Foreword

This is a supplement to the IHE Radiation Oncology Technical Framework. Each supplement undergoes a process of public comment and trial implementation before being incorporated into the volumes of the Technical Frameworks.

This supplement is published on November 10, 2023 for Public Comment. Comments are invited and can be submitted at https://www.ihe.net/Radiation_Oncology_Public_Comments. In order to be considered in development of the Trial Implementation version of the supplement, comments must be received by December 11, 2023.

This supplement describes changes to the existing technical framework documents.

“Boxed” instructions like the sample below indicate to the Volume Editor how to integrate the relevant section(s) into the relevant Technical Framework volume.

Amend Section X.X by the following:

Where the amendment adds text, make the added text bold underline. Where the amendment removes text, make the removed text bold strikethrough. When entire new sections are added, introduce with editor’s instructions to “add new text” or similar, which for readability are not bolded or underlined.

General information about IHE can be found at IHE.net. Information about the IHE Radiation Oncology domain can be found at IHE Domains. Information about the organization of IHE Technical Frameworks and Supplements and the process used to create them can be found at Profiles and IHE Process. The current version of the Radiation Oncology Technical Framework can be found at Radiation Oncology Technical Framework.
CONTENTS

55
Introduction to this Supplement ................................................................. 5
History of Changes .................................................................................... 5
Open Issues and Questions ....................................................................... 5
Closed Issues ......................................................................................... 6
60 IHE Technical Frameworks General Introduction ........................................... 7
9 Copyright Licenses .................................................................................. 7
10 Trademark ............................................................................................. 7
IHE Technical Frameworks General Introduction Appendices ....................... 8
Appendix A – Actors .................................................................................. 8
65 Appendix B – Transactions ........................................................................ 8
Appendix D – Glossary ............................................................................... 9

Volume 1 – Profiles .................................................................................. 10
X Brachy Treatment Delivery– Record Content (TDRC-Brachy) Profile .............. 10
X.1 TDRC-Brachy Actors, Transactions, and Content Modules ...................... 11
70 X.1.1 Actor Descriptions and Actor Profile Requirements ......................... 11
X.2 TDRC-Brachy Actor Options ............................................................... 11
X.3 TDRC-Brachy Required Actor Groupings .............................................. 12
X.4 TDRC-Brachy Overview ....................................................................... 12
X.4.1 Concepts ........................................................................................ 12
X.4.2 Use Cases ....................................................................................... 12
X.5 TDRC-Brachy Security Considerations ................................................. 12
X.6 TDRC-Brachy Cross Profile Considerations ........................................... 12

Volume 2 – Transactions ........................................................................ 13
3.Y1 Brachytherapy Treatment Record Storage [TDRC-Brachy-01] .................... 13
80 3.Y1.1 Scope ......................................................................................... 13
3.Y1.2 Actor Roles .................................................................................. 13
3.Y1.3 Referenced Standards .................................................................... 13
3.Y1.4 Messages ..................................................................................... 14
Appendices to Volume 2 ............................................................................ 15

Volume 3 – Content Modules ................................................................. 16
5 IHE Namespaces, Concept Domains and Vocabularies ............................... 16
5.1 IHE Radiation Oncology Namespaces ................................................... 16
5.2 IHE Radiation Oncology Concept Domains ............................................ 16
5.3 IHE Radiation Oncology Format Codes and Vocabularies ....................... 16
90 5.3.1 IHE Format Codes ....................................................................... 16
5.3.2 IHEActCode Vocabulary ............................................................... 16
5.3.3 IHERoleCode Vocabulary .............................................................. 16
6 Radiation Oncology HL7 V3 CDA Content Modules .................................... 16
7 Radiation Oncology DICOM Content Definition ........................................ 17
95 7.1 Conventions ..................................................................................... 17
7.2 General Definitions ............................................................................................................ 18
7.3 IOD Definitions .................................................................................................................. 18
  7.3.6 Treatment Record IODs ............................................................................................. 18
    7.3.6.1 Technique Specific RT Treatment Record ......................................................... 18
    7.3.6.2 RT Treatment Record for General Use .............................................................. 18
    7.3.6.3 RT Brachy Treatment Record ............................................................................ 18
      7.3.6.3.1 RT Brachy Treatment Record IOD ........................................................... 18
        7.3.6.3.1.1 Referenced Standards ............................................................................... 18
        7.3.6.3.1.2 IOD Definition ................................................................................... 19
105  7.4 Module Definitions ........................................................................................................ 19
  7.4.1 General Modules ........................................................................................................ 19
  7.4.2 Workflow-Related Modules ....................................................................................... 19
  7.4.11 Treatment Records ................................................................................................... 20
    7.4.11.6 RT Brachy Treatment Record .......................................................................... 20
110  7.4.11.6.1 Referenced Standards ..................................................................................... 20
    7.4.11.6.2 RT Brachy Session Record Module Definition for HDR and PDR .............. 20
Appendices to Volume 3 ............................................................................................................... 32

Volume 4 – National Extensions .......................................................................................... 33
4 National Extensions ............................................................................................................. 33
Introduction to this Supplement

This profile defines the content of the treatment record being transferred from the Treatment Delivery Device, Treatment Management System or other RT Brachy Treatment Record storage or producer to an interested consumer. This content can be available as soon as patient delivery is completed, and other computation and information gathering, and formatting can be performed to support the creation of the RT Brachy Treatment Record.

This content profile is motivated by medical physicists working with brachytherapy planning systems, who face an increasing demand from patient-care, data-quality and research perspectives to increase the usefulness, exchangeability and availability of clinical data across the various treatment planning systems.

The main role of this profile is to address a solution for such interoperability using the DICOM RT objects provided in their 1st generation.

History of Changes

<table>
<thead>
<tr>
<th>Date</th>
<th>Rev.</th>
<th>Author</th>
<th>Change Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>November 2023</td>
<td>1.0</td>
<td>IHE RO Technical Committee</td>
<td>Initial public comment publication</td>
</tr>
</tbody>
</table>

Open Issues and Questions

<table>
<thead>
<tr>
<th>#</th>
<th>Open Issue Description</th>
</tr>
</thead>
</table>
| 6   | 2018-11-21 WG-07 Brachy Subgroup: Added LDR Treatment Record Storage [BWF-11].
No specification is yet written for the Treatment Record Storage for Seeds. It was consensus that at least a manual recording would be useful. The main purpose is recording for archiving / charge capture. The record would be written by a TPS/TMS application.
Added a module definition by 7.4.11.6.2 RT Brachy Session Record Module for Seed treatment. Currently rules are a copy form HRD. They have to be revised all over the place but kept for convenience.
Rebecca Park will prepare a proposal for the meeting as a starting point.                                                                                                                                                                                                                                                                               |
| 8   | 2018-11-21 WG-07 Brachy Subgroup: The use of DICOM Unified Worklist suggested in some use cases will not be covered in this Profile. The use case section shall state that the use case variations using worklist are included only for illustration of the clinical environment.
The Technical Committee should look into eventually integrating Brachytherapy into TDW II, since the framework is basically the same. Alternatively, a separate TDW-style profiles could be written for Brachytherapy.                                                                                                                                                                                                                       |
## Closed Issues

<table>
<thead>
<tr>
<th>#</th>
<th>Closed Issue Description/Resolution</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Profile name: Brachytherapy Workflow Profile.</td>
</tr>
<tr>
<td></td>
<td>DONE: Changed to TDRC-Brachy</td>
</tr>
<tr>
<td>2</td>
<td>Content definition (based on different techniques)</td>
</tr>
<tr>
<td>3</td>
<td>Definition of the general workflow</td>
</tr>
<tr>
<td></td>
<td>2015-12-10; Various use case scenarios are described in X.4.2</td>
</tr>
<tr>
<td>4</td>
<td>Definition of the list of transactions (see Appendices A and B, X.1 Actors, Transactions …)</td>
</tr>
</tbody>
</table>
IHE Technical Frameworks General Introduction

The IHE Technical Frameworks General Introduction is shared by all of the IHE domain technical frameworks. Each technical framework volume contains links to this document where appropriate.

9 Copyright Licenses

IHE technical documents refer to, and make use of, a number of standards developed and published by several standards development organizations. Please refer to the IHE Technical Frameworks General Introduction, Section 9 - Copyright Licenses for copyright license information for frequently referenced base standards. Information pertaining to the use of IHE International copyrighted materials is also available there.

10 Trademark

IHE® and the IHE logo are trademarks of the Healthcare Information Management Systems Society in the United States and trademarks of IHE Europe in the European Community. Please refer to the IHE Technical Frameworks General Introduction, Section 10 - Trademark for information on their use.
IHE Technical Frameworks General Introduction Appendices

The IHE Technical Framework General Introduction Appendices are components shared by all of the IHE domain technical frameworks. Each technical framework volume contains links to these documents where appropriate.

Update the following appendices to the General Introduction as indicated below. Note that these are not appendices to this domain’s Technical Framework (TF-1, TF-2, TF-3 or TF-4) but rather, they are appendices to the IHE Technical Frameworks General Introduction located here.

Appendix A – Actors

Add the following new or modified actors to the IHE Technical Frameworks General Introduction Appendix A:

<table>
<thead>
<tr>
<th>New (or modified) Actor Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brachytherapy Treatment Record Producer</td>
<td>A system capable of producing a Brachytherapy Treatment Record.</td>
</tr>
<tr>
<td>Brachytherapy Treatment Record Consumer</td>
<td>A system capable of consuming a Brachytherapy Treatment Record.</td>
</tr>
</tbody>
</table>

Appendix B – Transactions

Add the following new or modified transactions to the IHE Technical Frameworks General Introduction Appendix B:

<table>
<thead>
<tr>
<th>New (or modified) Transaction Name and Number</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brachytherapy Treatment Record Storage [TDRC-BRACHY-01]</td>
<td>Stores a Brachytherapy Treatment Record</td>
</tr>
</tbody>
</table>
Appendix D – Glossary

Add the following new or modified glossary terms to the *IHE Technical Frameworks General Introduction Appendix D*:

No new glossary terms.
Volume 1 – Profiles

X Brachy Treatment Delivery– Record Content (TDRC-Brachy) Profile

The Brachytherapy Treatment Delivery Record Content (TDRC-Brachy) Profile specifies the content of the RT Brachy Treatment Record being transferred from the Brachytherapy Treatment Record Producer to the Brachytherapy Treatment Record Consumer.

This profile will ensure that the treatment record transferred from the producer will contain all necessary information. It will especially address the following:

1. This profile will make the therapists’ workflow easier and faster, because it incorporates all information needed for treatment delivery record keeping.
2. This will enable Therapists to concentrate more on finalizing the details of patient treatment rather dealing with technical issues.
3. Fewer uncertainties will also reduce the time physicists and therapists need to gather and review treatment delivery information.

TDRC-Brachy contributes to patient safety, because it will standardize the content which is reviewed to assess if delivery of treatment was accurate to the plan. The profile will not specify how to validate the treatment delivery parameters, but will ensure that they are available, and well-formatted. The applications involved are Treatment Management Systems, Treatment Delivery Systems and Quality Assurance Devices.

This profile is a content-focused profile. For further information on the context, see Section X.6 TDRC-Brachy Cross Profile Considerations
X.1 TDRC-Brachy Actors, Transactions, and Content Modules

Figure X.1-1 shows how this content profile is used in the exchanging of DICOM treatment records between actors that are identified as producers and actors that are identified as consumers.

The DICOM objects that are exchanged between producers and consumers have to implement the requirements listed in this profile in order to be IHE compliant.

Table X.1-1 lists the content module(s) defined in the TDRC-Brachy Profile. To claim support with this profile, an actor shall support all required content modules (labeled “R”) and may support optional content modules (labeled “O”).

<table>
<thead>
<tr>
<th>Actors</th>
<th>Content Modules</th>
<th>Optionality</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brachytherapy Treatment Record Producer</td>
<td>RT Brachy Treatment Record IOD</td>
<td>R</td>
<td>TDRC-Brachy-01</td>
</tr>
<tr>
<td>Brachytherapy Treatment Record Consumer</td>
<td>RT Brachy Treatment Record IOD</td>
<td>R</td>
<td>TDRC-Brachy-01</td>
</tr>
</tbody>
</table>

X.1.1 Actor Descriptions and Actor Profile Requirements

For Brachytherapy Treatment Record Consumers, the display requirements for dwell time and total dose contributions are not sufficiently met by just presenting the DICOM data. It must be converted as described in the notes in this section. An actor does not adhere to the profile unless the system provides the output in the prescribed format.

"Actors shall display total times and dwell times at the reference date and time of the delivery and not simple Control Point Values”.

X.2 TDRC-Brachy Actor Options

Options that may be selected for each actor in this profile, if any, are listed in the Table X.2-1. Dependencies between options, when applicable, are specified in notes.
Table X.2-1: TDRC-Brachy – Actors and Options

<table>
<thead>
<tr>
<th>Actor</th>
<th>Option Name</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>No options</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

235 **X.3 TDRC-Brachy Required Actor Groupings**

Not applicable.

**X.4 TDRC-Brachy Overview**

**X.4.1 Concepts**

The Brachytherapy Treatment Record Content Consumer receives the RT Brachy Treatment Record from a Brachytherapy Treatment Record Content Producer, which holds the treatment record or prepares it for transmission.

**X.4.2 Use Cases**

Not Applicable.

**X.5 TDRC-Brachy Security Considerations**

Not Applicable.

**X.6 TDRC-Brachy Cross Profile Considerations**

Not Applicable
Volume 2 – Transactions

Add section 3.Y

3.Y1 Brachytherapy Treatment Record Storage [TDRC-Brachy-01]

3.Y1.1 Scope

In the Brachytherapy Treatment Record Storage transaction, a Producer of an RT Treatment Record that incorporates the brachytherapy technique identified in Brachytherapy Treatment Record Storage [TDRC-Brachy-0], stores the plan to a Brachytherapy Treatment Record Consumer.

3.Y1.2 Actor Roles

<table>
<thead>
<tr>
<th>Actor:</th>
<th>Role:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brachytherapy Treatment</td>
<td>Creates a brachytherapy treatment record and stores/shares? it to/with? a</td>
</tr>
<tr>
<td>Record Producer</td>
<td>Brachytherapy Treatment Record Consumer</td>
</tr>
<tr>
<td>Brachytherapy Treatment</td>
<td>Accepts and stores the Brachytherapy Treatment Record from a Brachytherapy</td>
</tr>
<tr>
<td>Record Consumer</td>
<td>Treatment Record Producer</td>
</tr>
</tbody>
</table>

3.Y1.3 Referenced Standards

DICOM 2023b Edition. PS 3.3: RT Modules, PS 3.4: Storage Service Class.
3.Y1.4 Messages

![Interaction Diagram]

Figure 3.Y1.4-1: Interaction Diagram
Appendices to Volume 2

270  None
Volume 3 – Content Modules

5 IHE Namespaces, Concept Domains and Vocabularies

5.1 IHE Radiation Oncology Namespaces
None

5.2 IHE Radiation Oncology Concept Domains
None

5.3 IHE Radiation Oncology Format Codes and Vocabularies

5.3.1 IHE Format Codes
None

5.3.2 IHEActCode Vocabulary
None

5.3.3 IHERoleCode Vocabulary
None

6 Radiation Oncology HL7 V3 CDA Content Modules

285 No HL7 V3 CDA Content Modules defined.
7 Radiation Oncology DICOM Content Definition

DICOM Content Definitions constrain the use of instances of specific DICOM IODs (also referred to as DICOM objects). This typically means placing requirements on the creators of those instances, although requirements may also be placed on the receivers and users.

The most common such requirements are to:

- Make a module that is optional (U) in a DICOM IOD be required or conditional,
- Make an attribute that is optional (Type 3) in a DICOM Module be required or conditional,
- Require that an attribute that is optional (Type 3) in a DICOM Module be absent
- Constrain the content of an attribute to be empty
- Constrain the content of an attribute to be populated in a certain way, such as:
  - Constraining the value to be taken from a specific table
  - Constraining the value to be copied from a specific source
  - Constraining the value to encode certain information
- Require that an attribute be displayed/accessible to the operator

Reiterating DICOM requirements is kept to a minimum sufficient to provide context for the IHE requirements. Implementers are still required to be familiar with, and conform to, the underlying DICOM specification.

Content Definitions may be referenced from a Profile independent of transactions to constrain content without specifying the transport. Content Definitions may also be referenced from within a Transaction specification to constrain the content without duplicating the same constraint text across multiple related transactions.

For attributes that are optional, the creator is permitted but not required to include them, and the receiver is permitted but not required to ignore them.

7.1 Conventions

DICOM Conventions are defined in Appendix E to the IHE Technical Frameworks General Introduction.

Key to IHE-RO Column of requirements

- R+ = The requirement is an IHE extension of the DICOM requirements and needs to be displayed (note: when consumed!, not produced)
- R* = The attribute is required to be there but not required to be displayed
- R+* = The Requirement is an IHE extension of the DICOM requirements, but it is NOT required to be displayed
7.2 General Definitions

7.3 IOD Definitions

7.3.6 Treatment Record IODs

7.3.6.1 Technique Specific RT Treatment Record

7.3.6.2 RT Treatment Record for General Use

7.3.6.3 RT Brachy Treatment Record

7.3.6.3.1 RT Brachy Treatment Record IOD

7.3.6.3.1.1 Referenced Standards

DICOM 2023b Edition. PS 3.3
### 7.3.6.3.1.2 IOD Definition

<table>
<thead>
<tr>
<th>IE</th>
<th>Module</th>
<th>Reference</th>
<th>Usage</th>
<th>IHE-RO Usage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient</td>
<td>Patient</td>
<td>C.7.1.1</td>
<td>M</td>
<td>M</td>
</tr>
<tr>
<td></td>
<td>Clinical Trial Subject</td>
<td>C.7.1.3</td>
<td>U</td>
<td></td>
</tr>
<tr>
<td>C.Study</td>
<td>General Study</td>
<td>C.7.2.1</td>
<td>M</td>
<td>M</td>
</tr>
<tr>
<td></td>
<td>Patient Study</td>
<td>C.7.2.2</td>
<td>U</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Clinical Trial Study</td>
<td>C.7.2.3</td>
<td>U</td>
<td></td>
</tr>
<tr>
<td>8Series</td>
<td>RT Series</td>
<td>C.8.8.1</td>
<td>M</td>
<td>M</td>
</tr>
<tr>
<td></td>
<td>Clinical Trial Series</td>
<td>C.7.3.2</td>
<td>U</td>
<td></td>
</tr>
<tr>
<td>Equipment</td>
<td>General Equipment</td>
<td>C.7.5.1</td>
<td>M</td>
<td>M</td>
</tr>
<tr>
<td>Treatment</td>
<td>RT General Treatment Record</td>
<td>C.8.8.17</td>
<td>M</td>
<td>M</td>
</tr>
<tr>
<td>Record</td>
<td>RT Patient Setup</td>
<td>C.8.8.12</td>
<td>U</td>
<td></td>
</tr>
<tr>
<td></td>
<td>RT Treatment Machine Record</td>
<td>C.8.8.18</td>
<td>M</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Measured Dose Reference Record</td>
<td>C.8.8.19</td>
<td>U</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Calculated Dose Reference Record</td>
<td>C.8.8.20</td>
<td>U</td>
<td></td>
</tr>
<tr>
<td></td>
<td>RT Brachy Session Record</td>
<td>C.8.8.22</td>
<td>M</td>
<td>M</td>
</tr>
<tr>
<td></td>
<td>RT Treatment Summary Record</td>
<td>C.8.8.23</td>
<td>U</td>
<td></td>
</tr>
<tr>
<td></td>
<td>SOP Common</td>
<td>C.12.1</td>
<td>M</td>
<td>M</td>
</tr>
</tbody>
</table>

#### 7.4 Module Definitions

**7.4.1 General Modules**

*This section is present only to convey the envisioned section numbering.*

**7.4.2 Workflow-Related Modules**

*This section is present only to convey the envisioned section numbering.*
7.4.11 Treatment Records

7.4.11.6 RT Brachy Treatment Record

7.4.11.6.1 Referenced Standards
DICOM 2023b Edition PS 3.3

7.4.11.6.2 RT Brachy Session Record Module Definition for HDR and PDR

Key to IHE-RO Column of requirements
- R+ = The requirement is an IHE extension of the DICOM requirements and needs to be displayed (note: when consumed!, not produced)
- R* = The attribute is required to be there but not required to be displayed
- R+* = The Requirement is an IHE extension of the DICOM requirements, but it is NOT required to be displayed
- O+ = The attribute is optional but if there, it must be displayed.
- -* = The DICOM usage applies but the value does not need to be displayed

<table>
<thead>
<tr>
<th>Attribute Name</th>
<th>Tag</th>
<th>Type</th>
<th>IHE-RO</th>
<th>Attribute Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Referenced Fraction Group Number</td>
<td>(300C,0022)</td>
<td>3</td>
<td>R+</td>
<td>Identifier of Fraction Group within referenced RT Plan.</td>
</tr>
<tr>
<td>Number of Fractions Planned</td>
<td>(300A,0078)</td>
<td>2</td>
<td>R+</td>
<td>Total number of treatments (Fractions) planned for current Fraction Group.</td>
</tr>
<tr>
<td>Brachy Treatment Technique</td>
<td>(300A,0200)</td>
<td>1</td>
<td>-*</td>
<td>No need to display. Type of brachytherapy treatment technique.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Enumerated Values:</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>INTRALUMENARY</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>INTRACAVITARY</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>INTERSTITIAL</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>CONTACT</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>INTRAVASCULAR</td>
</tr>
<tr>
<td>Brachy Treatment Type</td>
<td>(300A,0202)</td>
<td>1</td>
<td>R+</td>
<td>Type of brachytherapy treatment. HDR or PDR only</td>
</tr>
<tr>
<td>Recorded Source Sequence</td>
<td>(3008,0100)</td>
<td>1</td>
<td>-</td>
<td>Sequence of Sources to be used within Application Setups. One or more Items shall be included in this Sequence.</td>
</tr>
<tr>
<td>Attribute Name</td>
<td>Tag</td>
<td>Type</td>
<td>IHE-RO</td>
<td>Attribute Description</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>-------------------</td>
<td>------</td>
<td>--------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>&gt;Source Number</td>
<td>(300A,0212)</td>
<td>1</td>
<td>-*</td>
<td>Identification number of the Source. The value of Source Number (300A,0212) shall be unique within the Recorded Source Sequence (3008,0100) in which it is created.</td>
</tr>
<tr>
<td>&gt;Source Type</td>
<td>(300A,0214)</td>
<td>1</td>
<td>-*</td>
<td>Type of Source.</td>
</tr>
<tr>
<td>&gt;Source Manufacturer</td>
<td>(300A,0216)</td>
<td>2</td>
<td>-*</td>
<td>Manufacturer of source.</td>
</tr>
<tr>
<td>&gt;Source Serial Number</td>
<td>(3008,0105)</td>
<td>2</td>
<td>R⁺</td>
<td>User-supplied identifier for the radioactive source model that was used for the source in the treatment plan of which this session record is based to. See Section C.8.8.15.15.</td>
</tr>
<tr>
<td>&gt;Source Model ID</td>
<td>(300A,021B)</td>
<td>3</td>
<td>-*</td>
<td>User-supplied identifier for the radioactive source model that was used for the source in the treatment plan of which this session record is based to. See Section C.8.8.15.15.</td>
</tr>
<tr>
<td>&gt;Source Description</td>
<td>(300A,021C)</td>
<td>3</td>
<td>-*</td>
<td></td>
</tr>
<tr>
<td>&gt;Source Isotope Name</td>
<td>(300A,0226)</td>
<td>1</td>
<td>-*</td>
<td></td>
</tr>
<tr>
<td>&gt;Source Isotope Half Life</td>
<td>(300A,0228)</td>
<td>1</td>
<td>-*</td>
<td>Half-life of Isotope (days).</td>
</tr>
<tr>
<td>&gt;Source Strength Units</td>
<td>(300A,0229)</td>
<td>1C</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>&gt;Reference Air Kerma Rate</td>
<td>(300A,022A)</td>
<td>1</td>
<td>-</td>
<td>Air Kerma Rate in air of Isotope specified at Source Strength Reference Date (300A,022C) and Source Strength Reference Time (300A,022E) (in µGy h⁻¹ at 1 m). Value shall be zero for non-gamma sources.</td>
</tr>
<tr>
<td>&gt;Source Strength</td>
<td>(300A,022B)</td>
<td>1C</td>
<td>-</td>
<td>Source Strength of Isotope at Source Strength Reference Date (300A,022C) and Source Strength Reference Time (300A,022E), in units specified in Source Strength Units (300A,0229). Required if the source is not a gamma-emitting (photon) source. See Section C.8.8.15.13.</td>
</tr>
<tr>
<td>&gt;Source Strength Reference Date</td>
<td>(300A,022C)</td>
<td>1</td>
<td>-</td>
<td>Reference date for Reference Air Kerma Rate (300A,022A) or Source Strength (300A,022B) of Isotope.</td>
</tr>
<tr>
<td>Attribute Name</td>
<td>Tag</td>
<td>Type</td>
<td>IHE-RO</td>
<td>Attribute Description</td>
</tr>
<tr>
<td>---------------------------------------------</td>
<td>----------------------</td>
<td>------</td>
<td>--------</td>
<td>--------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>&gt;Source Strength Reference Time</td>
<td>(300A,022E)</td>
<td>1</td>
<td>-</td>
<td>Reference time for Air Kerma Rate (300A,022A) or Source Strength (300A,022B) of Isotope.</td>
</tr>
<tr>
<td>Treatment Session Application Setup Sequence</td>
<td>(3008,0110)</td>
<td>1</td>
<td>-*</td>
<td>Sequence of Application Setups for RT Treatment Record for current RT Plan. One or more Items shall be included in this Sequence.</td>
</tr>
<tr>
<td>&gt;Current Fraction Number</td>
<td>(3008,0022)</td>
<td>2</td>
<td>R</td>
<td>Fraction number for this application setup.</td>
</tr>
<tr>
<td>&gt;Application Setup Type</td>
<td>(300A,0232)</td>
<td>1</td>
<td>-*</td>
<td>Type of Application Setup. Defined Terms: FLETCHER_SUIT DELCLOS BLOEDORN JOSLIN_FLYNN CHANDIGARH MANCHESTER HENSCHKE NASOPHARYNGEAL OESOPHAGEAL ENDOBRONCHIAL SYED_NEBLETT ENDORECTAL PERINEAL</td>
</tr>
<tr>
<td>&gt;Referenced Brachy Application Setup Number</td>
<td>(300C,000C)</td>
<td>3</td>
<td>-*</td>
<td>References application setup specified by Application Setup Number (300A,0234) in Application Setup Sequence (300A,0230) in RT Brachy Application Setups Module within referenced RT Plan.</td>
</tr>
<tr>
<td>&gt;Application Setup Name</td>
<td>(300A,0236)</td>
<td>3</td>
<td>-*</td>
<td>User-defined name for Application Setup.</td>
</tr>
<tr>
<td>&gt;Application Setup Manufacturer</td>
<td>(300A,0238)</td>
<td>3</td>
<td>-*</td>
<td>Manufacturer of Application Setup.</td>
</tr>
<tr>
<td>&gt;Template Number</td>
<td>(300A,0240)</td>
<td>3</td>
<td>-*</td>
<td>Identification number of the Template.</td>
</tr>
<tr>
<td>&gt;Template Type</td>
<td>(300A,0242)</td>
<td>3</td>
<td>-*</td>
<td>User-defined type for Template Device.</td>
</tr>
<tr>
<td>&gt;Template Name</td>
<td>(300A,0244)</td>
<td>3</td>
<td>-*</td>
<td>User-defined name for Template Device.</td>
</tr>
<tr>
<td>&gt;Application Setup Check</td>
<td>(3008,0116)</td>
<td>3</td>
<td>-*</td>
<td>Results of check-wire travel through all channels of current Application Setup.</td>
</tr>
<tr>
<td>Enumerated Values:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PASSED</td>
<td>Passed check</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FAILED</td>
<td>Failed check</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>UNKNOWN</td>
<td>Unknown status</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Attribute Name</td>
<td>Tag</td>
<td>Type</td>
<td>IHE-RO</td>
<td>Attribute Description</td>
</tr>
<tr>
<td>----------------------------------------------------</td>
<td>-------------------</td>
<td>------</td>
<td>--------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>&gt;Referenced Verification Image Sequence</td>
<td>(300C,0040)</td>
<td>3</td>
<td>-*</td>
<td>Sequence of verification images obtained during delivery of current beam. One or more Items are permitted in this Sequence. See Note 1.</td>
</tr>
<tr>
<td>&gt;&gt;Include Table 10-11 “SOP Instance Reference Macro Attributes”</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt;Total Reference Air Kerma</td>
<td>(300A,0250)</td>
<td>1</td>
<td>-</td>
<td>Total Reference Air Kerma for current Application Setup, i.e., the sum of the products of the Air Kerma Rates of each Source in each Channel with its respective Channel Time (µGy at 1 m). Value shall be zero for non-gamma sources.</td>
</tr>
<tr>
<td>&gt;Referenced Calculated Dose Reference Sequence</td>
<td>(3008,0090)</td>
<td>3</td>
<td>R*</td>
<td>Sequence of doses estimated for each treatment delivery. One or more Items are permitted in this Sequence.</td>
</tr>
<tr>
<td>&gt;&gt;Referenced Dose Reference Number</td>
<td>(300C,0051)</td>
<td>1</td>
<td>-*</td>
<td>Uniquely identifies Dose Reference specified by Dose Reference Number (300A,0012) in Dose Reference Sequence (300A,0010) in RT Prescription Module of referenced RT Plan. Required if Referenced Calculated Dose Reference Number (3008,0092) is not present. It shall not be present otherwise.</td>
</tr>
<tr>
<td>&gt;&gt;Referenced Calculated Dose Reference Number</td>
<td>(3008,0092)</td>
<td>1C</td>
<td>-*</td>
<td>Uniquely identifies Calculated Dose Reference specified by Calculated Dose Reference Number (3008,0072) within Calculated Dose Reference Sequence (3008,0070). Required if Referenced Dose Reference Number (300C,0051) is not present. It shall not be present otherwise.</td>
</tr>
<tr>
<td>&gt;&gt;Calculated Dose Reference Dose Value</td>
<td>(3008,0076)</td>
<td>1</td>
<td>-</td>
<td>Calculated Dose (Gy).</td>
</tr>
<tr>
<td>&gt;Current Fraction Number</td>
<td>(3008,0022)</td>
<td>2</td>
<td>R</td>
<td>Delivery Type of treatment. Defined Terms: TREATMENT normal patient treatment CONTINUATION continuation of interrupted treatment</td>
</tr>
<tr>
<td>&gt;Treatment Delivery Type</td>
<td>(300A,00CE)</td>
<td>2</td>
<td>R</td>
<td>Delivery Type of treatment. Defined Terms: TREATMENT normal patient treatment CONTINUATION continuation of interrupted treatment</td>
</tr>
<tr>
<td>Attribute Name</td>
<td>Tag</td>
<td>Type</td>
<td>IHE-RO</td>
<td>Attribute Description</td>
</tr>
<tr>
<td>----------------</td>
<td>-----</td>
<td>------</td>
<td>--------</td>
<td>-----------------------</td>
</tr>
<tr>
<td>&gt;Treatment Termination Status</td>
<td>(3008,002A)</td>
<td>1</td>
<td>R</td>
<td>Conditions under which treatment was terminated. Enumerated Values: NORMAL treatment terminated normally OPERATOR operator terminated treatment MACHINE machine terminated treatment for other than NORMAL condition UNKNOWN status at termination unknown</td>
</tr>
<tr>
<td>&gt;RT Treatment Termination Reason Code Sequence</td>
<td>(300A,0715)</td>
<td>3</td>
<td>R</td>
<td>Identifies the reason why the current treatment session has terminated. One or more items are permitted in this Sequence. The system must display the Code Meaning as a minimum.</td>
</tr>
<tr>
<td>&gt;Recorded Brachy Accessory Device Sequence</td>
<td>(3008,0120)</td>
<td>3</td>
<td>-*</td>
<td>Sequence of Brachy Accessory Devices associated with current Application Setup. One or more Items are permitted in this Sequence.</td>
</tr>
<tr>
<td>&gt;Referenced Brachy Accessory Device Number</td>
<td>(3008,0122)</td>
<td>2</td>
<td></td>
<td>Identification number of the Brachy Accessory Device. The value of Brachy Accessory Device Number (300A,0262) shall be unique within the Application Setup in which it is created.</td>
</tr>
<tr>
<td>&gt;&gt;Brachy Accessory Device ID</td>
<td>(300A,0263)</td>
<td>2</td>
<td></td>
<td>User or machine supplied identifier for Brachy Accessory Device.</td>
</tr>
<tr>
<td>&gt;&gt;Brachy Accessory Device Type</td>
<td>(300A,0264)</td>
<td>1</td>
<td></td>
<td>Type of Brachy Accessory Device. Defined Terms: SHIELD DILATATION MOLD PLAQUE FLAB</td>
</tr>
<tr>
<td>Attribute Name</td>
<td>Tag</td>
<td>Type</td>
<td>IHE-RO</td>
<td>Attribute Description</td>
</tr>
<tr>
<td>----------------------------------------------------</td>
<td>--------------------</td>
<td>------</td>
<td>--------</td>
<td>---------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>&gt;&gt;Brachy Accessory Device Name</td>
<td>(300A,0266)</td>
<td>3</td>
<td></td>
<td>User-defined name for Brachy Accessory Device.</td>
</tr>
<tr>
<td>&gt;Recorded Channel Sequence</td>
<td>(3008,0130)</td>
<td>1</td>
<td>-*</td>
<td>Sequence of Channels for current Application Setup. One or more Items shall be included in this Sequence.</td>
</tr>
<tr>
<td>&gt;&gt;Channel Number</td>
<td>(300A,0282)</td>
<td>1</td>
<td>-*</td>
<td>Identification number of the Channel. The value of Channel Number (300A,0282) shall be unique within the Application Setup in which it is created.</td>
</tr>
<tr>
<td>&gt;&gt;Referenced Channel Number</td>
<td>(0074,1406)</td>
<td>3</td>
<td>-*</td>
<td>The channel to be delivered, specified by the value of Channel Number (300A,0282) in referenced RT Plan.</td>
</tr>
<tr>
<td>&gt;&gt;Channel Length</td>
<td>(300A,0284)</td>
<td>2</td>
<td>-*</td>
<td>Length of Channel (mm). See RT Plan IOD.</td>
</tr>
<tr>
<td>&gt;&gt;Channel Effective Length</td>
<td>(300A,0271)</td>
<td>3</td>
<td>R+</td>
<td>Length of Channel (in mm) defined as the distance between the connector on the afterloader and the center of the distal-most possible position of the source. See Section C.8.8.15.16.</td>
</tr>
<tr>
<td>&gt;&gt;Channel Inner Length</td>
<td>(300A,0272)</td>
<td>2C</td>
<td>R+</td>
<td>The total physical inner length of channel (in mm). Specifies the distance between the connector on the afterloader and the end of the channel. Required if Channel Effective Length (300A,0271) is present. See Section C.8.8.15.16.1.</td>
</tr>
<tr>
<td>&gt;&gt;Afterloader Channel ID</td>
<td>(300A,0273)</td>
<td>3</td>
<td>R+</td>
<td>Identification of the Channel connection on the afterloader. See Section C.8.8.15.16.2.</td>
</tr>
<tr>
<td>&gt;&gt;Specified Channel Total Time</td>
<td>(3008,0132)</td>
<td>1</td>
<td>-</td>
<td>Total amount of time in seconds, scaled for the current source delivery strength and other delivery factors, specified to be delivered at the time of treatment between the first Control Point and the final Control Point for the current Channel. In the case of resuming a partially delivered treatment, the Specified Channel Total time will only include the remainder to be treated. See Section C.8.8.22.2.</td>
</tr>
<tr>
<td>&gt;&gt;Delivered Channel Total Time</td>
<td>(3008,0134)</td>
<td>1</td>
<td>-</td>
<td>Total amount of time in seconds actually delivered between Control Point 0 and final Control Point of the Brachy Control Point Delivered Sequence (3008,0160) for current Channel.</td>
</tr>
<tr>
<td>Attribute Name</td>
<td>Tag</td>
<td>Type</td>
<td>IHE-RO</td>
<td>Attribute Description</td>
</tr>
<tr>
<td>----------------------------------------------</td>
<td>--------------------------</td>
<td>------</td>
<td>--------</td>
<td>----------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>&gt;&gt;Source Movement Type</td>
<td>(300A,0288)</td>
<td>1</td>
<td>-*</td>
<td>Type of Source movement. Defined Terms: STEPWISE FIXED OSCILLATING UNIDIRECTIONAL</td>
</tr>
<tr>
<td>&gt;&gt;Specified Number of Pulses</td>
<td>(3008,0136)</td>
<td>1C</td>
<td>-</td>
<td>Number of Pulses specified per fraction for current Channel. Required if Brachy Treatment Type (300A,0202) is PDR. See Section C.8.8.22.1.</td>
</tr>
<tr>
<td>&gt;&gt;Delivered Number of Pulses</td>
<td>(3008,0138)</td>
<td>1C</td>
<td>-</td>
<td>Number of Pulses actually delivered per fraction for current Channel. Required if Brachy Treatment Type (300A,0202) is PDR. See Section C.8.8.22.1.</td>
</tr>
<tr>
<td>&gt;&gt;Specified Pulse Repetition Interval</td>
<td>(3008,013A)</td>
<td>1C</td>
<td>-</td>
<td>Pulse repetition interval (sec) specified for current Channel. Required if Brachy Treatment Type (300A,0202) is PDR. See Section C.8.8.22.1.</td>
</tr>
<tr>
<td>&gt;&gt;Delivered Pulse Repetition Interval</td>
<td>(3008,013C)</td>
<td>1C</td>
<td>-</td>
<td>Pulse repetition interval (sec) actually delivered for current Channel. Required if Brachy Treatment Type (300A,0202) is PDR. See Section C.8.8.22.1.</td>
</tr>
<tr>
<td>&gt;&gt;Referenced Measured Dose Reference Sequence</td>
<td>(3008,0080)</td>
<td>3</td>
<td>-*</td>
<td>Sequence of doses measured during treatment delivery, summed over entire session. One or more Items are permitted in this Sequence.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Referenced Dose Reference Number</td>
<td>(300C,0051)</td>
<td>1C</td>
<td></td>
<td>Uniquely references Dose Reference specified by Dose Reference Number (300A,0012) in Dose Reference Sequence (300A,0010) in RT Prescription Module of referenced RT Plan. Required if Referenced Measured Dose Reference Number (3008,0082) is not present. It shall not be present otherwise.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Referenced Measured Dose Reference Number</td>
<td>(3008,0082)</td>
<td>1C</td>
<td></td>
<td>References Measured Dose Reference specified by Measured Dose Reference Number (3008,0064) in Measured Dose Reference Sequence (3008,0010). Required if Referenced Dose Reference Number (300C,0051) is not present. It shall not be present otherwise.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Measured Dose Value</td>
<td>(3008,0016)</td>
<td>1</td>
<td>-</td>
<td>Measured Dose.</td>
</tr>
<tr>
<td>&gt;&gt;Referenced Calculated Dose Reference Sequence</td>
<td>(3008,0090)</td>
<td>3</td>
<td>-*</td>
<td></td>
</tr>
<tr>
<td>Attribute Name</td>
<td>Tag</td>
<td>Type</td>
<td>IHE-RO</td>
<td>Attribute Description</td>
</tr>
<tr>
<td>-----------------------------------</td>
<td>--------------------</td>
<td>------</td>
<td>--------</td>
<td>--------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Referenced Dose Reference</td>
<td>(300C,0051)</td>
<td>1C</td>
<td>-*</td>
<td>Uniquely identifies Dose Reference specified by Dose Reference Number (300A,0012) in Dose Reference Sequence (300A,0010) in RT Prescription Module of referenced RT Plan. Required if Referenced Calculated Dose Reference Number (3008,0092) is not present. It shall not be present otherwise.</td>
</tr>
<tr>
<td>Number</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt;&gt;&gt;Referenced Calculated Dose</td>
<td>(3008,0092)</td>
<td>1C</td>
<td>-*</td>
<td>Uniquely identifies Calculated Dose Reference specified by Calculated Dose Reference Number (3008,0072) within Calculated Dose Reference Sequence (3008,0070). Required if Referenced Dose Reference Number (300C,0051) is not present. It shall not be present otherwise.</td>
</tr>
<tr>
<td>Reference Number</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt;&gt;&gt;Calculated Dose Reference</td>
<td>(3008,0076)</td>
<td>1</td>
<td>-*</td>
<td>Calculated Dose (Gy).</td>
</tr>
<tr>
<td>Dose Value</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt;&gt;Recorded Source Applicator</td>
<td>(3008,0140)</td>
<td>3</td>
<td>R:+</td>
<td>Only one entry in the sequence</td>
</tr>
<tr>
<td>Sequence</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt;&gt;&gt;Referenced Source Applicator</td>
<td>(3008,0142)</td>
<td>2</td>
<td>R:+</td>
<td>Identification number of the Source Applicator. The value of Source Applicator Number (300A,0290) shall be unique within the Channel in which it is created.</td>
</tr>
<tr>
<td>Number</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt;&gt;&gt;Source Applicator ID</td>
<td>(300A,0291)</td>
<td>2</td>
<td>R+</td>
<td>User or machine supplied identifier for Source Applicator.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Source Applicator Type</td>
<td>(300A,0292)</td>
<td>1</td>
<td>-*</td>
<td>Type of Source Applicator. Defined Terms: FLEXIBLE RIGID</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Source Applicator Name</td>
<td>(300A,0294)</td>
<td>3</td>
<td>-*</td>
<td>User-defined name for Source Applicator.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Source Applicator Length</td>
<td>(300A,0296)</td>
<td>1</td>
<td>-*</td>
<td>Length of Source Applicator (mm), defined as the distance between the connector of the applicator and the distal-most position of the source.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Source Applicator Tip Length</td>
<td>(300A,0274)</td>
<td>2C</td>
<td>R*</td>
<td>Length of Source Applicator Tip (in mm), defined as the distance between the outer tip of the applicator and the center of the distal-most possible position of the source. Required if Channel Effective Length (300A,0271) is present. See Section C.8.8.15.16.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Source Applicator Manufacturer</td>
<td>(300A,0298)</td>
<td>3</td>
<td>-*</td>
<td>Manufacturer of Source Applicator.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Source Applicator Step Size</td>
<td>(300A,02A0)</td>
<td>1C</td>
<td>-*</td>
<td>Distance of path along channel (mm) between adjacent (potential) dwell positions. Required if Source Movement Type (300A,0288) is STEPWISE.</td>
</tr>
<tr>
<td>Attribute Name</td>
<td>Tag</td>
<td>Type</td>
<td>IHE-RO</td>
<td>Attribute Description</td>
</tr>
<tr>
<td>----------------</td>
<td>---------------------</td>
<td>------</td>
<td>--------</td>
<td>-----------------------</td>
</tr>
<tr>
<td>&gt;&gt;Transfer Tube Number</td>
<td>(300A,02A2)</td>
<td>2</td>
<td>-*</td>
<td>Identification number of the Transfer Tube. The value of Transfer Tube Number (300A,02A2) shall be unique within the Channel in which it is created.</td>
</tr>
<tr>
<td>&gt;&gt;Transfer Tube Length</td>
<td>(300A,02A4)</td>
<td>2C</td>
<td>-*</td>
<td>Length of Transfer Tube of current afterloading Channel (mm). Required if value Transfer Tube Number (300A,02A2) is not zero length.</td>
</tr>
<tr>
<td>&gt;&gt;Recorded Channel Shield Sequence</td>
<td>(3008,0150)</td>
<td>3</td>
<td>-*</td>
<td>Sequence of Channel Shields associated with current Channel. One or more Items are permitted in this Sequence. See RT Plan IOD for description of Channel Shields.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Referenced Channel Shield Number</td>
<td>(3008,0152)</td>
<td>2</td>
<td>-*</td>
<td>Identification number of the Channel Shield. The value of Channel Shield Number (300A,02B2) shall be unique within the Channel in which it is created.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Channel Shield ID</td>
<td>(300A,02B3)</td>
<td>2</td>
<td>-*</td>
<td>User or machine supplied identifier for Channel Shield.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Channel Shield Name</td>
<td>(300A,02B4)</td>
<td>3</td>
<td>-*</td>
<td>User-defined name for Channel Shield.</td>
</tr>
<tr>
<td>&gt;&gt;Referenced Source Number</td>
<td>(300C,000E)</td>
<td>1</td>
<td>-*</td>
<td>Uniquely identifies the referenced Source within the Recorded Source Sequence (3008,0100) for current Application Setup.</td>
</tr>
<tr>
<td>&gt;&gt;Safe Position Exit Date</td>
<td>(3008,0162)</td>
<td>1C</td>
<td>-*</td>
<td>Date on which the source(s) exited the safe. Required if Recorded Channel Sequence (3008,0130) is present and Brachy Treatment Type (300A,0202) is not MANUAL or PDR.</td>
</tr>
<tr>
<td>&gt;&gt;Safe Position Exit Time</td>
<td>(3008,0164)</td>
<td>1C</td>
<td>-*</td>
<td>Time on which the source(s) exited the safe. Required if Recorded Channel Sequence (3008,0130) is present and Brachy Treatment Type (300A,0202) is not MANUAL or PDR.</td>
</tr>
<tr>
<td>&gt;&gt;Safe Position Return Date</td>
<td>(3008,0166)</td>
<td>1C</td>
<td>-*</td>
<td>Date on which the source(s) returned to the safe. Required if Recorded Channel Sequence (3008,0130) is present and Brachy Treatment Type (300A,0202) is not MANUAL or PDR.</td>
</tr>
<tr>
<td>&gt;&gt;Safe Position Return Time</td>
<td>(3008,0168)</td>
<td>1C</td>
<td>-*</td>
<td>Time on which the source(s) returned to the safe. Required if Recorded Channel Sequence (3008,0130) is present and Brachy Treatment Type (300A,0202) is not MANUAL or PDR.</td>
</tr>
<tr>
<td>&gt;&gt;Number of Control Points</td>
<td>(300A,0110)</td>
<td>1</td>
<td>-*</td>
<td>Number of control points in Channel. For an N-segment Channel there will be 2N (stepwise movement) or N+1 (continuous movement) control points.</td>
</tr>
<tr>
<td>Attribute Name</td>
<td>Tag</td>
<td>Type</td>
<td>IHE-RO</td>
<td>Attribute Description</td>
</tr>
<tr>
<td>----------------------------------------------------</td>
<td>-------------------</td>
<td>------</td>
<td>--------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>&gt;&gt;Brachy Control Point Delivered Sequence</td>
<td>(3008,0160)</td>
<td>1</td>
<td>-</td>
<td>Sequence of machine configurations describing this Channel. The number of Items in this Sequence shall equal the value of Number of Control Points (300A,0110). See RT Plan IOD and Section C.8.8.22.1 for description of Brachy Control Point Delivered Sequence.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Referenced Control Point Index</td>
<td>(300C,00F0)</td>
<td>3</td>
<td>-</td>
<td>Index of current Control Point, starting at 0 for first Control Point.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Treatment Control Point Date</td>
<td>(3008,0024)</td>
<td>1</td>
<td>-</td>
<td>Date when the delivery of radiation at this control point began. For the final control point this shall be the Date when the previous control point ended.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Treatment Control Point Time</td>
<td>(3008,0025)</td>
<td>1</td>
<td>-</td>
<td>Time when the delivery of radiation at this control point began. For the final control point this shall be the Time when the previous control point ended.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Control Point Relative Position</td>
<td>(300A,02D2)</td>
<td>1</td>
<td>-</td>
<td>Distance in mm between current Control Point Position and the center of the distal-most possible Source position in current Channel. See RT Plan IOD. Display only a single dwell position for the two matching control point locations that define a dwell location.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;Override Sequence</td>
<td>(3008,0060)</td>
<td>3</td>
<td>-</td>
<td>Sequence of parameters that were overridden during the administration of the treatment immediately prior to the current control point. One or more Items are permitted in this Sequence.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Override Parameter Pointer</td>
<td>(3008,0062)</td>
<td>2</td>
<td>-</td>
<td>Contains the Data Element Tag of the Attribute that was overridden.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Operators' Name</td>
<td>(0008,1070)</td>
<td>2</td>
<td>-</td>
<td>Name of operator who authorized override.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Operator Identification Sequence</td>
<td>(0008,1072)</td>
<td>3</td>
<td>-</td>
<td>Identification of the operator who authorized override. Only a single Item is permitted in this Sequence.</td>
</tr>
</tbody>
</table>

>>>Include Table 10-1 “Person Identification Macro Attributes Description”
<table>
<thead>
<tr>
<th>Attribute Name</th>
<th>Tag</th>
<th>Type</th>
<th>IHE-RO</th>
<th>Attribute Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Override Reason</td>
<td>(3008,0066)</td>
<td>3</td>
<td></td>
<td>User-defined description of reason for override of parameter specified by Override Parameter Pointer (3008,0062).</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;Pulse Specific Brachy Control Point Delivered Sequence</td>
<td>(3008,0171)</td>
<td>3</td>
<td>RC*</td>
<td>Brachy Control Point Delivered Sequence for each PDR treatment pulse. Number of Items in the Sequence shall be equal to the Delivered Number of Pulses (3008,0138). Required if PDR</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Pulse Number</td>
<td>(3008,0172)</td>
<td>1</td>
<td></td>
<td>Identification Number of this delivered Pulse. The pulse numbers for a treatment start at 1 and increase monotonically by 1. A given SOP Instance might only contain some of the pulses of the given treatment.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Safe Position Exit Date</td>
<td>(3008,0162)</td>
<td>1</td>
<td></td>
<td>Date on which the source(s) exited the safe.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Safe Position Exit Time</td>
<td>(3008,0164)</td>
<td>1</td>
<td></td>
<td>Time at which the source(s) exited the safe.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Safe Position Return Date</td>
<td>(3008,0166)</td>
<td>1</td>
<td></td>
<td>Date on which the source(s) returned to the safe.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Safe Position Return Time</td>
<td>(3008,0168)</td>
<td>1</td>
<td></td>
<td>Time at which the source(s) returned to the safe.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;Brachy Pulse Control Point Delivered Sequence</td>
<td>(3008,0173)</td>
<td>1</td>
<td></td>
<td>List of control points for this pulse. See Section C.8.8.22.1.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Referenced Control Point Index</td>
<td>(300C,00F0)</td>
<td>3</td>
<td></td>
<td>Index of current Control Point, starting at 0 for first Control Point in this Sequence.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Treatment Control Point Date</td>
<td>(3008,0024)</td>
<td>1</td>
<td></td>
<td>Date when the delivery of radiation at this control point began. For the final control point, this shall be the Date when the previous control point ended.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Treatment Control Point Time</td>
<td>(3008,0025)</td>
<td>1</td>
<td></td>
<td>Time when the delivery of radiation at this control point began. As described in section X.1.1, display the calculated dwell time spent at the treatment position defined by two control points at the same (3008, 0025) Control Point Relative Position. For the final control point, this shall be the Time when the previous control point ended. Also display the total dwell time for each channel (catheter).</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;Control Point Relative Position</td>
<td>(300A,02D2)</td>
<td>1</td>
<td></td>
<td>Distance in mm between current Control Point Position and the center of the distal-most possible Source position in current Channel. Display only a single dwell position for the two matching control point locations that define a dwell location. See Section C.8.8.15.9.</td>
</tr>
<tr>
<td>Attribute Name</td>
<td>Tag</td>
<td>Type</td>
<td>IHE-RO</td>
<td>Attribute Description</td>
</tr>
<tr>
<td>----------------------------------------------------------</td>
<td>--------------------</td>
<td>------</td>
<td>--------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Override Sequence</td>
<td>(3008,0060)</td>
<td>3</td>
<td></td>
<td>Parameters which were overridden during the administration of the treatment immediately prior to the current control point. One or more Items are permitted in this Sequence.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Override Parameter Pointer</td>
<td>(3008,0062)</td>
<td>2</td>
<td></td>
<td>Data Element Tag of the Attribute that was overridden.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Operators' Name</td>
<td>(0008,1070)</td>
<td>2</td>
<td></td>
<td>Name of operator who authorized override.</td>
</tr>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Operator Identification Sequence</td>
<td>(0008,1072)</td>
<td>3</td>
<td></td>
<td>Identification of the operator who authorized override. Only a single Item is permitted in this Sequence.</td>
</tr>
</tbody>
</table>

>>>>>Include Table 10-1 “Person Identification Macro Attributes Description”

<table>
<thead>
<tr>
<th>Attribute Name</th>
<th>Tag</th>
<th>Type</th>
<th>IHE-RO</th>
<th>Attribute Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;&gt;&gt;&gt;&gt;Override Reason</td>
<td>(3008,0066)</td>
<td>3</td>
<td></td>
<td>User-defined description of reason for override.</td>
</tr>
</tbody>
</table>

**Notes**

1. The Referenced Verification Image Sequence (300C,0040) may contain either images taken specifically for verification of the brachy application setup or reference images used in place of verification images, as might be done in HDR treatment planning.

2. Treatment Termination Code (3008,002B) was previously included in this Module but has been retired. See <<immediately previous PS3.3 DICOM revision>>. The RT Treatment Termination Reason Code Sequence (300A,0715) and Machine-Specific Treatment Termination Code Sequence (300A,0716) should be used for machine readable codes and Treatment Termination Description (300A,0730) for human readable text respectively.
Appendices to Volume 3

None
Volume 4 – National Extensions

4 National Extensions
Not applicable.