions*.



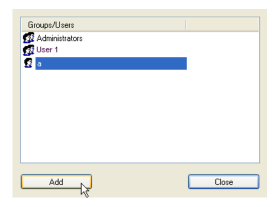
Below is an example of the (ACL) Access Control List Editor window that will be displayed.



In the window you can select a user (or a group of users) and allow or deny operations such as reading, editing, deleting, etc. Three quick settings buttons (*Allow Read*, *Allow Edit* and *Allow Delete*) assist you in the most frequently modified security settings.

The names of such operations refer to the item selected and directly convey the types of permission being managed. However please refer to your system administrator for more details.

It is also possible to add or remove users and groups from the list by clicking the *Add* or *Remove* buttons. When the *Add* button is selected, a related window will appear to allow the user or group selection.



Please note that the users and groups that can be added are only those registered for the ComPACS application. Please refer to your system administrator to modify this list.

Upon confirmation, it is also necessary to specify how the changes will be applied to the ComPACS database. Three different options are available:

Current Level Only — Modifies the permissions only for the selected level.

Merge at Sub-Levels — Modifies the permissions for the selected level and then merges the changes to all the sublevels (for example the sublevels of a study are its series and objects). The merge operation will preserve any additional and specific permission settings at the lower levels if not affected by the changes made (for example if it refers to different users or groups).

Overwrite Sub-Levels — Modifies the permissions for the selected level and then applies it to all the sublevels (for example the sublevels of a study are its series and objects), overwriting and prior permission settings that may have been set for such levels.

It is important to note that to set the permissions parameters, you must have Full Control Permission on the related object.

These permissions will refer to the item level (Patient, Study, Series or Object) you have selected but will necessarily be propagated to the upper levels to maintain consistency in the permission settings. For example setting a *Read* permission at the *Study* level requires at minimum a *List* permission at the *Patient* level.

Changing the permission settings will affect the possibility users will have to list, read, modify and delete the different objects in the ComPACS database. Please ensure to use this feature with great care.

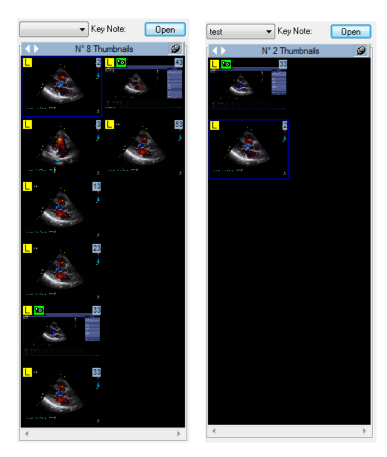
9. KEY IMAGE NOTES

9.1. REVIEWING A KEY IMAGE NOTE

If the Key Image Note module is licensed, a set of controls will be displayed in the control window:

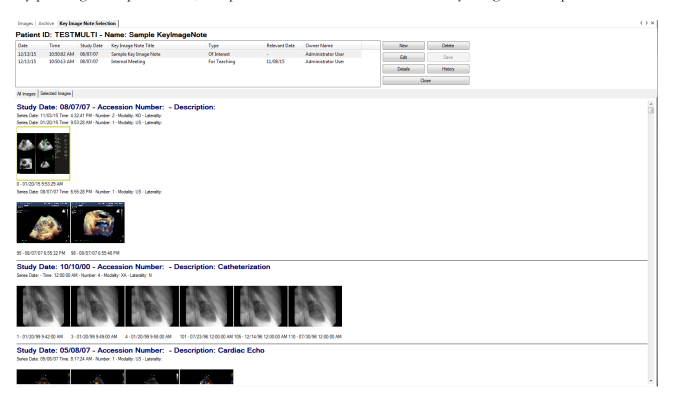


The drop down will list all Key Image Notes available. If selected, the thumbnails display will reflect the images referenced in the Key Image Note, even if belonging to different studies. All thumbnail navigation will now be within such Key Image Note set.



The user can always revert to the complete set of thumbnails, by removing the Key Image Note selection.

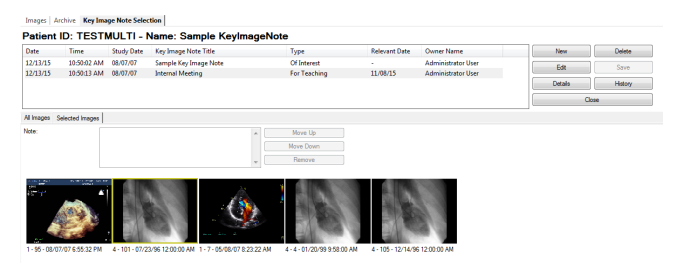
By pressing the Open button, it is possible to review and edit all the Key Image Notes opened:



At the top of the window, a list of Key Image Notes is displayed. If the title displays a "*" symbol at the end, the Key Image Note is opened for editing.

For the selected Key Image Note, the referenced images in the opened studies are identified by a yellow rectangle.

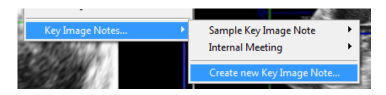
The "Selected Images Tab" displays only the images referenced in the selected Key Image Note.



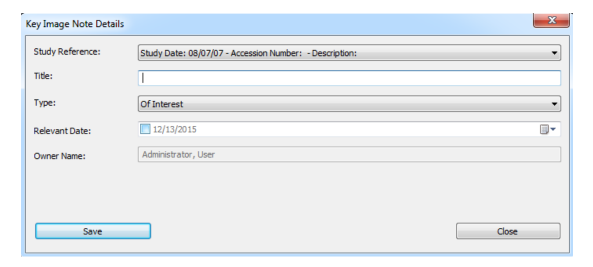
9.2. CREATING A NEW KEY IMAGE NOTE

It is possible to create a new Key Image Note in two different ways:

- By pressing the button in the Key Image Note Selection window.
- By right clicking on an image and selected the context menu item:



A dialog window will be displayed:



It is necessary to associate a *Title* and *Type* to the Key Image Note. If more studies are available for a patient it is also necessary to indicate the study in which this note will be stored.

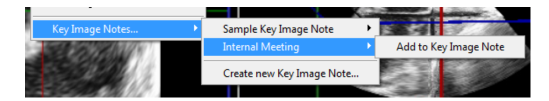
Optionally it is also possible to indicate a Relevant Date, useful to search for such note later.

9.3. EDITING A KEY IMAGE NOTE

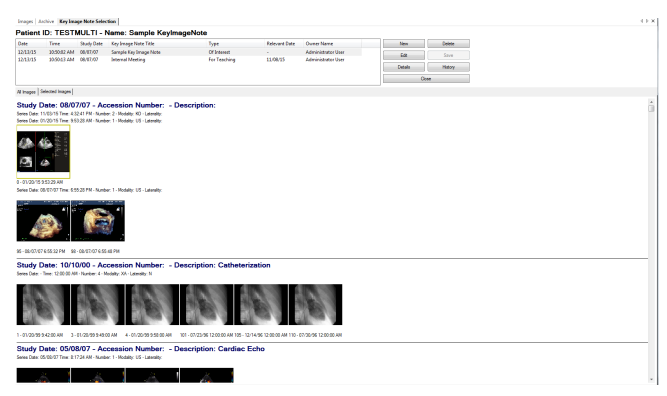
If the Key Image Note is not in editing mode, it is necessary to press the enter such mode. Only one user can edit the note at the same time.

Once all changes have been made, press the button to confirm the changes or the button to cancel the operation.

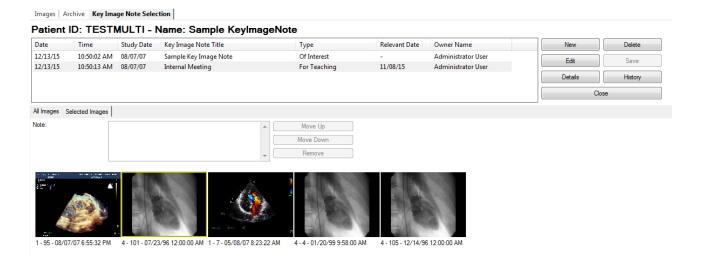
Images can be added or removed from the Key Image Note by right clicking on the related image and selected the corresponding context menu item:



Alternatively, it is possible to add or remove images by simply clicking on the related thumbnail in the Key Image Note Selection window. A yellow rectangle indicates if a specific image is referenced.



In the "Selected Images Tab" it is possible to associate notes to a specific image or change the display order, either by drag & drop, or using the *Move Up* and *Move Down* buttons. Images can also be removed using the *Remove* button.





To modify the content of the Key Image Note, it needs to be opened in editing mode.

9.4. OTHER OPERATIONS

The button opens a dialog displaying the history of the changes. In this dialog it is also possible to force the loading of a previous revision of the Key Image Note.

The button opens a dialog displaying the main Key Image Note information. In this dialog it is also possible to review the entire DICOM tag content or to modify the permissions to access the Key Image Note.

The button removes the Key Image Note from the archive. This operation will attempt to remove the entire Key Image Note chain, including prior revisions.

9.5. OPENING A KEY IMAGE NOTE

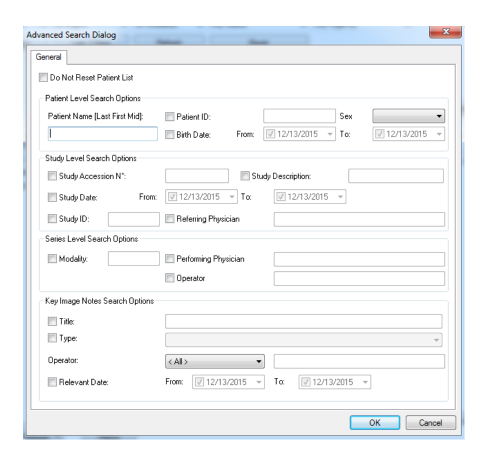
A Key Image Note can be opened and selected directly from the archive by right clicking on the Patient or Study Window:



A dialog window will list all the Key Image Notes associated to the selected item.

9.6. SEARCHING FOR A KEY IMAGE NOTE

It is possible to search the archive for specific Key Image Notes by using the Advanced Search dialog:



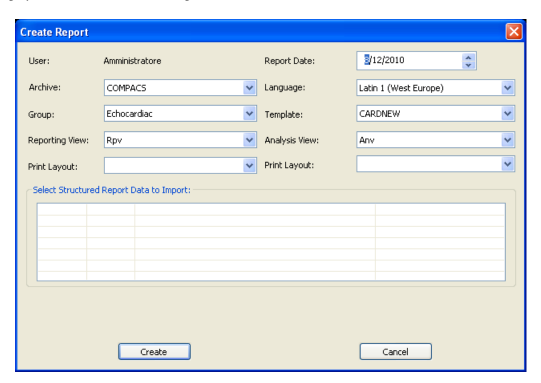
10. MANAGING EVIDENCE AND REPORTS

10.1. CREATING A NEW REPORT

To be able to create or open a report, you must first adequately configure your system. Please refer to your system administrator and to the *System Administrator's manuals* for more details.

In order to create a new report, a study first needs to be opened for review.

Then click the open/create report button on the toolbar 2 = 2 = 0. A dialog window will prompt you to enter a number of parameters.



Let us examine these parameters in detail:

Report Date — this is the date for the new report. It defaults to the current system date.

Archive — this is the archive in which the created report will be stored.

Language — this is the language used in the report. It defaults to the configured language for the client application entity to which your review station corresponds.

It is important to choose the correct language. Otherwise some special characters may not be stored correctly into the report.

Group

– report types are grouped to allow you to better identify and select the type required. This field allows you to select the group of report types you are interested in, for example *Echo Vascular* reports.

Template

– this field identifies the report type (or template) that you effectively need to create. For example a *Vascular Report*.

Reporting View

– the information stored in a report of a certain type (identified by its template) can be displayed in different ways (Views), if present, depending and optimized for reporting purposes based on the body part, user preferences, patient pathology, trial protocol, etc. A possible example is the *Carotid View* for a *Vascular Report*.

This field allows the selection of the default view for each report. Please note that this is just an initial selection that can then be changed dynamically during the processing of the report.

Print Layout

- under the *Reporting View* field it is possible to select the *Print* Layout. With this parameter you can define the final print layout of the information stored in a report of a certain type (identified by its template and reporting view).

This field allows the selection of the default print layout for each report. Please note that this is just an initial selection that can then be changed dynamically during the reporting process.

Analysis View

– the information stored in a report of a certain type (identified by its template) can be displayed in different ways (Views), if present, depending and optimized for image analysis purposes based on the body part, user preferences, patient pathology, trial protocol, etc. A possible example is the *Carotid View* for a *Vascular Report*.

This field allows the selection of the default view for each report. Please note that this is just an initial selection that can then be changed dynamically during the analysis process.

Print Layout

– under the *Analysis View* field it is possible to select the *Print* Layout. With this parameter you can define the final print layout of the information stored in an analysis report of a certain type (identified by its template and analysis view).

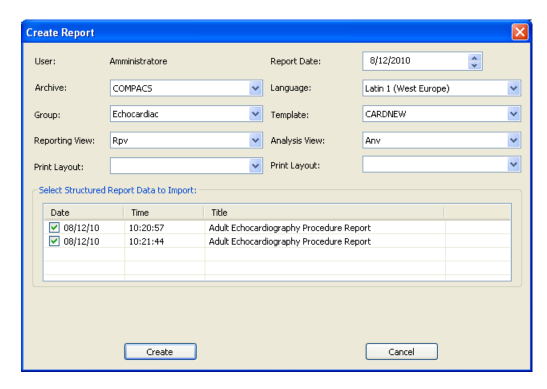
This field allows the selection of the default print layout for each analysis report. Please note that this is just an initial selection that can then be changed dynamically during the analysis process.

Once you have selected the right parameters you can create the report by clicking the *Create* button. Please note that this operation may take some time depending on the size and type of report being created.

The system stores the last settings used by each user in report creation. Therefore if you always create the same type of report it will be sufficient to directly click *Create*.

10.2. IMPORTING EXISTING REPORTS

If existing reports that have not yet been associated are detected, such reports will be listed in the dialog window. If checked, the content of these reports will be included and merged in the newly created report.



This is the typical scenario when a preliminary report has been generated from the modality and stored to the system. In this case the report window parameters will be automatically selected on the basis of the detected preliminary report type and configuration.

If only a single report is present and if ComPACS is capable of automatically and univocally assigning the necessary parameters, the report window dialog may not be displayed and the report object may be automatically opened for editing.

Such dialogs will always be displayed if more than one report needs to be merged to warn the user about such a condition. This operation should be performed with great care to make sure the data in the different reports does not conflict.

10.3. EDITING A REPORT

In order to open an existing report for editing, the report object needs to be opened and visible in the Report List control window, on the Data tab.



The operation is identical to the one performed to create a new report. Click the open/create report button on the toolbar on the toolbar.

If only a single, non-obsolete, report is present then the object will be automatically opened for editing.

If additional, non-obsolete, structured report objects have been stored to the system, the report window dialog described in the previous sections will be displayed. This will both warn the user about the existence of such additional reports, and it will also allow such reports to be imported and merged with the current report.

This typically occurs if the report is initialized prior to image and measurement acquisition. In this situation the measurements structured data will have to be imported when the initialized report is completed.

Importing and merging reports should be performed with great care to make sure the data in the different reports does not conflict.

10.4. OPENING REPORT DOCUMENTS

To be able to create or open a report it is first necessary to adequately configure your system. Please refer to your system administrator and to the *System Administrator's manuals* for more details.

In order to open a specific report document, a study containing a report needs to be opened for review. The available reports for the selected study are listed in the *Report List* control window, on the *Documents* tab:



Finalized reports and generally documents are stored as PDF files in DICOM Encapsulated PDF format. Such items are visible in the *Documents* tab and can be opened by a simple mouse click.

A DICOM Encapsulated PDF Report object may also be present if created by a third party system and then stored in the ComPACS Database.

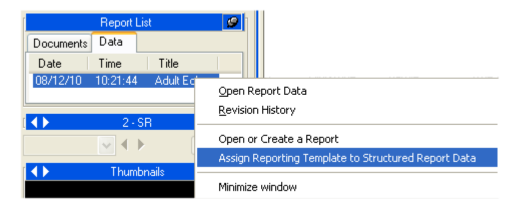
To display the PDF report it is necessary for the *Encapsulated Report Module* to be installed and licensed. Furthermore, Adobe® Acrobat PDF Reader needs also to be installed.

The report structured information is generally stored in a DICOM SR object and visible in the *Data tab*. These items can be opened by a simple mouse click and reviewed in depth with the specific analysis or reporting modules licensed. This is equivalent to right clicking on the report item and selecting *Open Report Data*.

If the report was not generated by a *ComPACS Review Station* or *ComPACS measurement import plugin*, but was received directly from a third party template, then upon clicking the report object, you will be prompted with a dialog window that will require the user to select the template and view information.

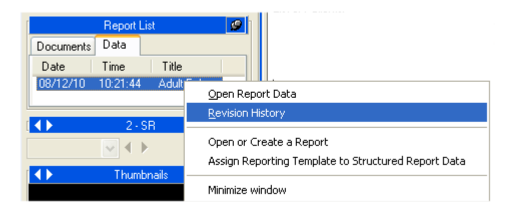
This operation is similar to report creation shown in the previous section but it does not lead to the generation of a new report, only to the association of the correct template and views.

It is also possible to manually associate a report to a different template. This can be performed by right clicking on the report item and selecting Assign Reporting Template to Structured Report Data.

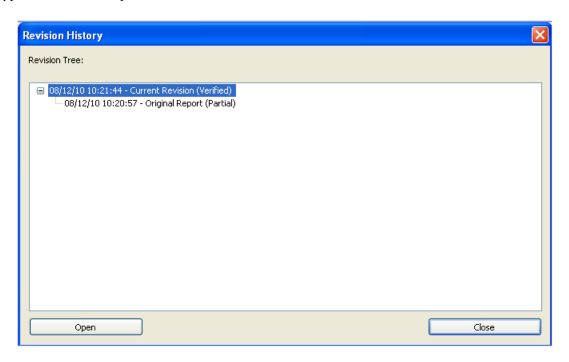


10.5. REPORT REVISION HISTORY

ComPACS keep track of report revisions. The *Report List* tabs will show only the most current revision of the documents and structured data opened for review. To examine any previous documents, right click the item and select *Revision History*.



A window will display in a hierarchical tree the revision history for the selected item. The report date, time, type and status is reported for each revision.



It is possible to open an obsolete document revision by selecting it and clicking on the *Open* button. These documents will be available for review but cannot be edited.

The revision history is limited to the object opened for review from the ComPACS archive. If only a limited set of objects is opened on the workstation, the items in the revision history shall be incomplete and may lead to an incorrect interpretation of the current report status.

Please ensure that all documents are always opened or that at least the most current report revision is always present in the opened list.

11. ECHO ANALYSIS

11.1. ACTIVATING OFF-LINE ECHO ANALYSIS

If ComPACS ECHO is enabled on your system, the following button will be visible in the toolbar, on the right hand side:



If the button is pressed down then the module is already active. Otherwise it can be activated by clicking the button with the mouse.

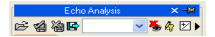
Upon activation, the module will request a runtime license from the ComPACS Server. If the license cannot be received, then the module will fail to activate and an error message will be displayed.

In case of concurrent licenses it is possible that all available licenses have already been requested and assigned to other users that are using the module at that time and that no licenses are left for activation. This is a likely cause for failure.

Activation failure can also occur if the network connection to the server is no longer available.

Please ensure that the module is released, by releasing the above button, after you have finished using the product so that the license is available for use by other users.

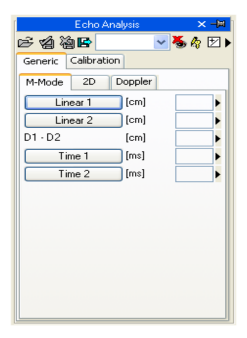
When the module is active, a window named "Echo Analysis" is added and visible in the control window. It can be opened or minimized with a single mouse left button click on the blue bar.



11.2. OFF-LINE MEASUREMENTS

The ECHO module provides the possibility of performing measurements on every type of echo image.

Simple measurements are always available in the Generic tab of the echo analysis window:



These are basic and generic M-Mode, 2D, and Doppler measurements for rapid verification purposes.

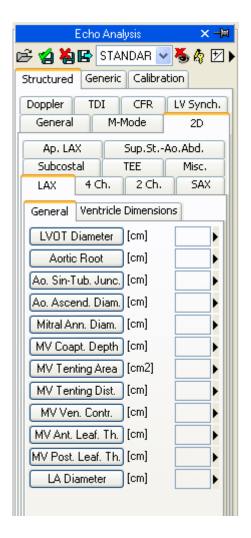
Generic measurements are only temporary. They are not stored to the ComPACS database and cannot be retained.

To access a more advanced measurement panel, open either an already existing report or create a new one. This operation can be performed by clicking on the toolbar icon as described in the "Managing Evidence and Report" chapter.

The analysis window will be extended with an additional *Structured* tab containing all the structured measurements. The appearance and content of the tab will depend on the reporting templates and analysis views configured by the system administrator for the specific workstation or user.



Structured measurements are stored in the DICOM SR object they refer to.



In this interface, only the windows corresponding to the data present in the open object will be filled (with numbers). To edit the opened report click the *Editing* icon:



To close and save the changes made click on the Close and Save changes icon:



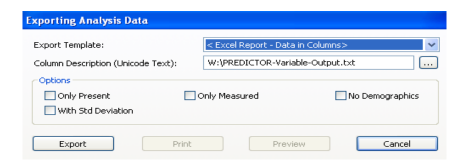
To close without saving any changes click on the Close discarding changes icon:



To export the data it is necessary to click the Export the report data to file icon:



An interface will be opened where you may select the data format and the file destination.



In the drop-down list, it is also possible to select an export template based on the installed configurations.

This functionality has been extended to export not only numerical values but also other field types (text, findings, dates, etc.). Furthermore the export also includes nested and repeated record structures (for example each phase of a echo stress study).

The template is a simple text file which lists, in separate lines, the fields which need to be exported. Each line has the following format:

FieldID; Label - Optional; Unit - Optional

If the label or the unit are not specified, the default values shall be used.

For repeated records, the list of fields need to be enclosed in the following statement:

BEGINGROUP(Record Field ID); Label – Optional; Index – Optional ... list of fields ... ENDGROUP(Record Field ID)

The index is optional and, if omitted, all records shall be exported.

Finally some special fields consist in DICOM fields. They are identified as follows:

\$(Group, Element); Label - Optional;



Editing operations will be described in the next chapters.

11.3. IMPORTED MEASUREMENTS

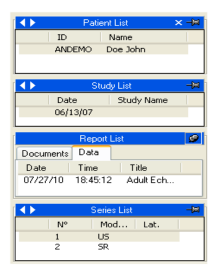
The ECHO module allows importing measurements taken from the ultrasound devices through DICOM SR. Therefore, the product offers the possibility to view the data measured on the platform as well as to insert new measurements, or correct the existing ones.

For the most common ultrasound platforms that do not comply to the DICOM SR standard, some ad hoc techniques are also available to import the measurements. The ComPACS architecture allows importing such measurements and transforming them into the DICOM SR format to assure maximum portability.

When a study is opened for review and a DICOM SR object is present, the *Report List* will display such element on the *Data* tab.

The *Data* tab initially represents the measurements performed on the modality, if any. However please note that the data object will also be revised to include off-line measurements or other reporting activities.

In the *Series List*, an item line with SR as the modality will also appear. This is the DICOM series containing the DICOM SR object elements.



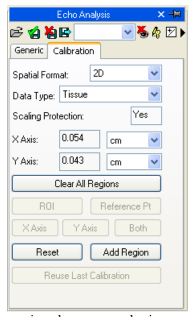
By clicking on the toolbar icon as described in the "Managing Evidence and Report" chapter, you can open existing reports for review and further analysis. This includes imported modality data.

Alternatively it is possible to specifically open an DICOM SR document and access the contained data by clicking on the selected SR item on the *Data* tab.

11.3.1 IMAGE CALIBRATION

The calibration information is usually contained in the DICOM file. The product exploits it to avoid a manual calibration and ensure greater accuracy. Therefore calibrating images in order to perform measurements is generally not necessary nor is it recommended.

By simply moving the mouse over the image, the calibration information will appear in the *Calibration* tab. This way it is possible to quickly verify if the image calibration is present or not.

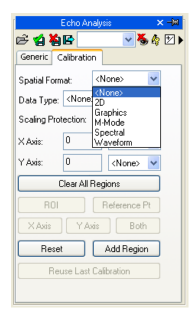


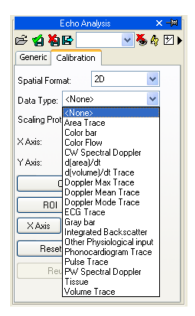
If no information is displayed when moving the mouse, the image needs to be calibrated.

Ultrasound images can contain multiple regions of calibration. Therefore it is possible that the calibration information differs in relation to the position of the mouse on the image. Also some regions may be calibrated whereas in others the calibration information may be absent.

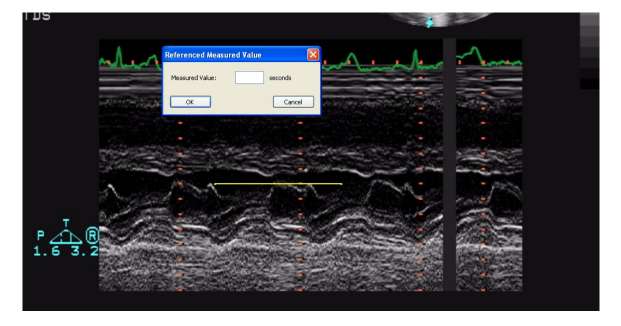
If the calibration information is absent, it is possible to manually calibrate the entire image or a region.

The first thing to do to calibrate an image is to choose the Spatial Format (2D, Doppler, M-Mode, etc...) and the type of data (Tissue, PW, CW, etc...). Using the two drop down menus in the *Calibration* tab:





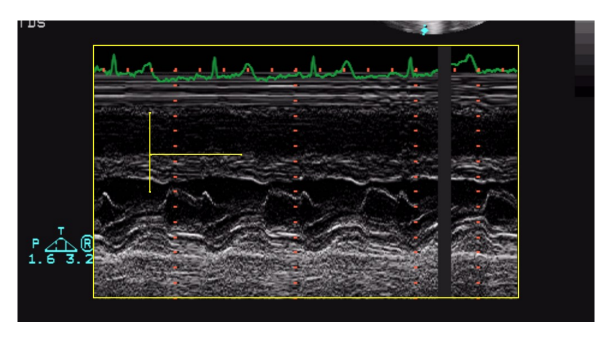
Next, establish (using the X axis and the Y axis) the calibration values to be assigned to the image. For instance, in case of a Doppler image, assign a time value on the X axis and a speed value on the Y axis. For 2D images, which have the same unit of measurement for both axis, it is also possible to calibrate both axis at once with a single diagonal segment using the *Both* button.



On the image, the mouse pointer will change into a pen and allow you to trace a segment. Click the left button once at the start point and then a second time at the end point. When finished, a dialogue window will prompt you to enter the dimension of the traced segment in the suitable unit of measurement.

It is also possible for the Doppler images to indicate the zero point position (the baseline) by the reference point button and then clicking it with the mouse.

Once the operation is performed on both axes you may decide if the calibration applies to the entire image (by clicking on *Add Region*) or only to a ROI (Region Of Interest). In the second case, select the ROI button and after tracing on the image the interest area, calibrate by clicking on the *Add Region* button.



When moving the mouse pointer on the calibrated region, you will see all the calibration information on the *Calibration* tab.

In the absence of a correct calibration all measurements on the images are invalid.

The program does not perform any control on the consistency of the calibration factor. It is <u>always</u> necessary to check the calculated calibration factor to verify whether it is admissible.

The accuracy of the calibration factor depends on the accuracy with which the segment is traced and it is inversely proportional to the pixel dimensions of the known structure. In the appendix, the reference accuracy and precision values of this approach are reported.

11.3.2 PERFORMING MEASUREMENTS

Now we are ready to perform some offline measurements on the images. ComPACS ECHO allows measuring 2D images, Doppler tracings, M-Mode images, and TDI images. For every data format there are specific measurements. For instance, in 2D the software allows measuring the length, area, perimeters, and calculates volumes. In Doppler, it allows measuring time, and speed, calculating flows, and other derived values.

Once the image is chosen, select the value to be measured on the measurement panel. In particular there are two panels: the *Generic* panel, which is always available, and the *Structured* panel which depends on the configuration set up and requires the opening of a report (intended as DICOM SR object).

The generic measurements are only temporary and will not be stored to a DICOM SR Object.

By clicking on the item, the mouse pointer becomes a pen. With a few mouse button clicks on the image, the software traces the measurement (in yellow) and saves it to the ComPACS Database in the DICOM SR format. The following picture shows an example of linear measurement on a 2D image:



To proceed, first select the desired measurement by clicking the chosen button in the control panel. Move the cursor to the starting point of the desired measurement and left-click once, the cursor will become a pen. Move the cursor to the end point of the measurement and left clicking once will terminate the measurement and populate the result in the chosen location. For measurements that are commonly performed in sequence, such as IVSD, LVID, LVPWD, each left mouse click will continue the sequence until the sequence is complete.

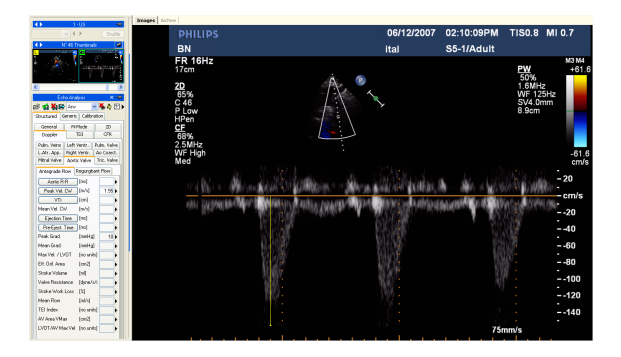
In more complex measurement case (i.e.: an area), an area point is fixed at every mouse left button click and the software creates a soft line passing through such points – "Spline" - to give it the harmonized shape. A double click terminates the measurement. In case of a tracing error, click the right mouse button to delete points and double right click to entirely cancel the measurement.

In the following example we have measured the left ventricle End Diastolic area in a 4 Chamber projection. ComPACS ECHO also automatically calculates the long axis and the Volume.

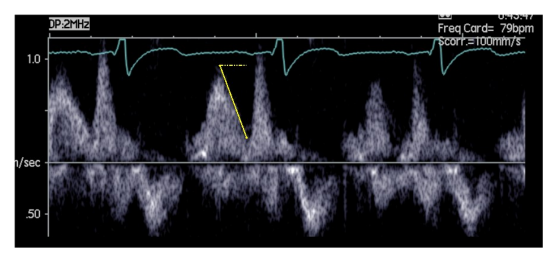


For velocity measurements as in the case of Doppler images, the software automatically identifies the baseline and in the case of a Doppler measurement where time is a factor, it creates a horizontal line.

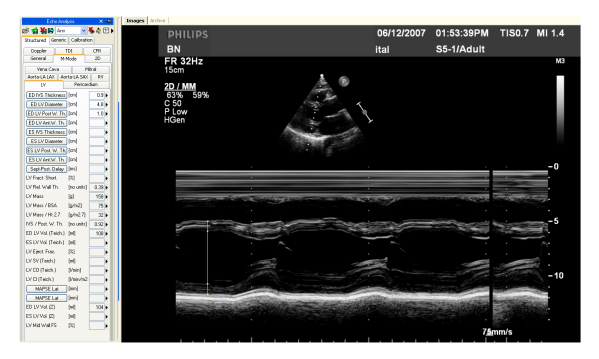
The following is an example of a velocity measurement on a Doppler image:



The next example is of a time measurement showing that even if the measurement is traced in diagonal way, the software takes the projection on the x axis.



M-Mode images have their own set of dedicated measurements. In the next example we see the intraventricular dimension measurements.



Not all the measurements are available on all types of images. For instance trying to perform a 2D area measurement on a Doppler image produces an error; in fact Doppler is not calibrated to make 2D area or length measurements. Or vice versa, if you try to measure a velocity on a 2D image, the software will again display an error message.

11.3.3 CALCULATIONS AND FORMULAS

The software not only allows performing direct measurements, but it is also capable of computing derived parameters by use of different mathematical formulas. For example, consider the left ventricle in the 4 chamber view. Once the area is measured in systole and diastole, the software computes the volumes with different methods (Area-length, Simpson) and then calculates the ejection fraction.



In the absence of a correct calibration all measurements on the images are invalid.

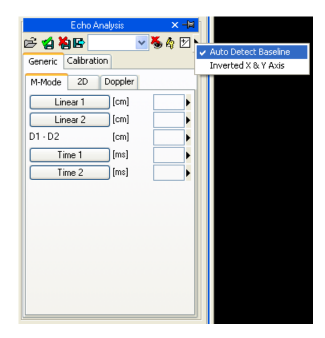
The program does not perform any control on the consistency of the calibration factor. It is <u>always</u> necessary to check the calculated calibration factor to verify whether it is admissible.

The program does not perform any control on the clinical validity of the measurement. The operator must visually verify the validity of any measurements taken.

11.4. MEASUREMENTS MANAGEMENT

The product allows for detailed management of the measurements taken.

It is possible to select the baseline and the axes during the measurement process. In fact, not all of the Doppler images have a baseline (the line of 0) set up correctly and a few 2D images have the X and Y axes reversed. In these situations, click the arrow on the right of the measurement panel and select the requested behavior with respect to such problems.

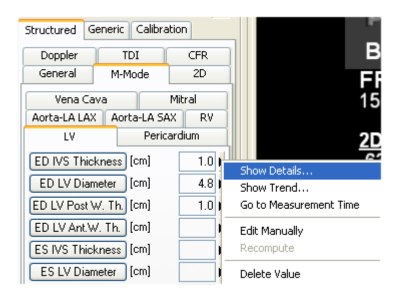


When a measurement is already present and re-executed, the software, by default, replaces the old value with the new one.

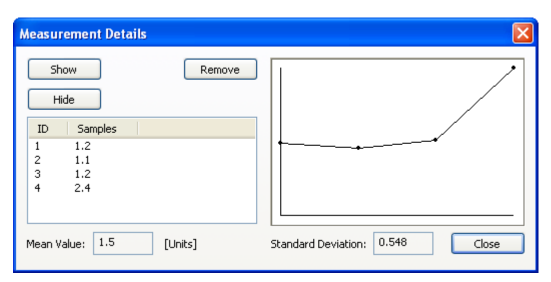
However more than one measurement on the same sample (or on various samples) can be performed and stored, calculating the mean value and the standard deviation. To perform multiple measurements and display the mean value click the button shown below. If this mode is active the button will turn yellow.



In the case of repeated measurements, the value shown nearby the item text is that of the mathematical average. However it is still possible to see all the measured samples. Alongside of every measurement is an arrow which allows accessing a management menu for the item.



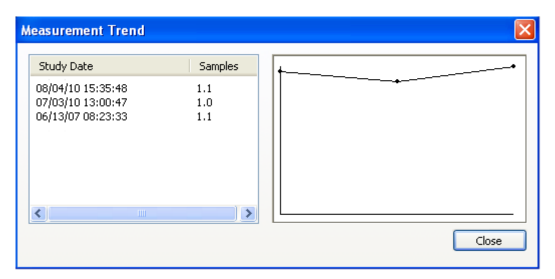
By clicking on the little arrow and then on *Show Details*, a window will be displayed in which all the samples taken for that item are shown together with the average and the standard deviation.



It is possible to delete a measured value if it seems incorrect. Simply select it and click on the *Remove* button. Use the buttons *Show* and *Hide* to show or hide the measurements on the image.

If echo analysis has been performed on the same patient in previous studies stored in the ComPACS database, you may view how a specific measurement field has varied over time.

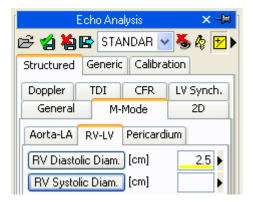
This operation can be performed by clicking on the *Show Trend* item in the menu. A window will be displayed in which the different values over time for the identified measurement are displayed both numerically and in a graphical format.



Another useful function is the ability to immediately display the image frame from which the measurement was taken. From the measurement menu, when clicking on the *Go to measurement time* option, the software automatically selects the image and the specific frame on which the measurement was performed.

The feature *Go to measurement time* is available only for off-line measurements taken with ComPACS ECHO.

Also by clicking on the *Edit Manually* button it is possible to manually insert the value instead of measuring it or calculating it. This value will be underlined in yellow to indicate it is a manually inserted measurement.



To show/hide the measurements on the images click on this icon:



To show the measurement graphics on all frames of the loop (and not just on the frame on which the measurement was taken) click on this icon:



To close and save the changes click on the Close and Save icon:



To close without saving the changes click on the Close Discarding changes icon:



11.5. CUSTOM SETTINGS

This component provides different user configurable settings, located in the *Program Settings* window, under the *Echo Analysis* tab.

Enable measurement average at startup

 Multiple measurements (if average is enabled)

 Do not display measurement label on image

 Invert X and Y axis

 Automatically detect Doppler baseline

 Use original pen cursor

[&]quot;Enable measurement average at startup" automatically enables the average function when the component is activated.

- "Multiple measurements (if average is enabled)" allows the user to measure a specific field multiple times, without having to press again the corresponding button. The measurement sequence can be interrupted by pressing the right mouse button.
- "Do not display measurement label on image" does not display the label and the measured value as an overlay on the image.
- "Invert X and Y axis" is used if the X and Y axis are to be interpreted differently in relation to the measurement structured menu. This option can be automatically enabled when the component is activated.
- "Automatically detect Doppler baseline" is used automatically detect the Doppler baseline and allow for faster, one-click, velocity measurements. This feature can be disabled if the DICOM image does not contain a correct encoding of the Doppler baseline correctly.
- "Use original pen cursor" uses the pen cursor of previous software revisions. Starting from 10.8.6 the pen cursor has been replaced with a sight view cursor.

12. ANGIOGRAPHIC ANALYSIS

12.1. ACTIVATING QCA II

If ComPACS QCA II is enabled on your system, the following button will be available on the toolbar, on the right hand side:



If the button is pressed down, then the module is already active. Otherwise it can be activated by clicking the button with the mouse.

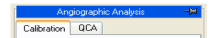
Upon activation the module will request a runtime license from the ComPACS Server. If the license cannot be received then the module will fail to activate and an error message will be displayed.

In case of concurrent licenses, it is possible that all available licenses have already been requested and assigned to other users that are using the module at that time and that no licenses are left for activation. This is a likely cause for failure.

Activation failure can also occur if the network connection to the server is no longer available.

Please ensure that the module is released, by releasing the above button, after you have finished using the product so that the license is available for use by other users.

When the module is active a window named "Angiographic Analysis" and containing a QCA tab is added and visible in the control window. It can be opened or minimized with a single left click on the blue bar.



12.2. ACTIVATING LVA

If ComPACS LVA is enabled on your system the following button will be available on the toolbar, on the right hand side:



If the button is pressed down, then the module is already active. Otherwise it can be activated by clicking the button with the mouse.

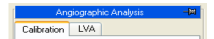
Upon activation the module will request a runtime license from the ComPACS Server. If the license cannot be received then the module will fail to activate and an error message will be displayed.

In case of concurrent licenses, it is possible that all available licenses have already been requested and assigned to other users that are using the module at that time and that no licenses are left for activation. This is a likely cause for failure.

Activation failure can also occur if the network connection to the server is no longer available.

Please ensure that the module is released, by releasing the above button, after you have finished using the product so that the license is available for use by other users.

When the module is active a window named "Angiographic Analysis" and containing an LVA tab is added and visible in the control window. It can be opened or minimized with a single left click of the mouse on the blue bar.



12.3. CALIBRATION

The calibration is an essential operation that must be carried out before any measurement or analysis on the images. This procedure leads to the computation of the zoom factor of the image by relating the measure in *pixels* of a known structure to its real measure in millimeters.

There are different ways to calibrate an image:

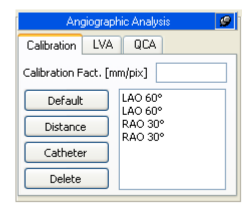
- default By selecting a pre-calculated (default) zoom factor because there are no known structures in the image, or the structure is significantly distorted (for example close to the image borders);
- distance By manually tracing a segment indicating the length of a structure of known size with the mouse (for example the distance between the markers in a catheter);
- catheter Automatically, based on a catheter (empty or contrast filled) of known diameter (for example 7 French).

Calibration can be carried out in all three ways by using the buttons in the *Calibration* tab on the *Angiographic Analysis* panel on the Control Window on the left.

In the absence of a correct calibration, all measurements on the images are invalid.

The program does not carry out any control on the consistency of the calibration factor. You must <u>always</u> check the calculated calibration factor to verify whether it is admissible.

12.3.1 DEFAULT CALIBRATION

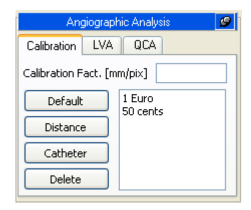


Default calibration can be carried out by clicking the *Default* button in the *Calibration* frame on the control window. In the white frame a list of names corresponding to the pre-set default calibration values are shown. Click with the left mouse button to select one. The corresponding calibration value will be displayed in the dedicated edit box above.

Alternatively, it is possible to directly set the calibration factor by selecting the related edit box with the mouse and by typing in the desired value.

The program does not carry out any control on the consistency of the calibration factor. The operator is entirely responsible for its value.

12.3.2 CALIBRATION BASED ON A KNOWN DISTANCE



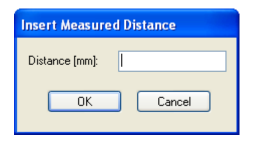
The calibration based on a known distance can be carried out by clicking the *Distance* button in the *Calibration* frame in the control. A list of names corresponding to the pre-set known distance values are shown in the white frame. Click with the left mouse button to select one.



On the image, click the left mouse button at one end of the segment to be measured and then release it. By moving the mouse it is possible to trace the desired segment. Click the left mouse button once again, the segment can be fixed. The related calibration factor will be automatically calculated and displayed in the suitable edit control. Alternatively, by right clicking the mouse button during segment tracing, the operation can be cancelled.

If a known distance has not been selected, a dialog box requesting the insertion of the length of the traced segment in millimeters will

be displayed. In this case the calibration factor will be then automatically calculated and displayed.

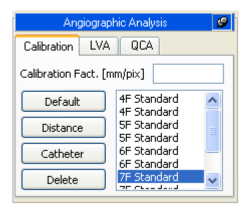


The program does not carry out any control on the consistency of the calibration factor. The operator is entirely responsible for its value.

It is necessary to calibrate on a known structure on the same plane as the ventricle to be measured.

The accuracy of the calibration factor depends on the accuracy with which the segment is traced and it is inversely proportional to the pixel dimensions of the known structure. In the appendix the reference accuracy and precision values of this approach are reported.

12.3.3 CATHETER BASED CALIBRATION



The calibration on a known catheter size can be carried out by clicking the *Catheter* button in the *Calibration* tab. In the white frame, a list of names corresponding to the pre-set known catheters are shown. By default, the catheter most recently used is selected and emphasized in blue If needed, select a different one with the left mouse button.

Now select two points along the catheter (preferably in the portion that is closest to the coronary lesion) by clicking the left mouse button on the image. The points are indicated by a yellow cross.



Around these points the program generates a region of interest in which the catheter skeleton and borders are automatically detected.

If the program does not detect the catheter skeleton or if the selected points are too close, an error is displayed by a message box. The operator will then have to select two different points or another image to perform the calibration.

When the analysis is completed, a results window will be shown in which the measured catheter borders are displayed in overlay. The image and the borders are zoomed to facilitate the verification of correctness of the measure. On the left, the calculated calibration factor and known diameter used are shown.

You must calibrate a catheter on the same plane as the ventricle to be measured. Also you calibrate a structure on the same plane as the ventricle to be measured.

The program does not carry out any control on the consistency of the calibration factor. The operator is entirely responsible of its value. For this reason you must <u>always</u> verify if the analysis is correct by comparing the measured profiles with those directly visible from the image.

The accuracy of the calibration factor depends on the image quality and varies if carried out on empty or contrast filled catheters. In the appendix, the reference accuracy and precision of this approach are discussed. Moreover the accuracy and precision depend on the overall correct system calibration to the images to be analyzed.

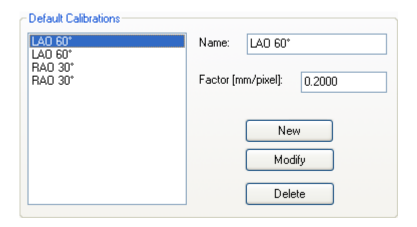
12.3.4 CALIBRATION SETTINGS

The calibration procedures require the presence of some reference information (default calibrations, known distances, known catheter dimensions). To avoid typing in these values for every computation, you may store this information and retrieve it when necessary.

To set these values, select the *Program Settings...* menu item from the *File menu*. These settings remain valid for all following activations of the program until they are modified again.

A dialog window will appear. Selecting the *Angiographic Analysis* tab, to set the required data using the following frames: *Default Calibrations*, *Reference Distances* and *Catheter Lists*. The procedure is similar for all frames (excepting, of course, the meaning of the values) and is as follows:

- The frame displays a list of names attributed to the values currently set.
- When a name in the window is selected, the name and the corresponding value are displayed on the right where they can be modified.
- To accept the modifications, click the *Modify* button.
- To remove some value from the list, click the *Delete* button.
- To set a new value and assign the name typed on the right, click the *New* button.

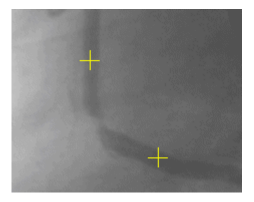


All changes will be applied when the *OK* button or the *Apply* button is pressed. Clicking the *Cancel* button will discard the changes.

12.4. CORONARY ANALYSIS

Having calculated the image calibration factor, it is possible to carry out the coronary analysis. Coronary stenosis measurements are carried out by selecting with the left mouse button, two points above and below the stenosis to be analyzed after having pressed the *2 Points* button in the *QCA* tab in the control window.

The selected points are indicated by yellow crosses. Around these points, the program generates a region of interest in which the centerline and the coronary borders are automatically determined. The program calculates the vessel diameter starting from the nearest points to the ones selected by the operator.



If the program cannot detect the coronary skeleton or if the selected points are too close, an error is reported. The operator will then have to select two further points or select a different image to analyze.

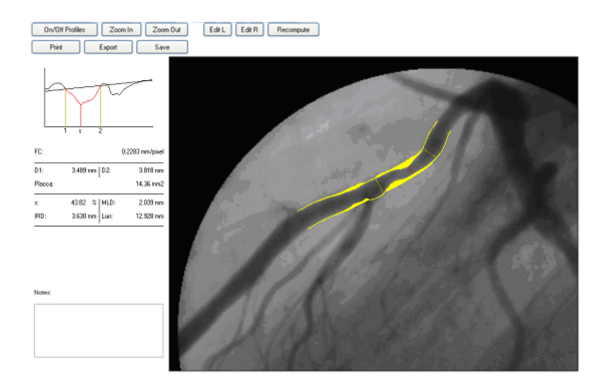
It is worth noting that the program finds the closest coronary path (in red) that connects the selected points. In some cases this may not be the desired path that needs to be analyzed. The problem can be overcome by indicating a third point in the path by activating the analysis with the *3 Points* button. In this case the intermediate point typically corresponds to the stenotic region or to a relevant way-point.

Finally you may carry out the analysis of a coronary vessel in which a stent has been inserted. In this case you need to click the *Stent* button and select four points on the vessel that will be indicated by yellow crosses. The first one corresponds to the start of the vessel we need to measure. The two intermediate points are needed to identify the beginning and the end of the region within the vessel in which the stent is present. The last one represents the end of the vessel.

The system will exclude the area in which the stent is present from the calculation to consider the profile as normal and will estimate three stenosis values: a proximal one within 5 mm from stent, an internal to stent itself, and one a distal one (within 5 mm from stent).

12.4.1 THE RESULTS WINDOW

When the analysis is concluded, a result window will appear displaying the detected borders in overlay on the analyzed image.



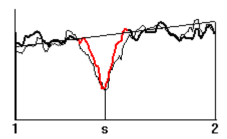
The image and the borders are zoomed to better visually verify the correctness of the measure. The coronary borders are displayed in yellow on the original image. The program also identifies the stenotic point of the vessel and the estimated normal profile. The former is indicated by a red segment perpendicular to the centerline whereas the latter is indicated in yellow. The region between the measured coronary borders and the normal profile is also filled with a yellow color.

On the left, the geometric analysis results are shown:

1) The calibration factor	(CF)
2) The proximal normal diameter	(D1)
3) The distal normal diameter	(D2)
4) The percentage of stenosis	(s)
5) The minimum luminal diameter	(MLD)
6) The interpolated reference diameter	(IRD)
7) The length of the stenosis	(LEN)

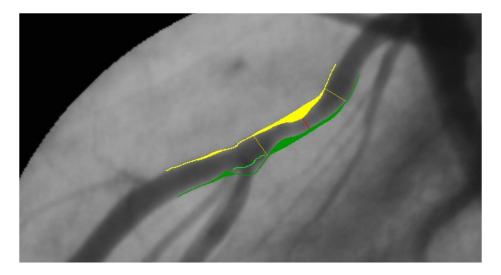
In the same window, a graph shows the diameter function of the analyzed vessel. The normal segments are identified by vertical segments labeled 1 and 2. The MLD is also identified by a vertical segment labeled s. Moreover an interpolated line indicates the estimated normal profile used in the computation of the percentage of stenosis.

The operator can modify the position of the segments 1, 2, and s by dragging them with the mouse on the graph window. The corresponding positions of such segments on the image are thus updated. When the mouse button is released, all measured parameters are recomputed on the basis of the new segment position.



Moreover, with a double click on the graph, the program will display the densitometric analysis results. The previous geometric analysis is reported in underlay for a direct comparison. A further mouse double click will return to the previous results display.

The software also allows you to manually modify the profiles in the situation where an error in the automatic estimate is discovered. This operation is performed by the buttons *Edit L* and *Edit R* according to what profile part you wish to modify. The area subject to modification will turn of green.



At this point the user must use the mouse to trace the part of the profile that was incorrect and click the *Recompute* button. The software will re-execute an estimate of the coronary profile on the basis of the provided information.

The window contains a set of further buttons with the following functions:

On/Off Profiles

It allows the activation/deactivation of the measured coronary borders overlay. It is thus possible to verify the automatic analysis by toggling the display on and off.

2 Zoom In It magnifies the image using interpolated zooming.

2 Zoom Out It reduces the image zoom factor using interpolated zooming.

• Print It prints a report of the results of the analysis complete of patient name and

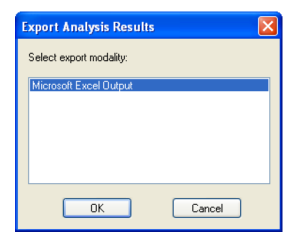
study and image references.

Export Exports the analysis results.

Save Generates a DICOM image containing the analysis results which can be

archived by the operator on the ComPACS Archive.

In particular the *Export* button allows you, depending on the installed export modalities, to export the analysis results to an external database, an Excel worksheet (if Microsoft Excel® is installed on the same PC), or event in proprietary or user defined formats.



On returning to the program interface, you may view the analysis results again by clicking the *Report* button. To remove the results, click the *Delete* button. The selection of a new image will also and automatically remove the results.

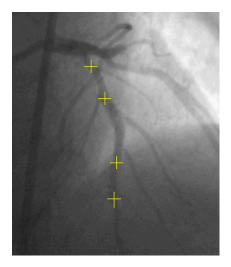
In the absence of a correct calibration, the measurements carried out on the images are invalid. It is always necessary to control the calculated calibration factor, the obtained measure and the borders traced to verify their admissibility.



In the appendix the accuracy and precision of the method are reported.

12.4.2 ANALYSIS AND STENT

To carry out the analysis of a coronary vessel in which a stent has been inserted, the same procedure for coronary analysis must be followed but clicking the *Stent* button instead of *2 Points* or *3 Points*.



To do so, select four points on the vessel that will be indicated by yellow crosses. The two intermediate points are needed to identify the beginning and the end of the region within the vessel in which the stent is present.

If the program cannot detect the coronary skeleton or if the selected points are too close, an error is reported in a message window. The operator will then have to select two further points or select a different image to analyze.

It is worth noting that the program finds the closest coronary path that connects the selected points (in red). In some cases this may not be the desired path that needs to be analyzed.

The stent position is used to remove the corresponding part of the vessel from the computation of the normal vessel borders. Moreover the program will calculate three stenosis values:

1) The stenosis within the stent (s)

2) The stenosis before the stent (p)

3) The stenosis after the stent (d)

For each stenosis, the percentage value, the absolute measured diameter, and the interpolated reference diameter are displayed. The stent position is indicated in a magenta color.

In the absence of a correct calibration, the measurements carried out on the images are invalid. It is always necessary to control the calculated calibration factor, the obtained measurement and the borders traced to verify their admissibility.



In the appendix the accuracy and precision of the method are reported.

12.5. VENTRICLE ANALYSIS

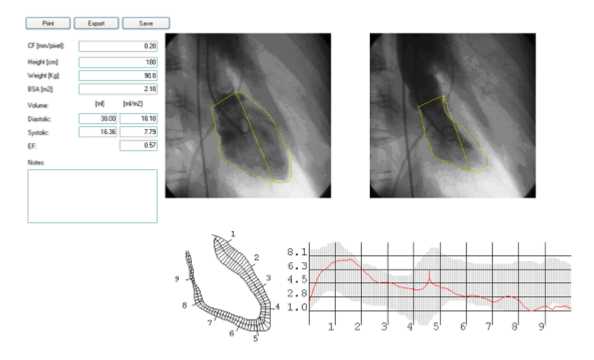
After having calculated the image calibration factor, it is possible to perform the left ventricle analysis with the following procedure.

- Select a diastolic image in the image window. Click the *Trace Diastolic* button in the *LVA* tab in the control window.
- Click the left mouse button on the image window and to trace the ventricle profile. When the button is released you may continue to trace the profile by clicking the left mouse button again or alternatively you may progressively delete the traced profile backwards by keeping the right mouse button pressed. Finally by double clicking, the traced profile will be fixed.
- Repeat the same procedure with the systolic image. Click the *Trace Systolic* button in the *LVA* tab in the control window. It is now possible to manually trace the systolic profile with the same procedure above.
- Click the *Analysis* button. In the main window, a tab called *LVA* with the results of the analysis (ventricular volumes, ejection fraction, etc...) will be displayed.

The ventricle profile tracing is not allowed if the calibration factor is null. You must calibrate the image before proceeding with the ventricle analysis.

The program does not perform any control on the consistency of the profiles traced by the operator. For this reason you must <u>always</u> verify if the traced profiles are correct by comparing the graphic trace in overlay.

An example of the result is shown below.



The window displays the ventricular volumes and ejection fraction calculated with the *Area-Length* method and the regional shortenings calculated with the *Centerline* method.

The volumes are expressed in ml and can be normalized with the body surface calculated from the patient height and weight. These values are taken from the main application and/or can be inserted/modified in this dialog window.

Regional shortenings are expressed as percentage values referred to the ventricle diastolic perimeter. In the background, in gray, is the normality region. It corresponds to the mean shortenings value +/- 2 units of standard deviation.

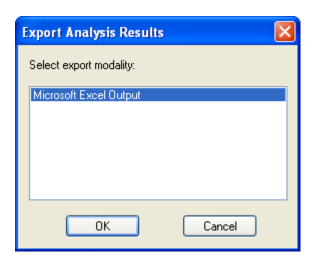
In this window it is also possible to insert annotations on the analysis and/or print a report. The latter is automatically set up and it reports patient information, study information, image information, the selected ventricle images, the results obtained and the possible annotations.

Using the *Save* button, the report will be generated as a DICOM image to allow the operator to archive it with the study images.

It is also possible, depending on the installed import and export modalities, to import patient weight and height information from an external database and/or to export the analysis results to the same database, an Excel worksheet (if Microsoft Excel® is installed on the same PC) or even in proprietary or user defined formats.

The import and export operations can be manually activated by clicking respectively the *Import* (not present if no import modalities are available) and *Export* buttons. The same operations can be

automatically activated when the dialog window is opened or closed. In this last case the user will be asked to confirm their activation.



Finally you may also perform the following operations:

- The profiles can be removed by clicking the *Delete* button.
- It is possible to display the previously selected systolic image by clicking the *Go to Systolic* button or by selecting the *Go to Diastolic* button to display the Diastolic image.

In the absence of a correct calibration and of a correct ventricle profile tracing, the measurements carried out on the images are invalid. It is necessary to control the calculated calibration factor, the obtained measure, and the profile traced to verify their admissibility.



In the appendix, the accuracy and precision of the method are discussed.

13. IVUS ANALYSIS

13.1. ACTIVATING IVUS ANALYSIS

If ComPACS IVUS is enabled on your system the following button will be available on the toolbar, on the right hand side:



If the button is pressed down, then the module is already active. Otherwise it can be activated by clicking the button with the mouse.

Upon activation the module will request a runtime license from the ComPACS Server. If the license cannot be received then the module will fail to activate and an error message will be displayed.

In case of concurrent licenses it is possible that all available licenses have already been requested and assigned to other users that are using the module at that time and that no licenses are left for activation. This is a likely cause for failure.

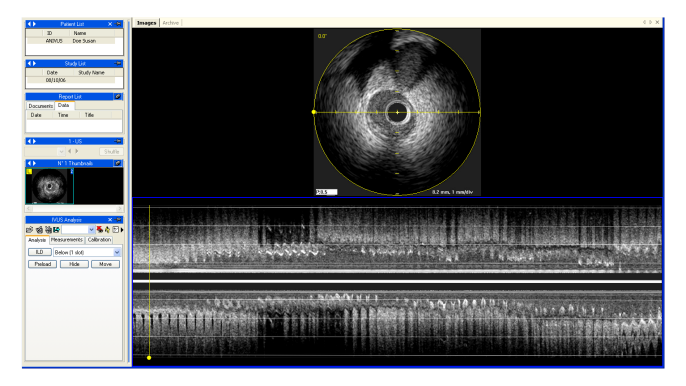
Activation failure can also occur if the network connection to the server is no longer available.

Please ensure that the module is released, by releasing the above button, after you have finished using the product so that the license is available for use by other users.

When the module is active, a window named "IVUS Analysis" is added and visible in the control window. It can be opened or minimized with a single left button click on the blue bar.



13.2. LONGITUDINAL RECONSTRUCTION (ILD)



If the DICOM object is an IVUS sequence, you may perform a longitudinal reconstruction in real time during loop playback by clicking the *ILD* button. The position and the image dimension depend on the drop-down window near the *ILD* button.

On the image, a yellow circle and a line show the undergoing reconstruction area and the sampling axis orientation. On the reconstructed image (ILD) a yellow line which shows the correspondence with the frame of the original sequence is shown. Such graphic information can be hidden with the *Hide* button.

The reconstruction is automatically performed during the clip sequence reproduction. It is possible to have it available all at once by clicking the *Preload* button.

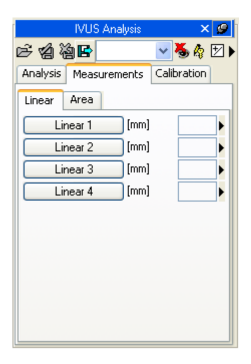
By clicking the *Move* button in the control window, the module allows a direct and intuitive modification of the sampling axis direction using the mouse. The reconstruction is then automatically updated, provided enough RAM is installed. Otherwise the file will have to be played back again for the 2D reconstruction update.

Similarly, by moving the yellow line along the image, it will be possible to move to the frame corresponding to the original image.

13.3. OFF-LINE MEASUREMENTS

The IVUS module offers the ability to perform measurements on IVUS images and the longitudinal reconstruction. The calibration information is usually contained in the DICOM file and the software uses it to avoid a manual calibration while at the same time ensure greater data accuracy. If the calibration information is absent, it is still possible to manually calibrate the image.

Simple generic measurements are always available in the Measurements tab of the IVUS Analysis window:



These are basic linear and area measurements for rapid verification purposes.

Generic measurements are only temporary. They are not stored to the ComPACS database and cannot be retained.

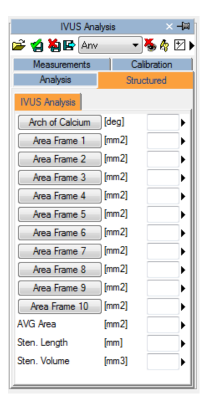
To access a more advanced measurement panel, open either an already existing report or create a new one. This operation can be performed by clicking on the toolbar icon as described in the "Managing Evidence and Report" chapter.

Structured measurement and reporting functionality is not available in the core IVUS module. IVUS II license or above is required.

The analysis window will be extended with an additional *Structured* tab containing all the structured measurements. The appearance and content of the tab will depend on the reporting templates and analysis views configured by the system administrator for the specific workstation or user.



Structured measurements are stored in the DICOM SR object they refer to.



In this interface, only the windows corresponding to the data present in the open object will be filled (with numbers). To edit the opened report click the *Editing* icon:



To close and save the changes made click on the Close and Save changes icon:

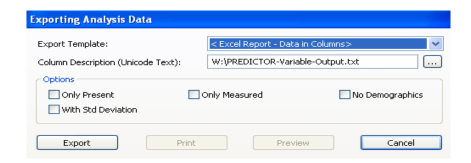


To close without saving any changes click on the Close discarding changes icon:



To export the data it is necessary to click the Export the report data to file icon:

An interface will be opened where you may select the data format and the file destination.



In the drop-down list, it is also possible to select an export template based on the installed configurations.

This functionality has been extended to export not only numerical values but also other field types (text, findings, dates, etc.). Furthermore the export also includes nested and repeated record structures.

The template is a simple text file which lists, in separate lines, the fields which need to be exported. Each line has the following format:

If the label or the unit are not specified, the default values shall be used.

For repeated records, the list of fields need to be enclosed in the following statement:

```
BEGINGROUP(Record Field ID); Label – Optional; Index – Optional ... list of fields ... ENDGROUP(Record Field ID)
```

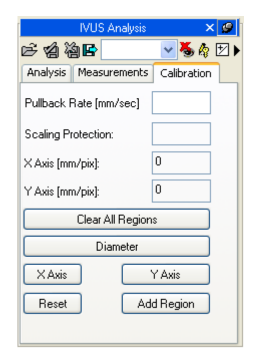
The index is optional and, if omitted, all records shall be exported.

Finally some special fields consist in DICOM fields. They are identified as follows:

\$(Group,Element); Label - Optional;

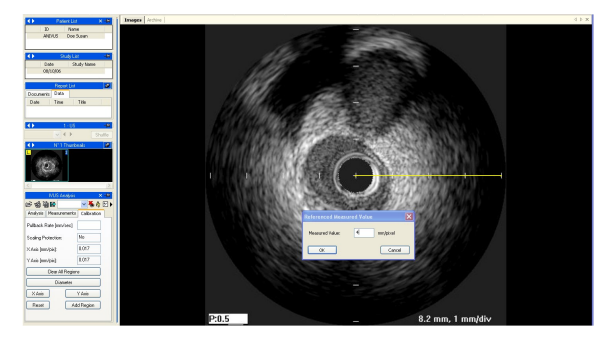
13.3.1 IMAGE CALIBRATION

To see if the image calibration is automatically imported or if it is absent, simply move the mouse cursor over the image. If present, the calibration information will be displayed in the *Calibration* tab.



If no information appears then the image needs a calibration. This operation must be executed very carefully because all subsequent measurements strictly depend on the calibration procedure.

To manually calibrate the image you have to determine the image axis values, using the X Axis and Y Axis buttons. The mouse pointer will change into a pen and it will allow you to trace a segment by clicking the mouse button in the center of the IVUS image and then a second time in the chosen axis end point. At the end, a dialog window will prompt you to type in the distance traced indicated in millimeters to perform calibration.



It is also possible to set the IVUS image diameter by clicking separately on the *Diameter* button. If such operation is not executed, the diameter is the greater value between the one identified by the X and the Y calibration.

Once this operation has been performed for both axes, the calibration can be accepted by clicking on the *Add Region* button. Now, when moving the mouse pointer over the calibrated area, you will see the calibration information displayed in the calibration tab.

In the absence of a correct calibration or pullback velocity, all measurements are invalid.

The program does not perform any control on the consistency of the calibration factor. The operator is therefore entirely responsible of its value. For this reason you must <u>always</u> verify if the analysis is correct by comparing the measured profiles with those directly visible from the image.

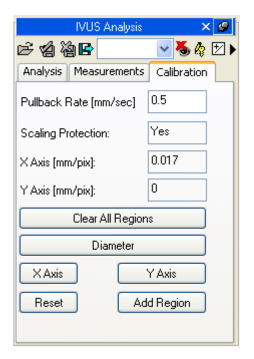
The accuracy of the calibration factor depends on accuracy of the operator and it depends on the image size in pixels

13.3.2 LONGITUDINAL RECONSTRUCTION CALIBRATION

The longitudinal reconstruction calibration factor is not directly set up but is imported by the images' reconstruction when performed. Therefore the IVUS images will have to be calibrated before performing the longitudinal reconstruction.

However, for the reconstruction to be fully calibrated, an additional parameter - the Pullback Speed – is required. Without this information it is not possible to perform measurements on the reconstructed image.

This parameter also is imported from DICOM (if present) but it can be manually set up in the *Calibration* tab after (and not before) the reconstruction has been created.



In the absence of a correct calibration or pullback velocity all measurements are invalid

The program does not perform any control on the consistency of the calibration factor. The operator is therefore entirely responsible of its value. For this reason you must <u>always</u> verify if the analysis is correct by comparing the measured profiles with those directly visible from the image.

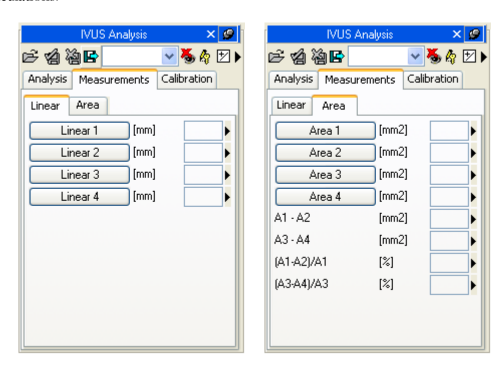
13.3.3 PERFORMING MEASUREMENTS

Now we are ready to perform some off-line measurements on the images. ComPACS IVUS allows measuring both the original IVUS image and the longitudinal reconstruction (ILD).

Once the image is chosen, select the value to be measured on the measurement panel. In particular there are two panels: the generic *Measurements* panel, which is always available, and the *Structured* panel which depends on the configuration set up and requires the opening of a report (intended as DICOM SR object).

The generic measurements are only temporary and will not be stored to a DICOM SR Object.

Once the user has selected the image, it is necessary to select the parameter requested in the measurement panel on the left. Possible measurements include: distances, areas, angles and computing derived calculations.

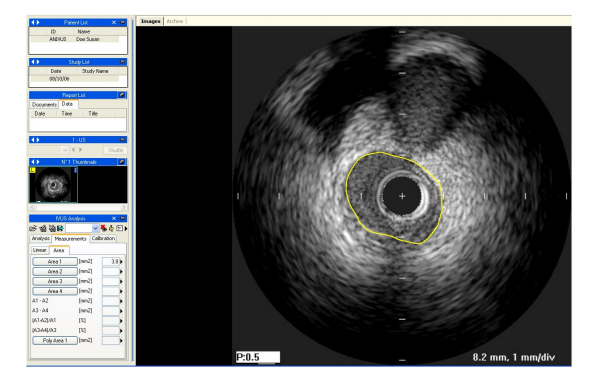


By clicking on the chosen button, the mouse pointer becomes a pen and with some simple clicks on the image, ComPACS traces the requested measurement.

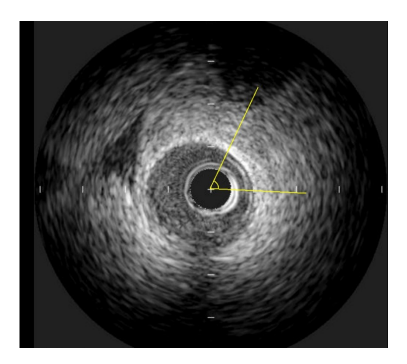
The following is an example of linear measurement on an IVUS image.



In case of area tracings, for each mouse click, a point of the area is fixed and the software creates a smooth line passing over these points — "spline". A double click sets the end of the measurement. In case of errors, you may delete points using the right mouse button or, with a double click, entirely cancel the measurement.



In case of angle measurements, the system automatically uses the center of the image as a reference point. It is therefore sufficient to click two points, moving clockwise, to measure an angle. In case of errors, you may delete a point using the right mouse button or, with a double click, entirely cancel the measurement.



13.3.4 CALCULATIONS AND FORMULAS

The software not only allows performing direct measurements, but it is also capable of computing derived parameters by use of different mathematical formulas. For example, if multiple area measurements are performed in different cross-sectional images over a manually inserted distance, it is possible to estimate the volume.

In the absence of a correct calibration or pullback velocity all measurements are invalid.

The program does not perform any control on the consistency of the calibration factor. The operator is entirely responsible of its value. For this reason you must <u>always</u> verify if the analysis is correct by comparing the measured profiles with those directly visible from the image.

The program does not perform any clinical validation of the measurement. The operator is therefore entirely responsible and must visually verify the accuracy of what has been measured.

13.4. MEASUREMENTS MANAGEMENT

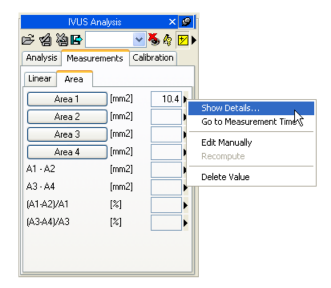
The software allows a detailed management of the measurements that have been taken.

When a measurement is already present and re-executed, the software, by default, replaces the old value with the new one.

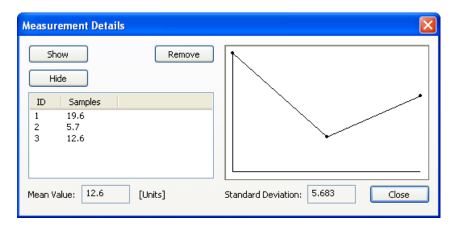
However, more than one measurement on the same sample (or on various samples) can be performed and stored, calculating the mean value and the standard deviation. To perform multiple measurements and display the mean value it is necessary to click the button shown below. If this mode is active, the button will turn yellow.

*

In cases of repeated measurements, the value shown near the item text is that of the mathematical average. However it is still possible to see all the measured samples. Alongside every measurement is an arrow that allows accessing a menu for that item.



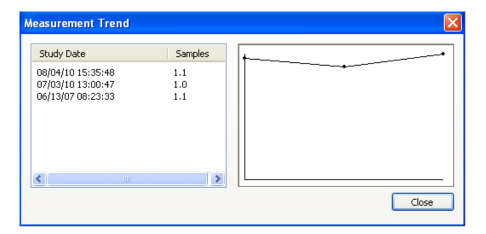
By clicking on the little arrow and then on *Show Details*, a window will be displayed in which all the samples taken for that item are shown together with the average and the standard deviation.



You may delete a measured value if it seems incorrect by simply selecting it and clicking the *Remove* button. Use the *Show* and *Hide* buttons to show or hide the measurements on the image.

If IVUS analysis has been performed on the same patient in previous studies stored in the ComPACS database, you may view how a specific measurement field has varied over time.

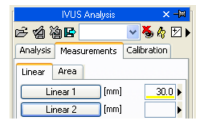
This operation can be performed by clicking on the *Show Trend* item in the menu. A window will be displayed in which the different values over time for the identified measurement are displayed both numerically and in a graphical format.



Another useful function is the ability to display the image frame from which the measurement was taken. Again on the *Measurement* menu, when clicking on the *Go to measurement time* option, the software automatically selects the image and the specific frame on which the measurement was performed.

The feature *Go to measurement time* is available only for off-line measurements taken with ComPACS IVUS.

Also by clicking on the *Edit Manually* button it is possible to manually insert the value instead of measuring it or calculating it. This value will be underlined in yellow to highlight it as an operator manually inserted measurement.



To show/hide the measurements on the images click on the icon:



To show the measurement on all frames of the loop (and not only on the frame on which the measurement was taken) click on the icon:



To close and save the changes click on the *Close and Save* icon:



To close without saving the changes click on the Close Discarding changes icon:



13.5. CUSTOM SETTINGS

This component provides different user configurable settings, located in the *Program Settings* window, under the *IVUS Analysis* tab.

- Enable measurement average at startup

 Multiple measurements (if average is enabled)

 Do not display measurement label on image

 Invert X and Y axis

 Use original pen cursor
- "Enable measurement average at startup" automatically enables the average function when the component is activated.
- "Multiple measurements (if average is enabled)" allows the user to measure a specific field multiple times, without having to press again the corresponding button. The measurement sequence can be interrupted by pressing the right mouse button.
- "Do not display measurement label on image" does not display the label and the measured value as an overlay on the image.
- "Invert X and Y axis" is used if the X and Y axis are to be interpreted differently in relation to the measurement structured menu. This option can be automatically enabled when the component is activated.
- "Use original pen cursor" uses the pen cursor of previous software revisions. Starting from 10.8.6 the pen cursor has been replaced with a sight view cursor.

14. GENERIC MEASUREMENTS

14.1. ACTIVATING OFF-LINE GENERIC MEASUREMENTS

If ComPACS Generic Measurements module is enabled on your system, the following button will be visible in the toolbar, on the right hand side:



If the button is pressed down then the module is already active. Otherwise it can be activated by clicking the button with the mouse.

Upon activation, the module will request a runtime license from the ComPACS Server. If the license cannot be received, then the module will fail to activate and an error message will be displayed.

In case of concurrent licenses it is possible that all available licenses have already been requested and assigned to other users that are using the module at that time and that no licenses are left for activation. This is a likely cause for failure.

Activation failure can also occur if the network connection to the server is no longer available.

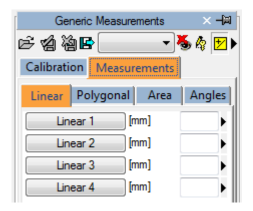
Please ensure that the module is released, by releasing the above button, after you have finished using the product so that the license is available for use by other users.

When the module is active, a window named "Generic Measurements" is added and visible in the control window. It can be opened or minimized with a single mouse left button click on the blue bar.



14.2. OFF-LINE MEASUREMENTS

The module provides the possibility of performing measurements on different types of images. Such simple measurements are always available in the *Measurements* tab of the analysis window:



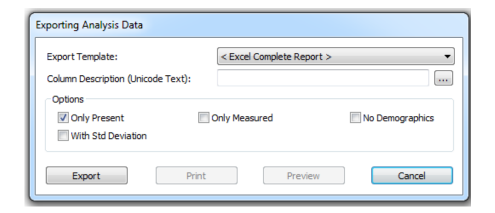
Linear, polygonal, area and angle measurements are available for rapid verification purposes.

Generic measurements are only temporary. They are not stored to the ComPACS database and cannot be retained.

To export the data it is necessary to click the Export the report data to file icon:



An interface will be opened where you may select the data format and the file destination.



In the drop-down list, it is also possible to select an export template based on the installed configurations.

This functionality has been extended to export not only numerical values but also other field types (text, findings, dates, etc.). Furthermore the export also includes nested and repeated record structures (for example each phase of a echo stress study).

The template is a simple text file which lists, in separate lines, the fields which need to be exported. Each line has the following format:

```
FieldID; Label - Optional; Unit - Optional
```

If the label or the unit are not specified, the default values shall be used.

For repeated records, the list of fields need to be enclosed in the following statement:

```
BEGINGROUP(Record Field ID); Label – Optional; Index – Optional ... list of fields ... ENDGROUP(Record Field ID)
```

The index is optional and, if omitted, all records shall be exported.

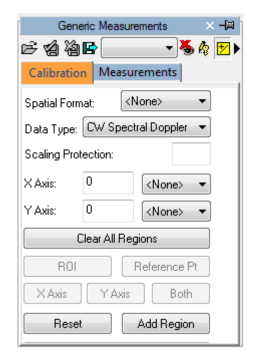
Finally some special fields consist in DICOM fields. They are identified as follows:

```
$(Group, Element); Label - Optional;
```

14.2.1 IMAGE CALIBRATION

The calibration information is usually contained in the DICOM file. The product exploits it to avoid a manual calibration and ensure greater accuracy. Therefore calibrating images in order to perform measurements is generally not necessary nor is it recommended.

By simply moving the mouse over the image, the calibration information will appear in the *Calibration* tab. This way it is possible to quickly verify if the image calibration is present or not.



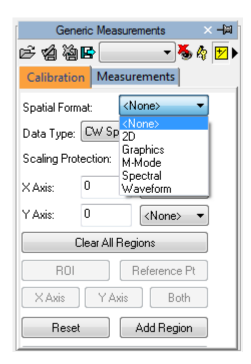
If no information is displayed when moving the mouse, the image needs to be calibrated.

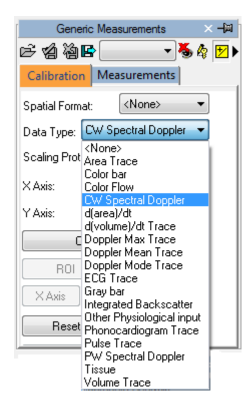
Calibration for radiology images (DICOM Pixel Spacing) is supported. It is however important that this calibration is done on the receiving surface which is not necessarily in the same plane and orientation as the structure that is measured. This may lead to inaccurate results. Verifying the calibration (or manually recalibrating) using a structure of known size in the same plane and orientation is recommended.

Ultrasound images can contain multiple regions of calibration. Therefore it is possible that the calibration information differs in relation to the position of the mouse on the image. Also some regions may be calibrated whereas in others the calibration information may be absent.

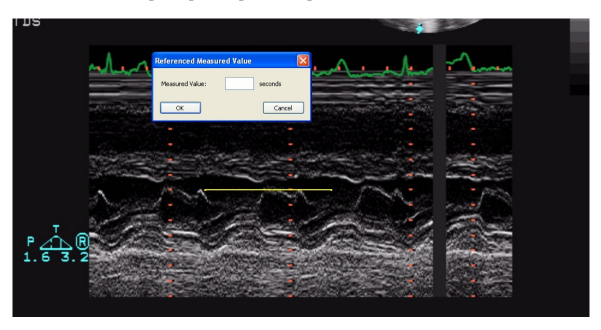
If the calibration information is absent, it is possible to manually calibrate the entire image or a region.

The first thing to do to calibrate an image is to choose the Spatial Format (2D, Doppler, M-Mode, etc...) and the type of data (Tissue, PW, CW, etc...). Using the two drop down menus in the *Calibration* tab:





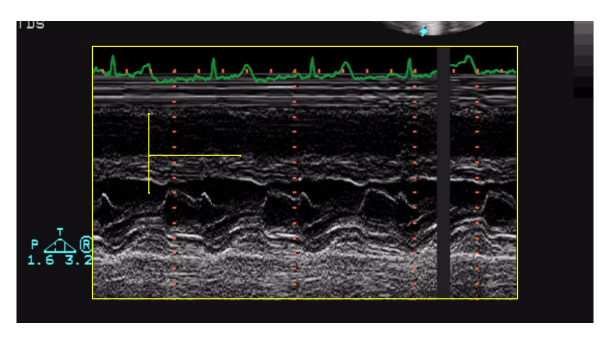
Next, establish (using the X axis and the Y axis) the calibration values to be assigned to the image. For instance, in case of a Doppler image, assign a time value on the X axis and a speed value on the Y axis. For 2D images, which have the same unit of measurement for both axis, it is also possible to calibrate both axis at once with a single diagonal segment using the *Both* button.



On the image, the mouse pointer will change into a pen and allow you to trace a segment. Click the left button once at the start point and then a second time at the end point. When finished, a dialogue window will prompt you to enter the dimension of the traced segment in the suitable unit of measurement.

It is also possible for the Doppler images to indicate the zero point position (the baseline) by the reference point button and then clicking it with the mouse.

Once the operation is performed on both axes you may decide if the calibration applies to the entire image (by clicking on *Add Region*) or only to a ROI (Region Of Interest). In the second case, select the ROI button and after tracing on the image the interest area, calibrate by clicking on the *Add Region* button.



When moving the mouse pointer on the calibrated region, you will see all the calibration information on the *Calibration* tab.

In the absence of a correct calibration all measurements on the images are invalid.

The program does not perform any control on the consistency of the calibration factor. It is <u>always</u> necessary to check the calculated calibration factor to verify whether it is admissible.

The accuracy of the calibration factor depends on the accuracy with which the segment is traced and it is inversely proportional to the pixel dimensions of the known structure. In the appendix, the reference accuracy and precision values of this approach are reported.

Now we are ready to perform some offline measurements on the images. Once the image is chosen, select the value to be measured on the measurement panel.

The generic measurements are only temporary and will not be stored to a DICOM SR Object.

By clicking on the item, the mouse pointer becomes a pen. With a few mouse button clicks on the image, the software traces the measurement (in yellow).

To proceed, first select the desired measurement by clicking the chosen button in the control panel. Move the cursor to the starting point of the desired measurement and left-click once, the cursor will become a pen. Move the cursor to the end point of the measurement and left clicking once will terminate the measurement and populate the result in the chosen location.

In more complex measurement case (i.e.: an area), an area point is fixed at every mouse left button click and the software creates a soft line passing through such points – "Spline" - to give it the harmonized shape. A double click terminates the measurement. In case of a tracing error, click the right mouse button to delete points and double right click to entirely cancel the measurement.

Not all the measurements are available on all types of images. For instance trying to perform a 2D area measurement on a Doppler image produces an error; in fact Doppler is not calibrated to make 2D area or length measurements.

14.2.3 CALCULATIONS AND FORMULAS

The software not only allows performing direct measurements, but it is also capable of computing derived parameters by use of different mathematical formulas.



In the absence of a correct calibration all measurements on the images are invalid.

The program does not perform any control on the consistency of the calibration factor. It is <u>always</u> necessary to check the calculated calibration factor to verify whether it is admissible.

The program does not perform any control on the clinical validity of the measurement. The operator must visually verify the validity of any measurements taken.

14.3. MEASUREMENTS MANAGEMENT

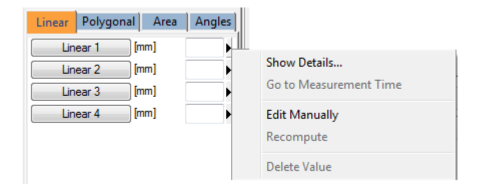
The product allows for detailed management of the measurements taken.

When a measurement is already present and re-executed, the software, by default, replaces the old value with the new one.

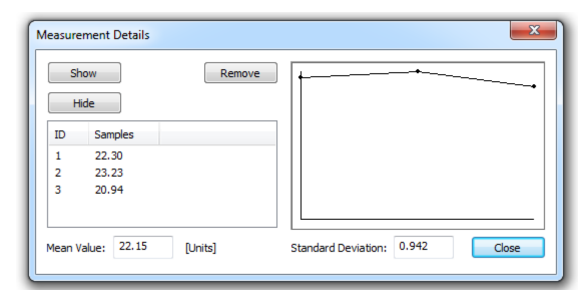
However more than one measurement on the same sample (or on various samples) can be performed and stored, calculating the mean value and the standard deviation. To perform multiple measurements and display the mean value click the button shown below. If this mode is active the button will turn yellow.



In the case of repeated measurements, the value shown nearby the item text is that of the mathematical average. However it is still possible to see all the measured samples. Alongside of every measurement is an arrow which allows accessing a management menu for the item.



By clicking on the little arrow and then on *Show Details*, a window will be displayed in which all the samples taken for that item are shown together with the average and the standard deviation.



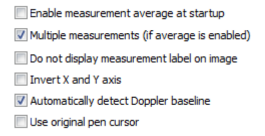
It is possible to delete a measured value if it seems incorrect. Simply select it and click on the *Remove* button. Use the buttons *Show* and *Hide* to show or hide the measurements on the image.

Another useful function is the ability to immediately display the image frame from which the measurement was taken. From the measurement menu, when clicking on the *Go to measurement time* option, the software automatically selects the image and the specific frame on which the measurement was performed.

The feature *Go to measurement time* is available only for off-line measurements taken with ComPACS.

14.4. CUSTOM SETTINGS

This component provides different user configurable settings, located in the *Program Settings* window, under the *Generic Measurements* tab.



- "Enable measurement average at startup" automatically enables the average function when the component is activated.
- "Multiple measurements (if average is enabled)" allows the user to measure a specific field multiple times, without having to press again the corresponding button. The measurement sequence can be interrupted by pressing the right mouse button.
- "Do not display measurement label on image" does not display the label and the measured value as an overlay on the image.
- "Invert X and Y axis" is used if the X and Y axis are to be interpreted differently in relation to the measurement structured menu. This option can be automatically enabled when the component is activated.
- "Automatically detect Doppler baseline" is used automatically detect the Doppler baseline and allow for faster, one-click, velocity measurements. This feature can be disabled if the DICOM image does not contain a correct encoding of the Doppler baseline correctly.
- "Use original pen cursor" uses the pen cursor of previous software revisions. Starting from 10.8.6 the pen cursor has been replaced with a sight view cursor.

15. PHILIPS QLAB

15.1. ACTIVATING PHILIPS QLAB

If Philips QLAB integrated in ComPACS Workstation is enabled on your system, the following button will be available on the toolbar, on the right hand side:



If the button is pressed down, then the module is already active. Otherwise it can be activated by clicking the button with the mouse.

Upon activation the module will request a runtime license from the ComPACS Server. If the license cannot be received then the module will fail to activate and an error message will be displayed.

In case of concurrent licenses it is possible that all available licenses have already been requested and assigned to other users that are using the module at that time and that no licenses are left for activation. This is a likely cause for failure.

Activation failure can also occur if the network connection to the server is no longer available.

Please ensure that the module is released, by releasing the above button, after you have finished using the product so that the license is available for use by other users.

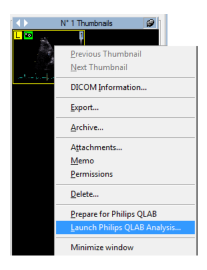
Philips QLAB, in addition to the core viewer, comprises of different analysis modules that can be separately licensed. The specific modules licensed for your workstation can be viewed by clicking on the arrow to the right hand side of the activation button. A drop down window will be displayed, listing the available modules.



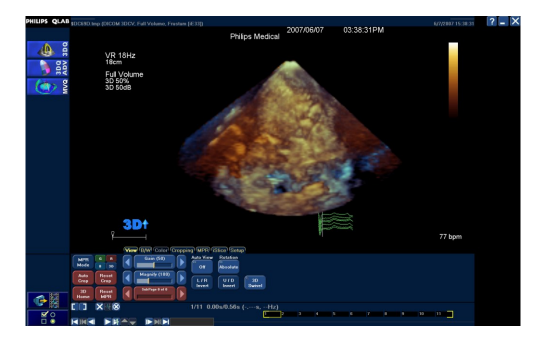
In this window you may to select or de-select the modules that you wish to activate. When clicking on the QLAB button only the license for the selected modules will be requested.

15.2. LAUNCHING PHILIPS QLAB

The Philips QLAB plug-in can be activated simply by right clicking in the thumbnail window on the item to be analyzed. The following menu will be displayed:

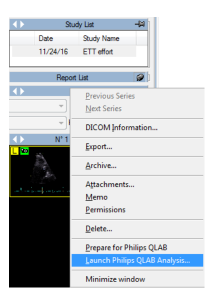


By selecting Launch Philips QLAB Analysis the plugin will be immediately opened on such selected image.



Alternatively, you may add the selected image to the analysis queue. This can be used if more than one item needs to be passed to the QLAB application.

Furthermore, by right clicking on the series window, you may directly analyze an entire series. The following menu will be used:



When working within the QLAB application, the content of the screen can be saved, either as an image or as a loop, using the following button:



The saved object will be automatically stored to the archive together with the original image analyzed.

Similarly it is also possible to export DICOM Structured Report data using the following button:



The report data will also be automatically imported in the opened report, if there is one open for editing at the time the data is exported.

This guide describes only how the Philips QLAB application can be launched. For usage information, cautions and warnings, and any other instruction please refer to the QLAB user guides and documentation.

16. INVIA CORRIDOR 4DM

16.1. ACTIVATING INVIA CORRIDOR 4DM

If INVIA Corridor 4DM is integrated in ComPACS Workstation is enabled on your system, the following button will be available on the toolbar, on the right hand side:



If the button is pressed down, then the module is already active. Otherwise it can be activated by clicking the button with the mouse.

Upon activation the module will request a runtime license from the ComPACS Server. If the license cannot be received then the module will fail to activate and an error message will be displayed.

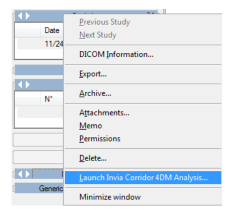
In case of concurrent licenses it is possible that all available licenses have already been requested and assigned to other users that are using the module at that time and that no licenses are left for activation. This is a likely cause for failure.

Activation failure can also occur if the network connection to the server is no longer available.

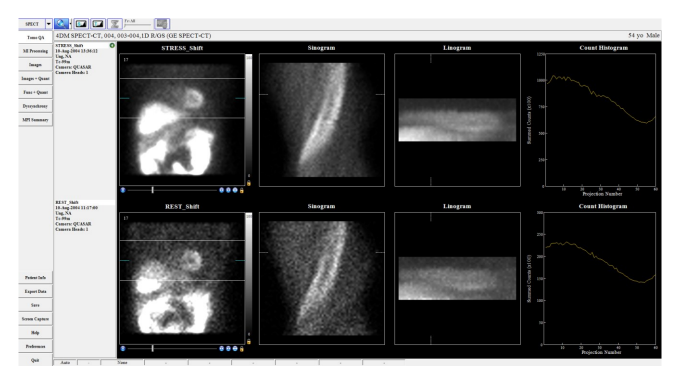
Please ensure that the module is released, by releasing the above button, after you have finished using the product so that the license is available for use by other users.

16.2. LAUNCHING INVIA CORRIDOR 4DM

The INVIA Corridor 4DM plug-in can be activated simply by right clicking in either the patient window or the study window on the item to be analyzed. Typically this operation is performed at the study level.



By selecting Launch Invia Corridor 4DM Analysis the plugin will be immediately opened on such selected image.



When working within the INVIA Corridor 4CM application, secondary captures or performed analysis can be saved. The saved object will be automatically stored to the archive together with the original image analyzed.

This guide describes only how the INVIA Corridor 4DM application can be launched. For usage information, cautions and warnings, and any other instruction please refer to the INVIA Corridor 4DM user guides and documentation.

17. GE ECHOPAC PLUGIN

17.1. ACTIVATING GE ECHOPAC PLUGIN

If General Electric EchoPAC Plugin integrated in ComPACS Workstation is enabled on your system, the following button will be available on the toolbar, on the right hand side:



If the button is pressed down, then the module is already active. Otherwise it can be activated by clicking the button with the mouse.

Upon activation the module will request a runtime license from the ComPACS Server in addition to the local workstation license provided by General Electric. If the license cannot be received then the module will fail to activate and an error message will be displayed.

In case of concurrent licenses it is possible that all available licenses have already been requested and assigned to other users that are using the module at that time and that no licenses are left for activation. This is a likely cause for failure.

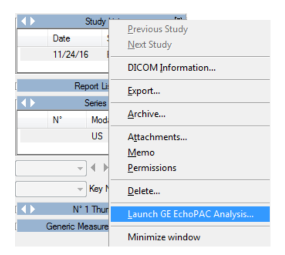
Activation failure can also occur if the network connection to the server is no longer available.

Please ensure that the module is released, by releasing the above button, after you have finished using the product so that the license is available for use by other users.

17.2. LAUNCHING GE ECHOPAC PLUGIN

The EchoPAC plug-in can be activated simply by right clicking in either the patient window, the study window, the series window or even the thumbnail window on the item to be analyzed. Typically this operation is performed at the study level.

The following menu will be displayed:



By selecting Launch GE EchoPAC Analysis... the plugin will be immediately opened on such selected item.

When working within the EchoPAC application, the content of the screen can be saved, either as an image or as a loop, using the following button:



The saved object will be automatically stored to the archive together with the original image analyzed.

If any measurement is performed in the application, automatically a DICOM Structured Report will be saved once the user exits the application. This will allow the results of the measurements to be imported into the report.

This guide describes only how the GE EchoPAC Plugin application can be launched. For usage information, cautions and warnings, and any other instruction please refer to the EchoPAC Plugin user guides and documentation.

18. TOMTEC ANALYSIS

18.1. ACTIVATING TOMTEC ANALYSIS

If TomTec Analysis modules integrated in ComPACS Workstation are enabled on your system, the following button will be available on the toolbar, on the right hand side:



If the button is pressed down, then the module is already active. Otherwise it can be activated by clicking the button with the mouse.

Upon activation the module will request a runtime license from the ComPACS Server in addition to the license provided by the TomTec licensing server. If the license cannot be received then the module will fail to activate and an error message will be displayed.

In case of concurrent licenses it is possible that all available licenses have already been requested and assigned to other users that are using the module at that time and that no licenses are left for activation. This is a likely cause for failure.

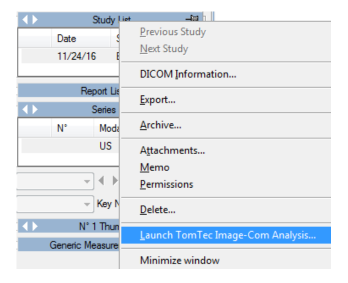
Activation failure can also occur if the network connection to the server is no longer available.

Please ensure that the module is released, by releasing the above button, after you have finished using the product so that the license is available for use by other users.

TomTec, in addition to the core Image-Com viewer, comprises of different analysis modules that can be separately licensed. The specific modules licensed for your workstation can be viewed within the TomTec application once the TomTec viewer is launched.

18.2. LAUNCHING TOMTEC ANALYSIS

The TomTec Analysis plug-in can be activated simply by right clicking in either the patient window, the study window, the series window or even the thumbnail window on the item to be analyzed. Typically this operation is performed at the study level.



By selecting Launch TomTec Image-Com Analysis... the plugin will be immediately opened on such selected item.

When working within the TomTec application, the content of the screen can be saved, either as an image or as a loop, using the following button:



The saved object will be automatically stored to the archive together with the original image analyzed.

If any measurement is performed in the application, automatically a DICOM Structured Report will be saved once the user exits the application. This will allow the results of the measurements to be imported into the report.

This guide describes only how the TomTec Image-Com application can be launched. For usage information, cautions and warnings, and any other instruction please refer to the TomTec user guides and documentation.

19. MEDIS ANALYSIS

19.1. ACTIVATING MEDIS ANALYSIS

If Medis Analysis modules integrated in ComPACS Workstation are enabled on your system, the following button will be available on the toolbar, on the right hand side:



If the button is pressed down, then the module is already active. Otherwise it can be activated by clicking the button with the mouse.

Upon activation the module will request a runtime license from the ComPACS Server in addition to the license provided by the Medis licensing system. If the license cannot be received then the module will fail to activate and an error message will be displayed.

In case of concurrent licenses it is possible that all available licenses have already been requested and assigned to other users that are using the module at that time and that no licenses are left for activation. This is a likely cause for failure.

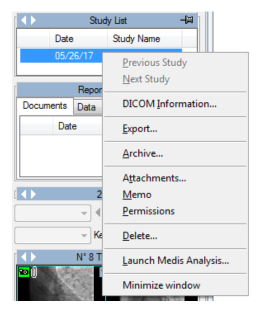
Activation failure can also occur if the network connection to the server is no longer available.

Please ensure that the module is released, by releasing the above button, after you have finished using the product so that the license is available for use by other users.

Medis, in addition to the core Medis Suite viewer, comprises of different analysis modules that can be separately licensed. The specific modules licensed for your workstation can be viewed within the Medis application once the Medis Suite viewer is launched.

19.2. LAUNCHING MEDIS ANALYSIS

The Medis Analysis plug-in can be activated simply by right clicking in either the patient window, the study window, the series window or even the thumbnail window on the item to be analyzed. Typically this operation is performed at the study level.



By selecting Launch Medis Analysis... the plugin will be immediately opened on such selected item.

When working within the Medis application, the content of the screen can be saved, either as an image or as a loop.

The saved object will be automatically stored to the archive together with the original image analyzed.

If any measurement is performed in the application, automatically a DICOM Structured Report will be saved once the user exits the application. This will allow the results of the measurements to be imported into the report.

This guide describes only how the Medis Suite application can be launched. For usage information, cautions and warnings, and any other instruction please refer to the Medis user guides and documentation.

20. EPSILON ECHO INSIGHT ANALYSIS

20.1. ACTIVATING EPSILON ANALYSIS

If Epsilon Echo Insight Analysis module integrated in ComPACS Workstation is enabled on your system, the following button will be available on the toolbar, on the right hand side:



If the button is pressed down, then the module is already active. Otherwise it can be activated by clicking the button with the mouse.

Upon activation the module will request a runtime license from the ComPACS Server in addition to the license provided by the Epsilon licensing system. If the license cannot be received then the module will fail to activate and an error message will be displayed.

In case of concurrent licenses it is possible that all available licenses have already been requested and assigned to other users that are using the module at that time and that no licenses are left for activation. This is a likely cause for failure.

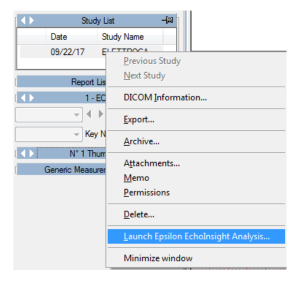
Activation failure can also occur if the network connection to the server is no longer available.

Please ensure that the module is released, by releasing the above button, after you have finished using the product so that the license is available for use by other users.

Epsilon, in addition to the core Epsilon viewer, comprises of different analysis modules that can be separately licensed. The specific modules licensed for your workstation can be viewed within the Epsilon application once the Epsilon viewer is launched.

20.2. LAUNCHING ECHO INSIGHT ANALYSIS

The Echo Insight Analysis plug-in can be activated simply by right clicking in either the patient window, the study window, the series window or even the thumbnail window on the item to be analyzed. Typically this operation is performed at the study level.



By selecting Launch Epsilon Echo Insight Analysis... the plugin will be immediately opened on such selected item.

When working within the Epsilon application, the content of the screen can be saved, either as an image or as a loop.

The saved object will be automatically stored to the archive together with the original image analyzed.

If any measurement is performed in the application, automatically a DICOM Structured Report will be saved once the user exits the application. This will allow the results of the measurements to be imported into the report.

This guide describes only how the Epsilon application can be launched. For usage information, cautions and warnings, and any other instruction please refer to the Epsilon user guides and documentation.

21. CIRCLE CVI42 ANALYSIS

21.1. ACTIVATING CVI42 ANALYSIS

If Circle CVI42 Analysis module integrated in ComPACS Workstation is enabled on your system, the following button will be available on the toolbar, on the right hand side:



If the button is pressed down, then the module is already active. Otherwise, it can be activated by clicking the button with the mouse.

Upon activation the module will request a runtime license from the ComPACS Server in addition to the license provided by the Circle licensing system. If the license cannot be received then the module will fail to activate and an error message will be displayed.

In case of concurrent licenses it is possible that all available licenses have already been requested and assigned to other users that are using the module at that time and that no licenses are left for activation. This is a likely cause for failure.

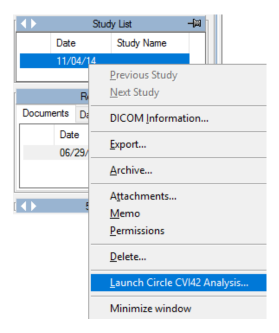
Activation failure can also occur if the network connection to the server is no longer available.

Please ensure that the module is released, by releasing the above button, after you have finished using the product so that the license is available for use by other users.

CVI42, in addition to the core application, comprises of different analysis modules that can be separately licensed. The specific modules licensed for your workstation can be viewed within the CVI42 application once the it is launched.

21.2. LAUNCHING CVI42 ANALYSIS

The CVI42 Analysis plug-in can be activated simply by right clicking in the study window on the study to be analyzed.



By selecting Launch Circle CVI42 Analysis... the plugin will be immediately opened on such selected study. At first launch, the user will need to authenticate also in CVI42.

When working within the CVI42 application the results will be stored directly to the ComPACS server and added to the analyzed study.

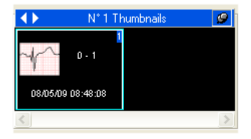
This guide describes only how the Circle CVI42 application can be launched. For usage information, cautions and warnings, and any other instruction please refer to the Circle CVI42 user guides and documentation.

22. ECG MANAGEMENT

22.1. REVIEW OF DICOM ECG

In order to review DICOM ECG in ComPACS, the ECG Viewer module needs to be licensed. The ECG viewer does not require any user manual activation: the license is automatically activated at login.

If a DICOM ECG object is present for the selected study, a thumbnail similar to the one shown below will be visible in the *Thumbnails* window.



Clicking the thumbnail will open the ECG DICOM object for review.

At the top of the ECG display window, if the ECG is opened in full screen mode, a set of controls are visible:



The Sequence drop down control lists all the ECG sequences available. The Display Format drop down control allows you to choose the preferred display format. The Report Format drop down control allows you to choose the preferred format for report generation. For certain multi-page reports it is possible to choose between a lead based or a time based layout: in the former each page will contain a subset of the leads whereas in the latter each page will contain a subset of the timeline.

The Speed drop down and the Calibration drop down define the horizontal and vertical calibration.

The Filter drop down defines the post-processing filter applied, if any.

The *Print* button allows you to print the selected ECG sequence in real scale units in the Report Format.

The Report button allows you to generate a report or open the most recent report if already existing.

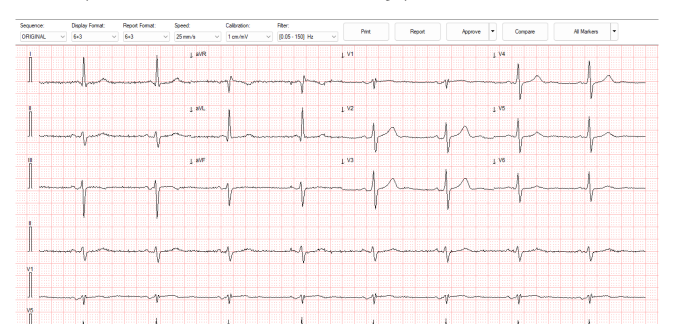
The *Approve* button allows you to directly generate and approve the report based on the imported interpretation and the default signature method for the selected template. The associated drop-down menu allows the user selection of the signature method.

The *Compare* button allows you to compare different ECG traces for the same patient, if such traces have all been previously opened for review.

The All Markers button displays the detected fiducial points, if present in the DICOM file. These points will be used for median beat alignment in overlay comparison mode.

By using the drop down menu associated to the *All Markers* button, it is possible to limit the display to the specific type of fiducial point of interest: for example *P Onset* or *QRS Onset*. The button name will change to reflect the selection.

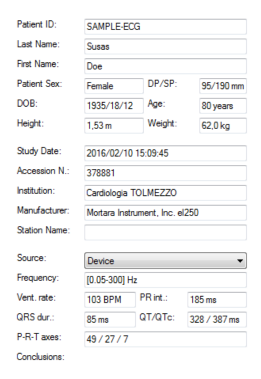
Immediately below the user controls the ECG traces are displayed.



A horizontal slider may be present if the entire ECG trace timeline cannot be displayed at one time. By moving the slider it is possible to review the entire timeline.

A vertical slider may be present if all traces cannot be displayed at one time. By moving the slider it is possible to view the entire list of traces, on the basis of the selected display format.

Instead, on the right, information about the patient, study and ECG object is displayed, including report information. Please note that some information may not be visible if not present in the DICOM file.



ECG objects can also be processed with the same tools described in this manual for image objects. For example the features related to zooming, exporting and storing are all available.

22.2. ACTIVATING ECG ANALYSIS AND REPORTING

If the ECG analysis and reporting module is enabled on your system, the following button will be visible in the toolbar, on the right hand side:



If the button is pressed down then the module is already active. Otherwise it can be activated by clicking the button with the mouse.

Upon activation, the module will request a runtime license from the ComPACS Server. If the license cannot be received, then the module will fail to activate and an error message will be displayed.

In case of concurrent licenses it is possible that all available licenses have already been requested and assigned to other users that are using the module at that time and that no licenses are left for activation. This is a likely cause for failure.

Activation failure can also occur if the network connection to the server is no longer available.

Please ensure that the module is released, by releasing the above button, after you have finished using the product so that the license is available for use by other users.

When the module is active, a window named "ECG Analysis" is added and visible in the control window. It can be opened or minimized with a single mouse left button click on the blue bar.



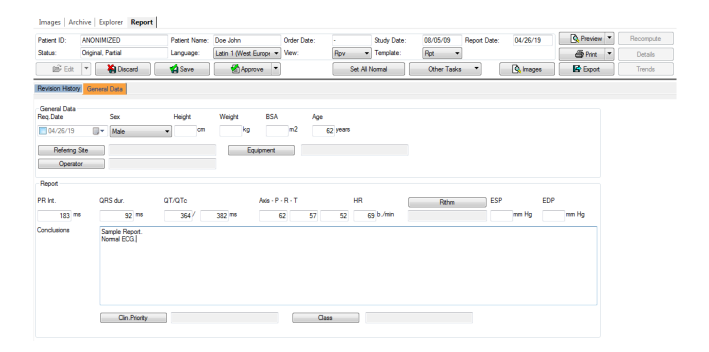
22.3. REPORTING

The ECG preliminary report, imported from the ECG device, is displayed on the right hand side of the ECG traces.

Depending on the ECG system used, this portion can either be empty or contain automated diagnosis performed by the software on board of the device or by centralized automated diagnosis software (for example by HES – Hannover EKG System)

The program does not perform any control on the clinical validity of the imported or computed automated diagnosis. The operator must verify its validity.

By clicking on the *Report* button, a new report will be created or, if a report already exists, such report will be opened in editing mode.



The *Approve* button allows you to generate and approve the report based on the imported interpretation and the default signature method for the selected template. The associated drop-down menu allows the user selection of the signature method.

Please refer to the "Reporting" Chapter for more information.

22.4. COMPARING

The Compare button allows you to compare different ECG traces for the same patient, if such traces have all been previously opened for review.

A set of additional controls will be displayed.



Each drop down control allows the selection of the specific ECG study and sequence to be compared. Up to four different traces can be selected.

The screen will be divided vertically in different sections to reflect the different ECG traces.

The Overlay button, if pressed, allows to overlay the traces being compared with each trace shown with a different color.

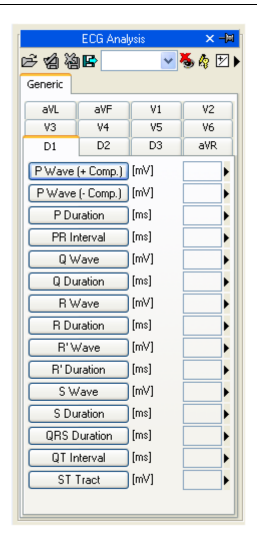
The Complexes button, if pressed, compares the different complexes if present.

22.5. OFF-LINE MEASUREMENTS

The ECG module provides the possibility of performing measurements on DICOM ECG objects.

Measurements are available in the Generic tab of the ECG Analysis window.

Generic measurements are only temporary. They are not stored to the ComPACS database and cannot be retained.



To access a more advanced or customized measurement panel, open either an already existing DICOM SR ECG report or create a new one. This operation can be performed by clicking on the toolbar icon as described in the "Managing Evidence and Report" chapter.

The analysis window will be extended with an additional *Structured* tab containing all the structured measurements. The appearance and content of the tab will depend on the reporting templates and analysis views configured by the system administrator for the specific workstation or user.



Structured measurements are stored in the DICOM SR object they refer to.

In this interface, only the windows corresponding to the data present in the open object will be filled (with numbers). To edit the opened report click the *Editing* icon:



To close and save the changes made click on the *Close and Save changes* icon:



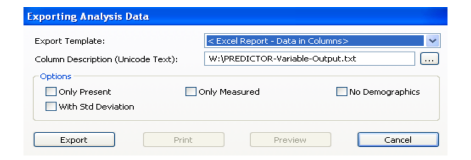
To close without saving any changes click on the Close discarding changes icon:



To export the data it is necessary to click the Export the report data to file icon:



An interface will be opened where you may select the data format and the file destination.



In the drop-down list, it is also possible to select an export template based on the installed configurations.

This functionality has been extended to export not only numerical values but also other field types (text, findings, dates, etc.). Furthermore the export also includes nested and repeated record structures.

The template is a simple text file which lists, in separate lines, the fields which need to be exported. Each line has the following format:

FieldID; Label - Optional; Unit - Optional

If the label or the unit are not specified, the default values shall be used.

For repeated records, the list of fields need to be enclosed in the following statement:

BEGINGROUP(Record Field ID); Label – Optional; Index – Optional ... list of fields ... ENDGROUP(Record Field ID)

The index is optional and, if omitted, all records shall be exported. Finally some special fields consist in DICOM fields. They are identified as follows:

\$(Group, Element); Label - Optional;

22.5.1 ECG TRACE CALIBRATION

The calibration information is contained in the DICOM ECG file. The product exploits it to avoid a manual calibration and ensure greater accuracy. Therefore calibrating ECG files in order to perform measurements is not required.

22.5.2 PERFORMING MEASUREMENTS

Now we are ready to perform some offline measurements on the ECG traces.

Once the ECG object is chosen, select the value to be measured on the measurement panel. In particular there are two panels: the *Generic* panel, which is always available, and the *Structured* panel which depends on the configuration set up and requires the opening of a report (intended as DICOM SR object).

The generic measurements are only temporary and will not be stored to a DICOM SR Object.

By clicking on the item, the mouse pointer becomes a pen. With a few mouse button clicks on the ECG trace, the software traces the measurement (in blue) and saves it to the ComPACS Database in the DICOM SR format. The following picture shows an example of time measurement:



To proceed, first select the desired measurement by clicking the chosen button in the control panel. Move the cursor to the starting point of the desired measurement and left-click once, the cursor will become a pen. Move the cursor to the end point of the measurement and left clicking once will terminate the measurement and populate the result in the chosen location.

22.5.3 CALCULATIONS AND FORMULAS

The software not only allows performing direct measurements, but it is also capable of computing derived parameters by use of different mathematical formulas.



In the absence of a correct calibration all measurements on the images are invalid.

The program does not perform any control on the consistency of the calibration factor. It is <u>always</u> necessary to check the calculated calibration factor to verify whether it is admissible.

The program does not perform any control on the clinical validity of the measurement. The operator must visually verify the validity of any measurements taken.

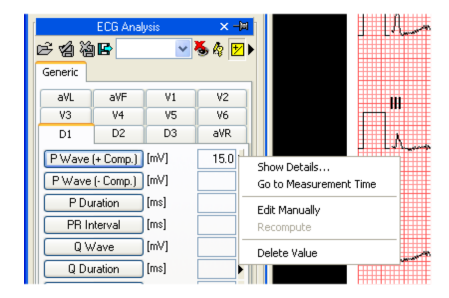
The product allows for detailed management of the measurements taken.

When a measurement is already present and re-executed, the software, by default, replaces the old value with the new one.

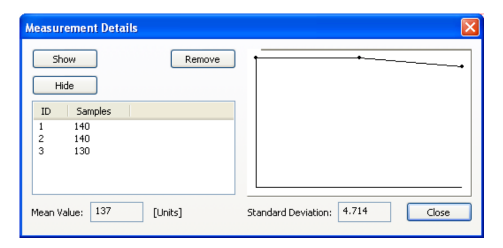
However more than one measurement on the same sample (or on various samples) can be performed and stored, calculating the mean value and the standard deviation. To perform multiple measurements and display the mean value click the button shown below. If this mode is active the button will turn yellow.



In the case of repeated measurements, the value shown nearby the item text is that of the mathematical average. However it is still possible to see all the measured samples. Alongside of every measurement is an arrow which allows accessing a management menu for the item.



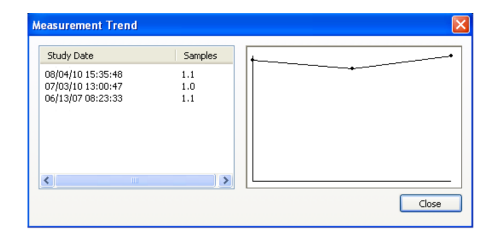
By clicking on the little arrow and then on *Show Details*, a window will be displayed in which all the samples taken for that item are shown together with the average and the standard deviation.



It is possible to delete a measured value if it seems incorrect. Simply select it and click on the *Remove* button. Use the buttons *Show* and *Hide* to show or hide the measurements on the image.

If ECG analysis has been performed on the same patient in previous studies stored in the ComPACS database, you may view how a specific measurement field has varied over time.

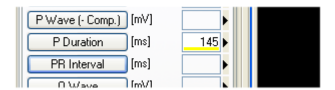
This operation can be performed by clicking on the *Show Trend* item in the menu. A window will be displayed in which the different values over time for the identified measurement are displayed both numerically and in a graphical format.



Another useful function is the ability to immediately display the ECG trace from which the measurement was taken. From the measurement menu, when clicking on the *Go to measurement time* option, the software automatically selects the trace on which the measurement was performed.

The feature *Go to measurement time* is available only for off-line measurements taken with ComPACS ECG.

Also by clicking on the *Edit Manually* button it is possible to manually insert the value instead of measuring it or calculating it. This value will be underlined in yellow to indicate it is a manually inserted measurement.



To show/hide the measurements on the images click on this icon:



To fix the measurement graphics on the display:



To close and save the changes click on the *Close and Save* icon:

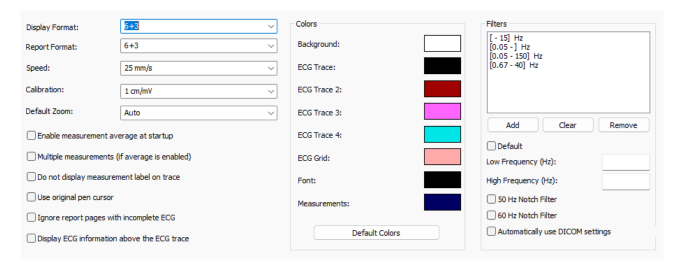


To close without saving the changes click on the *Close Discarding changes* icon:



22.7. CUSTOM SETTINGS

This component provides different user configurable settings, located in the *Program Settings* window, under the *ECG* tab.



The default display format, report format, time scale, calibration and zoom to use for review and reporting can be indicated.

"Enable measurement average at startup" automatically enables the average function when the component is activated.

"Multiple measurements (if average is enabled)" allows the user to measure a specific field multiple times, without having to press again the corresponding button. The measurement sequence can be interrupted by pressing the right mouse button.

"Do not display measurement label on trace" does not display the label and the measured value as an overlay on the ECG trace.

"Use original pen cursor" uses the pen cursor of previous software revisions. Starting from 10.8.6 the pen cursor has been replaced with a sight view cursor.

"Ignore report pages with incomplete ECG" does not include in the ECG report any additional page in which the ECG samples do not cover the entire page timeline.

"Display ECG Information above the ECG trace" displays the information section above the ECG trace instead of the default right position.

It is also possible to customize the colors used to display the ECG traces and the list of post-processing filters available to users.

23. REPORTING

23.1. ACTIVATING COMPACS REPORTING

To be able to create or open a report, it is first necessary to adequately configure your system. Please refer to your system administrator and to the *System Administrator's manuals* for more details.

If ComPACS Reporting is enabled on your system, the following button will be visible in the toolbar, on the right hand side:



If the button is pressed then the module is already active. Otherwise it can be activated by pressing the button with the mouse.

Upon activation, the module will request a runtime license from the ComPACS Server. If the license cannot be received, then the module will fail to activate and an error message will be displayed.

In case of concurrent licenses, it is possible that all available licenses have already been requested and assigned to other users that are using the module at that time and that no licenses are left for activation. This is a likely cause for failure.

Activation failure can also occur if the network connection to the server is no longer available.

Please ensure that the module is released by releasing the button above, after you have finished using the product so that the license is also released for use by other users.

When the module is active each time a report is opened or created a reporting window will be displayed.

23.2. CREATING OR OPENING A REPORT

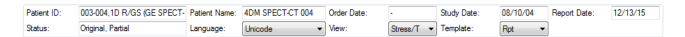
In order to create or open a report, a study first needs to be opened for review.

Then, press the report button on the toolbar $2 = 10 \times 4$. There can be different situations:

- a) There are no reports for the selected study. A new report is created. A dialog window will allow you to select all the necessary parameters.
- b) Only one report exists for the selected study and the report type is automatically identified. The report is opened for editing. Please note that obsolete reports (i.e. reports that are marked as predecessor of other reports) are not included in the report count.
- c) Only one report exists for the selected study but the report type cannot be automatically identified. A dialog window will ask the user to associate the report to one of the configured templates.
- d) More reports exist for the selected study. A dialog window will be displayed, asking the user to confirm the creation of a new report and the importing of the previous reports.

Please refer to the "Managing Evidence and Reports" chapter for more detailed information.

Once the report is opened, a new *Report* tab window will be opened in the monitor identified for this operation. At the top of the window, a brief report summary will be presented.



The summary will present the patient ID, name, order date, study date, report date, report status (if modified, verified, finalized, etc.), language (character set used) and the Reporting Views and Print Templates currently selected.

The Reporting Views and Print Templates can also be changed dynamically, similarly to the character set used, simply by selecting from the drop down menus displayed above. The other fields are read only and managed directly by the software.

Newly generated reports will be opened directly in *Edit* mode. Existing reports are instead opened by

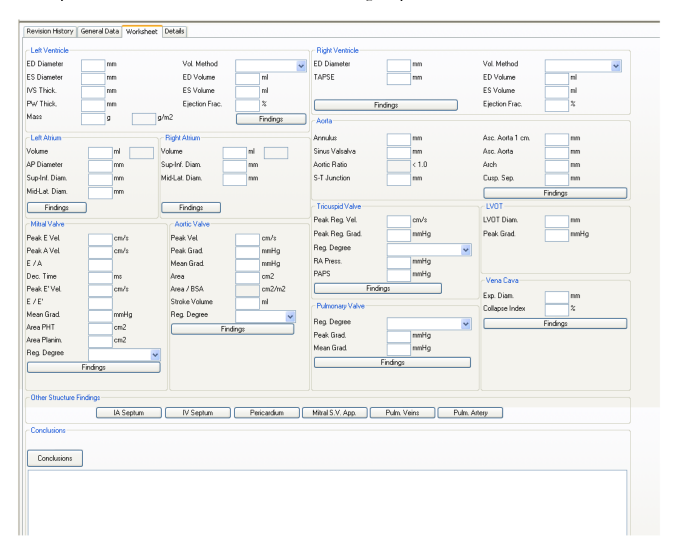
default in *Viewing Only* mode. To edit them, you simply need to press the have sufficient privileges to perform this operation on the selected report.

If a report has already been approved and signed, it is no longer possible to edit its content. A new cancellation report or replacement reports needs to be created, by clicking the arrow to the left of the button and selecting the appropriate item from the drop down menu:



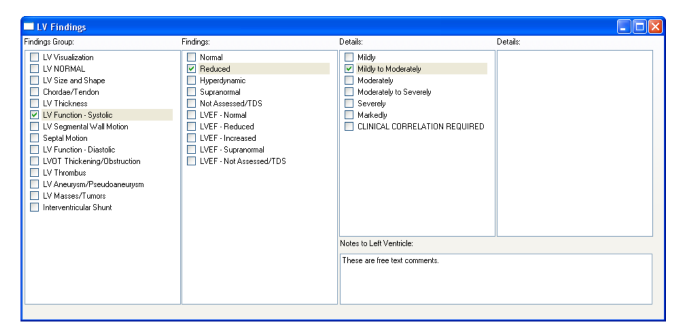
Once a cancellation report is approved, it is no longer possible to continue the same reporting chain. A new chain of reports has to be created in such scenario.

Below you will see a number of sub-windows, also arranged by tabs.

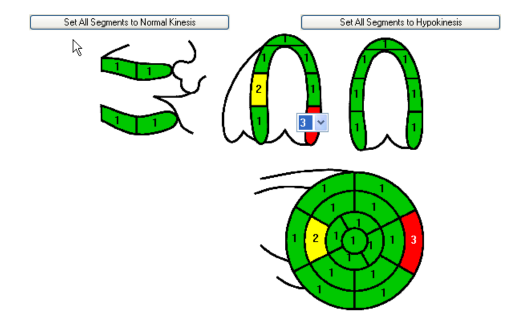


The first tab is always present and conveys the *Revision History* information about the report. The layout of the remaining tabs varies depending on the specific template and view selected. A possible example is depicted in the following picture.

In the example, you can see a worksheet with different measurement fields and with buttons named after anatomical structures. By pressing these buttons, pop-up windows will be displayed to allow structured reporting selections.

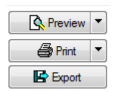


Some boxes for free text or graphical items can also be present.

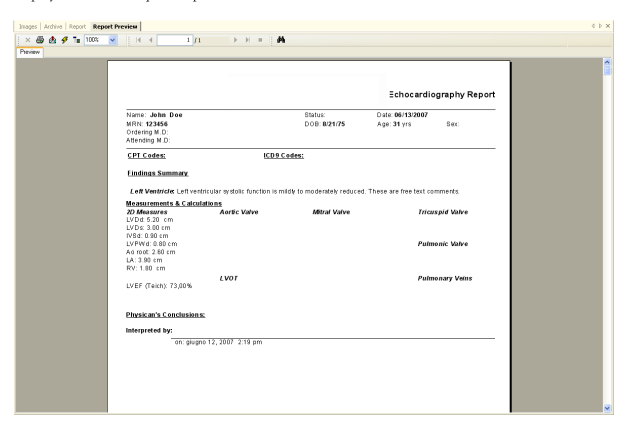


The structure of the report menus depends entirely on the template and the related *Context Groups* and *Reporting Views* selected. What is presented in this guide is just a possible example.

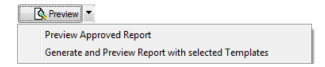
23.3. PREVIEWING AND PRINTING THE REPORT



It is possible to display a preview of the report by pressing the *Preview* button. An additional tab will be displayed with the requested preview.



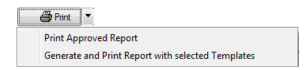
If the report has been approved, the preview will display the approved PDF document. Instead, if it is being created or edited, the preview will be dynamically updated every time some changes are made to the structured reporting selections or in the measurements.



It is also possible to display the preview of the approved report, re-generated with a different set of templates by selecting the related item in the button drop down menu.

In a multi-monitor setting, the preview could be shown side by side to the structured reporting view.

The *Print* button allows you print the report to a system printer. For approved reports the PDF document shall be printed. For non-approved reports, the selected templates will be used for report generation and printing.



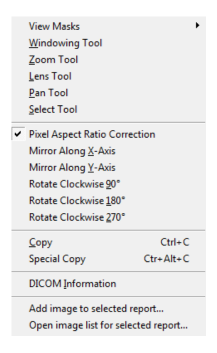
Also in this case it is possible to print the approved report, re-generated with a different set of templates by selecting the related item in the button drop down menu.

The *Export* button regenerates and exports the report to common electronic formats such as *PDF*, *RTF*, *HTML*, *Excel*, etc.

23.4. ADVANCED OPERATIONS

23.4.1 ADDING IMAGES

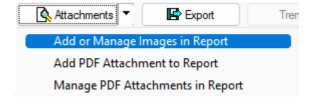
When a report is open for editing, it is possible to add images to the report by right clicking on the image window and selecting "Add image to selected report..." from the context menu:



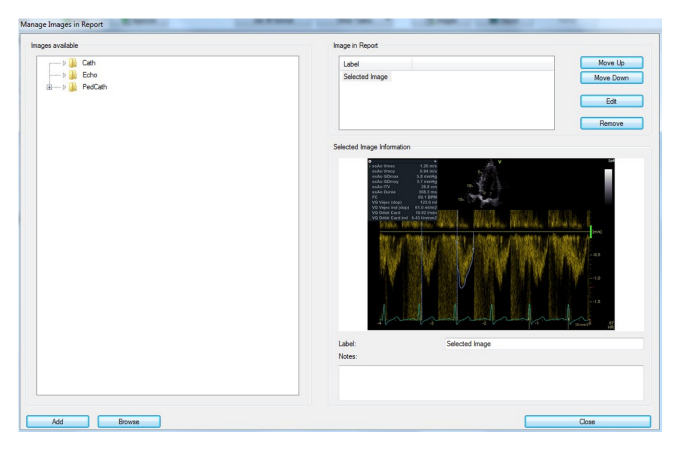
A dialog will be displayed to confirm the image selection:



It will be possible to add the entire view currently displayed, just the selected image or simply a region of interest. The added image will also require a label and optional notes.



From the same context menu or using the above context menu in the report view, it is possible to review the images present in the report, re-order them or remove them from the report.



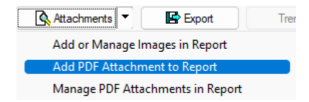
In this window it is also possible to add images from a file library or browse for specific images to add. The file library is built dynamically from the "ReportImages" folder, located in the parent folder of the application executable.

By pressing the button it is also possible to edit the image using the operating system default image editor.

The reporting templates also need to modify to reflect how the report images will be displayed in the report printout.

23.4.2 ADDING ATTACHMENTS

When a report is open for editing, it is possible to add attachments to the report by clicking on the following context menu:



The user can then select the attachment in the dialog window. It is possible to select attachments from DICOM Encapsulated PDF objects in the same study or from external PDF files. Optionally only a partial set of pages can be attached.

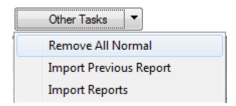


From the same button or context menu, it is possible to review the attachments present in the report, re-order them or remove them from the report.

23.4.3 ALL NORMAL STATEMENTS

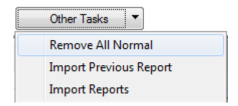
Reports for normal patients can be created quickly using the button which sets the default normal values for the report being edited.

It is also possible to remove such selection by using the "Other Tasks" drop down menu and selecting "Remove All Normal" item.



23.4.4 IMPORTING PREVIOUS REPORTS

Using the "Other Tasks" drop down menu and selecting "Import Previous Reports" or "Import Reports" items it is possible to import a report a previous study of the same patient in order to avoid entering a similar diagnosis.

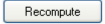


If the "Import Previous Report" item is selected, the system will automatically import the most recent compatible (of the same type) report.

If the "Import Reports" item is selected a dialog will be displayed to allow the manual selection of the report to import.

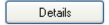
The specific fields to be imported are to be configured by an application specialist.

23.4.5 RECOMPUTING VALUES

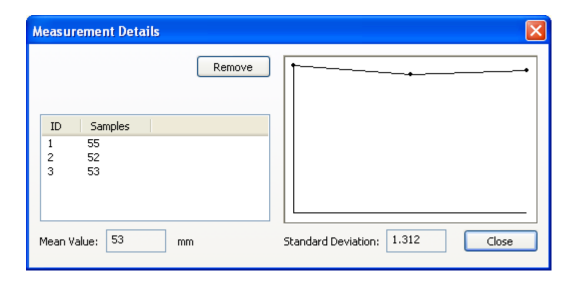


The Recompute button is activated when the mouse cursor is positioned on a measurement field with an associated formula. By clicking the button, the formula will be executed and, if applicable, the selected field value updated.

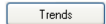
23.4.6 MEASUREMENT DETAILS



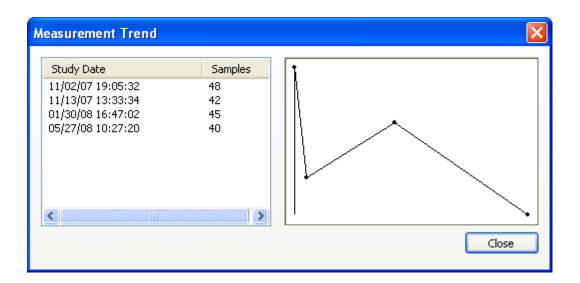
The Details button is activated when the mouse cursor is positioned on a measurement field. By clicking the button, a window will display all the different measured or imported samples for the selected field, their mean value and the standard deviation.



23.4.7 MEASUREMENT TRENDS



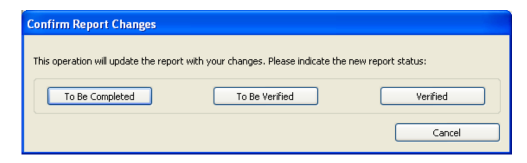
The Trends button is activated when the mouse cursor is positioned on a measurement field. By clicking the button, a window will display the trend over time for the selected field, considering all previous reports for the selected patient.



23.5. CONFIRMING OR DISCARDING CHANGES



If in Edit mode, you need to decide what to do with the changes applied before closing the report. If you want to discard the changes made since the last edit operation or just discard a new report, simply press the Discard button. If instead you want to save the changes but leave the report open for editing, press the Save button; a dialog will prompt you for additional information.



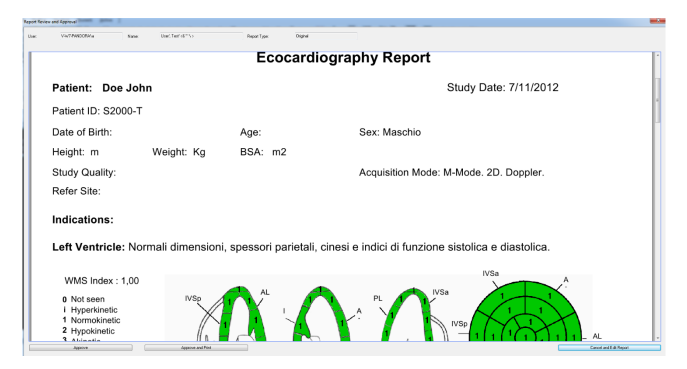
Please press the button to confirm your choice.

To Be Completed	- This is used if you need to temporarily stop the reporting process in the middle of a non complete report.
To Be Verified	 This is used if the report is complete but you do not have the authority to verify/validate its content from a medical standpoint.
Verified	- This is used if the report is complete and is to be verified from a medical standpoint. The report can still be edited pending final approval by an authorized user.

Finally, if you want approve and sign the report, press the *Approve* button to sign using the default signature method associated to the report template or use the associated drop-down menu to select a specific signature method.

ComPACS default signature method is the Electronic Signature, i.e. username and password. However, different custom signature method plugins, such as smart-card based digital signature, are available.

A window will display the final, signed, report in PDF format for your review and final confirmation.



Information about the user currently logged on, his or her name and the type of report (Original, Replacement or Cancellation) is also provided.

Depending on your privileges, it is possible that only a subset of these options is available to you.



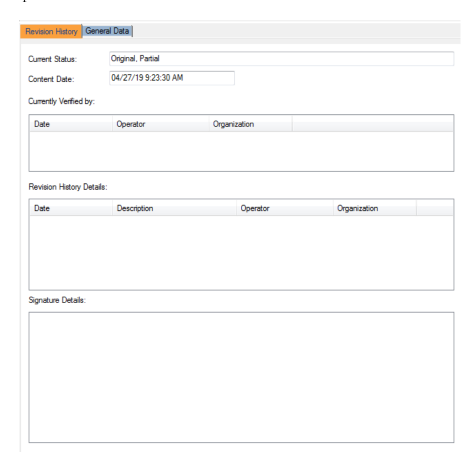
After Approval no further changes are allowed on the report which is definitively archived.

Upon approval, the system generates a PDF version of the report with the print template in use at the time of finalization. This PDF file, encapsulated in a DICOM object, is stored for future reference in the exact content and layout.

If the print templates are modified, the report content and layout in preview may change even after finalization. It is important to always reference the PDF version for the content and layout of the report issued.

23.6. REVISION HISTORY

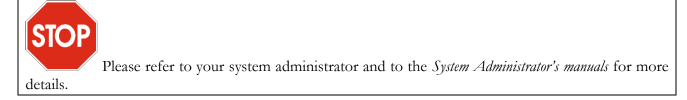
The revision history tab is common to all templates and shows the current status of the report as well as the history of the changes that occurred, the user responsible for such changes and the date/time at which they took place.



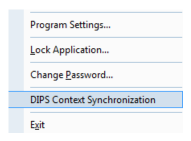
If a digital signature has been applied to the report document, the details about the signature process as well as possible additional information, is also listed.

24. CONTEXT SYNCHRONIZATION

ComPACS Workstation can be synchronized with a compatible third party system. This allows the application to automatically change patient context if such changes occur in a third party system or vice-versa.



If this capability is enabled, a context synchronization item will be displayed in the *File* menu, before the *Exit* item.



If this capability is active, the menu item will be checked. If unchecked, the workstation will not synchronize with the third party application.

When the menu item is selected, the synchronization status is modified.

Warning messages will be provided when the workstation loses context synchronization or when it is not capable of changing patient context.



25. APPENDIX A

25.1. POST MARKETING SURVEILLANCE



This software is subject to post marketing surveillance in order to detect or become aware of, as rapidly as possible, real or potential hazards for the patient and/or operator due to incorrect functioning of the software and consequently resolving such problems in the fastest and most efficient way.

For this reason, if the operator notices any incorrect functioning of the software, reduction or limitation in its performance, inaccuracy or inadequacy of the labeling or operator's manuals, that may lead to a real or potential hazard for the patient and/or operator, we kindly request to be immediately informed directly or through our representatives by filling in the following form or sending a communication conveying the same information. All the information related to the software (name, version, license code) are available in the *about window* of the program and in the installation CD. In this way, we can undertake all adequate preventive and corrective actions with the greatest efficiency and speed.

25.2. Hazard Notification Form

Send to:

MEDIMATIC S.R.L.

Quality Assurance Function
Viale Cembrano 4C
I-16148 Genova ITALY
E-mail: <u>regulatory@medimatic.com</u> , Certified E-mail: <u>medimatic@legalmail.it</u>
oftware Name:
oftware Name:
ntegrated Module(s) Name(s):
icense Code(S/N):
Description of the real/potential hazards:
Notes and suggestions:
Person/Department to Contact:
Address:
Phone: Fax:
E-Mail:
Date: Signature:



26. APPENDIX B

26.1. ADDITIONAL ARCHIVES

The login configuration defines the main archive. In addition, secondary archives per each configuration can be configured using the *Program Settings* item of the *File* menu and going to the *Archive* tab. This way, ComPACS can perform simultaneous queries on multiple archives instead of a single one.

The first choice is to determine if this configuration is valid only for the current workstation (identified by its *Workstation AE* Title) or to *All others*, meaning all the workstations for which no dedicated configuration has been defined. The *Add Workstation* and *Delete Workstation* respectively allow the creation or removal of dedicated workstation settings.

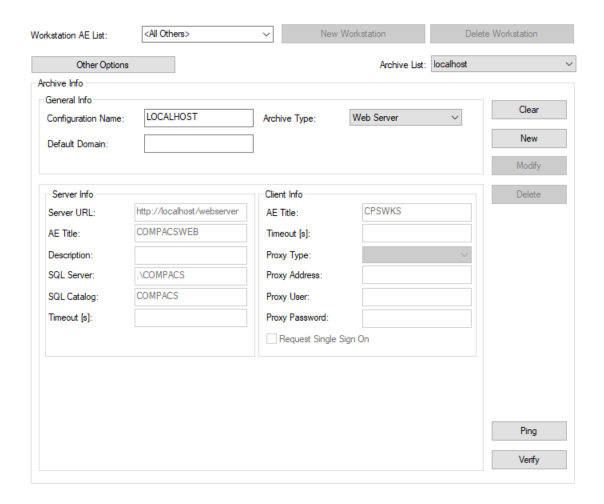
Then in this window you need to initially assign the following parameters:

Configuration Name - A name identifying this specific archive configuration

Archive Type - The type of archive you are connecting to.

Default Domain - The default domain the users logging into this archive belong to.

Default Archive - If this is the default configuration to be prompted upon startup.



There currently two different *Archive Types* available: a) archives which represent ComPACS systems Web Server and, b) Third party archives accessible via DICOM network protocol (DICOM).

To connect to a ComPACS archive you need to specify the following additional information:

Server URL	- This is the URL of the Web Server. Typically http:// <machinename>/webserver.</machinename>
AE Title	- The AE Title of the Server
Description	- A free text description (does not need to match the server side description).
SQL Server	- The name of the Microsoft® SQL Server instance on which the ComPACS database is installed.
SQL Catalog	- The name of the catalog within the above SQL server in which the database is installed.
Timeout	- The default value automatically prompted is recommended.

and for the Client:

AE Title - The AE title of the workstation. This needs to have been already

configured with the Web & Server Tools.

Proxy Type - The type of proxy used.

Request Single Sign On - This parameter can be checked if the workstation is to automatically

authenticate the user on the basis of the current Windows logged on

user.

To connect to a DICOM archive you need to specify the following additional information:

Host Name/IP - dns name or IP of the server on which the DICOM Server is running.

AE Title - The AE Title of the DICOM Server

Description - A free text description.

Port - The port on which the server is listening for incoming connections.

Timeout - The association timeout that the server allows for a connection

(expressed in seconds).

Max PDU - The maximum PDU size that the server expects. The value is

expressed in Kb.

<DICOM Prop.> - Check boxes for the different DICOM services that the server

provides.

< Advanced > - Advanced DICOM settings, recompression formats for C-STORE

operations and logging functions.

And for the *Client*:

AE Title - The AE Title of the DICOM Server

Port - The port on which the server is listening for incoming connections.

Timeout - The association timeout in attempting a connection (expressed in

seconds).

Max PDU - The maximum PDU size to propose in DICOM associations. The

value is expressed in Kb.



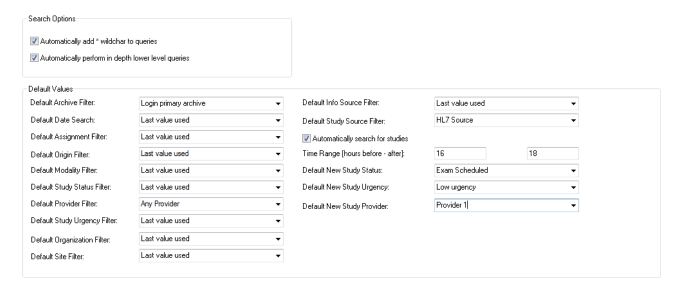
set.

The Verify button allows you to perform a C-ECHO to see if the parameters are correctly

Once set, apply the parameters by clicking the *New* button. The software will put the registered archive in the field labeled *Archive List*. This method allows you to add several archives or to edit already defined ones.

26.2. OTHER OPTIONS

The Other Options button allows the configuration of the default archive settings.



It is possible to indicate whether wildcard characters are to be automatically added to search fields. Please note that this feature does not apply to ID fields such as Patient ID or Accession Number.

It is also possible to indicate the default filtering values for the Archive Search window, the Information Source provider to use and the default provider and status for new created studies. For most fields the system can also retain the last settings used for each user.



27. APPENDIX C

27.1. ECHO ANALYSIS ACCURACY AND PRECISION

In this appendix the terms accuracy and precision have the following meaning:

Accuracy - the absolute value of the difference between the real measure and the mean of the estimated measurements;

Precision - the standard deviation of the estimated measurements.

The following table reports the mean values and precision of an experiment in which repeated measurements of geometrical artificial structures in an ultrasound DICOM image.

Measurement	Accuracy	Precision	Calibration factor
Linear measurements	0.25 mm	0.83 mm	1 mm /pixel
Polygonal Areas	0.28 mm^2	1.02 mm^2	1 mm /pixel
Velocity	0.001 m/s	0.007 m/s	0.1 m/s / pixel
Time	0.1 ms	0.40 ms	0.5 ms /pixel

In the absence of a correct calibration, the measurements carried out on the images are invalid. It is always necessary to control the calculated calibration factor, the obtained measure and the borders traced to verify their admissibility.

The accuracy and precision depend on the user pixel selection and consequently on the image calibration factor. Generally for the measurements we suggest to apply a zoom factor on the image to reduce the effect of the introduced mistake from the manual indication of the distance to be measured.

Velocities are generally measured with a single click. This justifies the greater accuracy and precision.



28. APPENDIX D

28.1. ANGIOGRAPHIC ANALYSIS ACCURACY AND PRECISION

In this appendix the terms accuracy and precision have the following meaning:

Accuracy - the absolute value of the difference between the real measure and the mean of the estimated measurements;

Precision - the standard deviation of the estimated measurements.

28.2. KNOWN DISTANCE BASED CALIBRATION

The following table reports the mean values and precision of an experiment in which repeated measurements of known distances of different dimensions have been carried out on the same image. The table underlines a relation between the size of the measured structure in the image and the precision of the measurement; when the size in pixels increases (in the experiment this also corresponds to an increase in the real dimension) the measurement precision increases.

Square Side Length	Accuracy	Precision
4 mm	0.0004 mm/pixel	0.00196 mm/pixel
8 mm	0.0004 mm/pixel	0.001497 mm/pixel
12 mm	0.0004 mm/pixel	0.001497 mm/pixel
16 mm	0.0004 mm/pixel	0.001497 mm/pixel
20 mm	0.0004 mm/pixel	0.001497 mm/pixel
40 mm	0.0004 mm/pixel	0.0008 mm/pixel

For this reason it is better to carry out calibration on structures with sufficiently large pixel dimensions (dimensions in the image to be analyzed). Alternatively it is possible to apply a zoom factor to the image to reduce the error introduced by the manual selection of the distance to be measured.

28.3. CATHETER BASED CALIBRATION

The following table reports the results of a calibration experiment on a set of contrast filled catheters of different size and vendor.

Contrast filled		
Accuracy Precision		
0.0024 mm	0.0009 mm	

28.4. CORONARY ANALYSIS

The following table reports the results of an in vitro analysis of a set of phantoms of different diameters:

Dimensions	X-ray Images		
Size	Accuracy	Precision	
	mm	mm	
5.0 mm	0.00143	0.01349	
4.5 mm	0.00684	0.02083	
4.0 mm	0.00245	0.01854	
3.5 mm	0.00274	0.02282	
3.0 mm	0.00391	0.02812	
2.5 mm	0.00001	0.02370	
2.0 mm	0.00066	0.02994	
1.5 mm	0.00336	0.03547	
Mean Value	0.003905	0.00724	

28.5. VENTRICULAR VOLUME

The following table reports the results obtained by using the program to measure an elliptic synthetic image in order to estimate the error due to the manual profile tracing:

	Accuracy	Precision
Diastolic Volume	0.97 ml	0.9618 ml
Systolic Volume	0.19 ml	0.3928 ml

It is clear that the errors are negligible not only with respect to the measure carried out, but also with respect to the intrinsic errors in the applied *Area-Length* method for volume calculation.

28.6. BODY SURFACE

The formula used to compute the body surface is:

$$BSA = 71.84 \cdot W^{0.425} \cdot H^{0.725}$$

where W corresponds to the *weight* in Kg, H corresponds to the Height in cm and the calculated body surface is cm^2 .

28.7. REGIONAL SHORTENING

The following table shows the error in the detection of the regional shortenings on a synthetic image:

	Accuracy	Precision
Regional Shortening	0.3922 %	0.7072 %

In the absence of a correct calibration, the measurements carried out on the images are invalid. It is always necessary to control the calculated calibration factor, the obtained measure and the borders traced to verify their admissibility.



29. APPENDIX E

29.1. IVUS ANALYSIS ACCURACY AND PRECISION

In this appendix the terms accuracy and precision have the following meaning:

Accuracy - the absolute value of the difference between the real measure and the mean of the estimated measures;

Precision - the standard deviation of the estimated measures.

The following table reports the mean values and precisions of an experiment in which repeated measures of known distances of different dimensions have been carried out on the DICOM file.

Measurement	Accuracy	Precision
Horizontal segment	0.01 mm	0.030 mm
Vertical segment	0.01 mm	0.030 mm
Diagonal segment	0.03 mm	0.045 mm
Areas	0.05 mm^2	0.086 mm^2

For these measurements it is suggested to apply a zoom factor to the image to reduce the error introduced by the manual selection of the distance to be measured.

The following table reports the mean values and precisions of an experiment in which repeated measures of known distances and areas into the image longitudinal reconstructed.

Measurement	Accuracy	Precision
Horizontal segment	0.09 mm	0.03 mm
Vertical segment	0.01 mm	0.03 mm
Diagonal segment	0.09 mm	0.03 mm
Areas	0.02 mm^2	0.06 mm^2



30. APPENDIX F

30.1. GENERIC MEASUREMENTS ACCURACY AND PRECISION

In this appendix the terms accuracy and precision have the following meaning:

Accuracy - the absolute value of the difference between the real measure and the mean of the estimated measurements;

Precision - the standard deviation of the estimated measurements.

The following table reports the mean values and precision of an experiment in which repeated measurements of geometrical artificial structures in an ultrasound DICOM image.

Measurement	Accuracy	Precision	Calibration factor
Linear	0.33 mm	0.723 mm	1 mm / pixel
Polygonal Area	0.28 mm^2	1.020 mm ²	1 mm / pixel
Angle	0.04 °	0.126 °	

In the absence of a correct calibration, the measurements carried out on the images are invalid. It is always necessary to control the calculated calibration factor, the obtained measure and the borders traced to verify their admissibility.

The accuracy and precision depend on the user pixel selection and consequently on the image calibration factor. Generally for the measurements we suggest to apply a zoom factor on the image to reduce the effect of the introduced mistake from the manual indication of the distance to be measured.