RAYCARE 5A

IHE-RO Integration Statement

Declaration of conformity



 $Complies with 93/42/EEC \, Medical \, Device \, Directive \, as \, amended \, by \, M1 \, to \, M5. \, A \, copy \, of the \, corresponding \, Declaration \, of \, Conformity \, is \, available \, on \, request.$

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INTEGRATION STATEMENT

1.1 INTRODUCTION

This IHE Integration Statement is prepared and published to describe the conformance of RayCare with the IHE-RO Technical Framework. It identifies specific IHE-RO capabilities that are supported in terms of actors and integration profiles described in the technical frameworks of each domain.

Users familiar with these concepts can use Integration Statements to determine the integration of RayCare with complementary systems and what clinical and operational benefits such integration might provide. This Integration Statements is intended to be used in conjunction with the RayCare DICOM Conformance Statements.

1.2 PRODUCT VERSION

This Integration Statement is valid for RayCare 5A.

1.3 ABOUT THE TESTING PROCESS

IHE provides a process for vendors to test their implementations of IHE actors and integration profiles. The IHE testing process, culminating in a multi-party interactive testing event called the Connect-a-thon, provides vendors with valuable feedback and provides a baseline indication of the conformance of their implementations. The process is not intended to independently evaluate, or ensure, product compliance. In publishing the results of the Connect-a-thon and facilitating access to vendors' IHE Integration Statements, IHE and its sponsoring organizations are in no way attesting to the accuracy or validity of any vendor's IHE Integration Statements or any other claims by vendors regarding their products.

RaySearch Laboratories has done further validation beyond the IHE-RO Connect-a-thon to ensure that RayCare conforms to applicable standards. This Integration Statement shall not be used a guarantee that RayCare will work in any environment or with any external system.

1.4 REFERENCES

Reference	Resource
DICOM	https://www.dicomstandard.org/
IHE	https://www.ihe.net/
IHE-RO	https://www.ihe.net/ihe_domains/radiation_oncology/

1.5 DEFINITIONS

Term	Meaning		
CE	CE marking is a certification mark that indicates conformity with health, safety, and environmental protection standards for products sold within the European Economic Area (EEA)		
DICOM	Digital Imaging and Communications in Medicine		
EEC	European Economic Community		
IHE	Integrating the Healthcare Enterprise		
IHE-RO	Integrating the Healthcare Enterprise Radiation Oncology		
TDW-II	Treatment Delivery Workflow-II (IHE-RO Profile)		

1.6 INTEGRATION STATEMENT

IHE-Integration Statement	Date	13 September 2021					
Vendor	Product Name	Version					
RaySearch Laboratories	RayCare	5A					
This product implements all transactions required in the IHE Technical Framework to support the IHE Integration Profiles, Actors and Options listed below:							
Integration Profiles Implemented	Actors Implemented	Options Implemented					
TDW-II	Treatment Management System	Retain Original Treatment Records					
	Object Storage	Retain Original Treatment Records					
Links to Standards Conformance Statements for the Implementation							
DICOM IHE	https://www.raysearchlabs.com/product-configurations/						
Links to general information on IHE							
In North America: www.ihe.het	In Europe: www.ihe-europe.org	In Japan: www.jira-net.or.jp/ihe-j					



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