Treatment Data and Technical Process Challenges for Practical Big Data Efforts in Radiation Oncology

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40	Abstract
41	The term <i>Big Data</i> has come to encompass a number of concepts and uses within medicine.
42	This paper lays out the relevance and application of large collections of data in the radiation
43	oncology community. We describe the potential importance and uses in clinical practice. The
44	important concepts are then described and how they have been or could be implemented are
45	discussed. Impediments to progress in the collection and use of sufficient quantities of data are
46	also described. Finally, recommendations for how the community can move forward to achieve
47	the potential of Big Data in radiation oncology are provided.
48	
49	Introduction

To the clinician, it often seems that we have too much and too little data at the same time. We 50 spend more time than we would like at computer terminals entering or reading data. Perhaps 51 52 it would be better stated that we would like the *data* we input to be transformed into

information that we can use. This is the aspect of *Big Data* that this manuscript addresses.
Computerized data handling has been an integral part of our field since the introduction of
computerized treatment planning and record and verify systems. The question is, now that
there are highly successful algorithms for using computerized data to make models for
predictive purposes, can the radiation oncology community harness our data for our patients'
benefit?

Pan et al. have provided a very clear picture of the difficulties that we face in collecting and using data in the clinic [1]. The questions we must answer are: (a) is it worth making an effort to improve the situation, (b) what are the details of the clinical data environment that need to be addressed, and (c) how do we accomplish our goals? An AAPM Science Council Focused Research Meeting (FOREM) meeting, jointly sponsored with vendors, was held in Ann Arbor in May of 2017, to address these questions. In this publication we provide an overview and summary of the answers that emerged.

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67 Motivation for embracing Big Data

a. Need to learn from and adapt to emerging therapies such as genetics, immunotherapy

It is now commonly understood that the explosion of data and knowledge that has resulted from genomics will have a great impact on all areas of cancer care, including radiation therapy. A patient's genetic profile may play an important role in how they will react to certain agents or in their ability to repair radiation damage [2]. The tumor's genetic profiles (since many tumors have a multitude of different mutations) is increasingly being used to determine the best therapy or combination of therapies [3].

- Immunotherapy is another area of increasing importance. The ability to use different aspects
 of the immune system to target tumor cells is an area of great current interest [4].
- 77 Radiation oncology is not alone in the interest and need for better data on patients' genetic
- profiles. NIH has been working with a number of groups to establish a workable solution in

- 79 order to avoid the current problems such as laboratory-dependent formats, text-based storage,
- and lack of centralized storage in current electronic health records (EHR) [5].
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82 b. Cancer as chronic disease and multiple care givers

As cancer therapy becomes more effective, more and more patients are living longer. As a 83 result, the extent and complexity of information which needs to be tracked to improve 84 understanding of outcomes is increasing. For example, for patients who are essentially cancer 85 free, monitoring risk for treatment-related complications when their long term home location 86 87 based follow up is not at the treatment center is a challenge. Parry et al. estimate that there 88 will be 18 million cancer survivors by 2020 [6]. In addition, there are the increasing numbers of 89 patients who survive longer than ever due to improvements in targeted therapies, better 90 imaging and better methods for localizing dose [7]. These advances can lead to improved local control and better control of oligometastases. The upshot is that as the number of patients 91 92 who suffer cancer-related health consequences increases over time, the more likely it is that 93 they will see a wider spectrum of specialists and in a larger number of clinical settings, interacting with a large variety of recording-keeping systems. 94

95 Even just considering the electronic health records, there are no general standards for the 96 selection and formatting of data to be recorded. Different vendors, different institutions, 97 different departments and even different physicians have different methods which are often not compatible. Finally, even within well-structured organizations, much of the data exist 98 99 within text documents. Lack of standards for which data elements to gather, inconsistent 100 processes for entry and variability among commercial systems for aggregation and reporting increase the likelihood that physicians and staff will miss information or have incorrect 101 102 information regarding a patient's health and/or treatments that could potentially affect 103 decisions.

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105 c. Comparative Effectiveness Research

In the last decade, comparative effectiveness research (CER) has come to be seen as an important and necessary adjunct to randomized clinical trials (RCT) [8]. In CER, two different therapies or tests that are already accepted are compared, whereas RCT's focus on comparing a new to a current therapy. The Patient Centered Outcomes Research Institute cites CER as its primary method of research. Given the relatively small numbers of cancer patients that are enrolled in RCT's (approximately 3%), the need to use the information that is available through CER is understandable.

Comparative effectiveness research can be tailored along a spectrum of methods ranging from essentially an RCT to a comparison of current clinical practice with an integrated practice beyond the current norm. A recent paper by Fiore et al. looked at four different trials that sought to use only data in the current EHR's [9]. Their conclusions included: "We find that EHRbased clinical trials are feasible but pose limitations on the questions that can be addressed, the processes that can be implemented, and the outcomes that can be assessed."

Clearly, for progress to be made using CER practical methods for the easy and accuratecollection of data and for the sharing of data must be available in clinics.

121 d. Quality Improvement and Error Detection

122 The past few years have seen an explosion in the use of data to reduce errors in radiation 123 therapy. ASTRO and AAPM have implemented the Radiation Oncology—Incident Learning System (RO-ILS) that relies on data submitted to it to develop a shared learning platform. 124 While this system is not "big data" in the sense that it is in text format and is a relatively small 125 126 amount of data, it does count in our definition of transforming data to information. In 127 particular, the system is set up to provide users with more knowledge about the sources of 128 errors and how best to avoid them. Another area is in artificial intelligence applications of error 129 detection. For example, Kalet et al. successfully mined an OIS to develop a probabilistic model 130 of the contributing factors to errors [10].

131 e. Modeling in Radiation Oncology

132 Perhaps the most widespread use of data in radiation oncology is in modeling. The examples are too numerous to list, but some of the most impactful models are the QUANTEC models, 133 134 outcomes, tumor control probabilities, equivalent uniform dose, and biologically effective dose 135 [11]. As construction of Big Data Analytics Resource Systems (BDARS) aggregating a wider range of health care information (e.g. labs, medications, genomics, demographics, patient 136 reported outcomes (PROs) etc.) expands, more comprehensive models are progressing beyond 137 138 dose metrics alone [12-14]. In addition, heuristic type models have been constructed for automating the objectives of inverse planning and library-based contouring. A promising area 139 for the more conventional use of big data is in machine learning for automated contouring. In 140 141 this application, images that have been segmented by experts are fed into a machine learning 142 algorithm and image features that predict the true contours are selected to produce anatomical contour models. 143

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146 State of the Data

147 One of the most important concepts is that Big Data, in most cases, implies more data than may 148 be obtained by any single institution. In order to use machine learning or any modeling 149 techniques, there must be enough data to (a) build the model, (b) test the model, and (c) 150 validate the model. Optimally, validation (c) can be done with data from a different institution 151 in order to account for hidden variables that may not be appreciated [15]. In addition, as our 152 ability to differentiate patients improves, e.g. genomics and radiomics, the number of patients 153 suitable for any given model decreases, thereby increasing our need for more comprehensive 154 capture of intra-institutional data as well as for multi-institutional data. This has critical implications for how organizations cooperate. Whereas success in medical research in the past 155 has favored very large single institutions that can develop a critical mass of knowledge and 156 157 resources in close physical proximity, diffuse networks of institutions able to generate and 158 share information will have an advantage in the future.

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In addition to the need for broad (many patients) data sources, we also need deep 160 (relationships among key data elements) sources. Systems promoted as big data sources may in 161 162 fact be shallow, capturing only a few data elements for a large number of patients. For 163 example, some data sources draw upon billing records or imaging records for a large number of patients, but lack depth needed to enable linkage to diagnosis, treatment, dosimetric or 164 outcomes details. Another impediment to obtaining the "deep" type of data is that sources 165 often dump unstructured, "as is", data into data lakes where key data elements and 166 relationships can in principle be extracted, but in practice carry a high overhead for extraction. 167 Challenges for ensuring depth in aggregation of key data elements needed for radiation 168 169 oncology fall into four categories

- Access Staff possessing both domain knowledge of radiation oncology and of
 informatics need access to query data bases in source systems to construct
 functional big data repositories.
- Data Integrity Data elements that may not require accurate entry to enable
 treatment but are vital for correctly identifying specific patient groups in practice
 quality improvement (PQI) and research efforts require changes in clinical processes
 to assure validity. This often implies a cultural shift to prioritize recording data in
 recoverable formats.
- Data Structure The cost of free text is high. Lack of standardized structure for entry undermines ability to automate extraction of key data elements from text fields such as notes. To assure accurate, high volume, electronic extraction of key data elements standardized methods for encoding key data elements need to be defined and implemented in clinical processes.
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- 187 large commercial and governmental datasets such as the National Cancer Database188 Base.
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191 **Process and system changes**

- 192 In reviewing current practices, a number of obstacles stand in the way of obtaining the amount
- and quality of data needed to make substantial progress. The following outline provides a view
- that is geared towards identifying means of overcoming them.
- 195 (1) Failure to collect necessary structured data
- 196 (2) Lack of data standardization
- 197 (3) Inability of different electronic data systems to communicate.
- 198 Within each of these broad categories, it is useful to provide a finer-grained view of how
- 199 different aspects of our clinical and electronic environments contribute to the overall difficulty
- in achieving the data collection and use that we seek.
- 201 (1.a) Commercial System Databases

Focus for development of commercial systems that store the range of data needed for clinical 202 data repositories is often on the user interfaces rather than on the back-end databases. The 203 situation is similar to a clinical focus on data required to treat the day's patients and support 204 205 billing documentation with few resources devoted to standardizations and optimizations to 206 increase big data extractions. Individual systems may use multiple loosely connected databases, 207 complex compound keys, lack of indexing, poorly designed schema, lack reasonable security, or 208 use non-standard database technologies. Vendors may also refuse to provide end-users access 209 to extract their own data. Some commercial systems are much better than others, so end user 210 experience is variable.

211 (1.b) Diagnosis and staging

212 Correct usage and quantified entry of diagnosis and staging information is central to many PQI and research efforts. For example, incorrect entry of primary disease codes (e.g. prostate 185, 213 214 C61) when treating subsequent bone (C79.51), brain (C79.31) or lung (C78.00) metastasis and 215 failure to utilize functionality in radiation oncology information systems (ROIS) to connect primary and metastatic diagnosis undermine the ability to use these codes to correctly identify 216 patient groups by codes. Failure to utilize functionality in ROIS connecting treatment courses to 217 218 these codes undermines ability to connect treatment elements (e.g. DVH metrics) to patients. 219 The cost of not taking a few seconds to select ICD-O (International Classification of Diseases for Oncology) values linked to ICD9 (International Classification of Disease, revision 9) and ICD10 220 (International Classification of Disease revision 10) in the ROIS means that subsequent 221 questions about disease site location become prohibitively expensive to answer because of the 222 223 manual effort required to retrospectively revisit the chart. When survival information is obtained from EHRs, failure to utilize functionality in ROIS to enter staging information 224 undermines ability to factor staging into survival, recurrence and other factors. Typically, EHRs 225 do not have functionality for quantifying diagnosis and staging information according to 226 guidelines (e.g. AJCC, FIGO) or to connect primary and metastatic disease. On the other hand, 227 228 ROIS generally do, but frequently this functionality is not utilized fully as part of clinical practice.

229 (1.c) Outcomes

230 Patient outcomes such as toxicity and disease site status (e.g. recurrence) are frequently 231 entered into electronic records as free text using unstandardized terminology. This renders them unavailable for automated electronic extraction. Lack of standardizations 1) for which 232 toxicities are routinely measured, 2) how treatment site categorizations are named (e.g. breast 233 tangents, breast tangents plus supra-clavicular field, breast tangents plus supra-clavicular field 234 plus internal mammary node field, etc), 3) how categorizations for disease site status are 235 named (e.g. no-evidence-of-disease, local recurrence) or 4) in use of regular schemas for text 236 237 representation of these key data elements prevent this information from being used to its full 238 value in routine characterization of outcomes for treated patients.

239 (1.d) "As-Treated Plan Sums"

240 To assess correlation of outcomes with dose volume histogram (DVH) metrics, it is necessary to first create treatment plan sums corresponding to the plans and number of fractions treated, 241 reflecting boosts, plan revisions and incomplete treatments. When these "as treated" plan 242 243 sums (ATPSs) are created as part of routine practice, then automated solutions for calculating dose-volume histograms metrics becomes possible. Unfortunately, often these are not created 244 as part of routine practice, with the result that they must be constructed retrospectively, ad-245 hoc, preventing systematic, automated aggregation. Currently no major commercial system, to 246 our knowledge, has a standard means for reporting ATPSs. 247

248 (2.a) Prescriptions

Electronic prescription summaries that defined dose levels, target structures, number of treatments, fractionation groups (e.g first course, plan revision, boosts, etc) and connection to target structures, organs at risk, treated plans and DVH metrics have been developed by a few researchers [16,17]. These custom solutions were developed to fill the void left by commercial ROISs. Recently ASTRO has suggested a baseline set of guidelines for information that should be included in prescriptions to promote standardization [18]. Similar to ATPSs, commercial solutions and clinical processes often lack ability to retrospectively extract this key information.

256 (2.b) Key Treatment Parameters

Ensuring ability to identify which patients were treated with special technologies and details of those treatments is important to being able to prove their efficacy. Examples include breath hold technologies, radio frequency or radio-opaque fiducials used for positioning, immobilization devices, etc. However, commercial systems and clinical approaches to utilizing those systems are frequently inadequate for retrospectively gathering this data.

262 (3.a) Integration of Treatment Planning System (TPS) with ROIS

263 If systems do not use a common database for TPS and ROIS it is difficult to unambiguously 264 move from the ROIS record of plans actually treated back to specific plans, plan sums and DVH 265 curves in the TPS. Some vendors may even discard DICOM Unique Identifiers for plans from the 266 TPS. 267 (3.b) Integration with EHR

ROIS and TPS systems typically do not integrate with EHR's. Connections may be made through medical record numbers and inferences around dates recorded in respective systems. This is an area where Health Level 7 (HL7) Fast Healthcare Interoperability Resources (FHIR) could significantly improve integration.

272 (3.c) Integration with specialty systems

Treatment devices other than conventional linear accelerators (e.g. brachytherapy, particles, specialty accelerators, MR guided linacs) may provide minimal details back to the ROIS or may use specialty tables in the ROIS that do not integrate well with tables used to manage external beam therapies. This limits the range of questions around treatment details for these specialty modalities that can be addressed at large scale for all patients treated.

278 (3.d) Integration with institutional registry data

Institutions with the American College of Surgeons Commission on Cancer and National Comprehensive Cancer Network (NCCN) designations are required to have medical registries that follow up on cancer patients. Registries document demographics, diagnosis, staging, survival, cause of death and other factors. Registry data is rarely linked to radiation oncology data repositories.

284 (3.e) Integration with public databases

Institutional registries supply data to state registries. Published state analyses are,
unfortunately, many years behind current practice. Although state registries have high volumes
of patients, there is no simple means to connect back to patients to check on the validity of the
data or to investigate impact of cofactors on outcomes tracked in the registries.

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290 Access and Extraction Issues

As radiation oncology has developed, a number of structural issues have arisen that limit clinicians', caregivers' and researchers' access to the data that we do have. Access requires

several key elements: knowledge of the format and schema of the stored data, software thatcan identify and extract the data, and permissions to view and extract the data.

Figure 1 illustrates the level of detail that is needed regarding the treatment of rectal carcinoma patients under three RTOG studies. To combine the data from these trials requires knowledge of how the problem is framed (which clinical data are needed, what are the key elements of those data), how the data are formatted (type of value, allowed values, units, standards if applicable), and the specific software needed to access the data (SQL, RDF triples, spreadsheets).

The issue of framing the medical problem is difficult but rewards are high. The DICOM standard 301 302 (and its radiation therapy extension) has achieved such success in large part due to its 303 structuring of what an imaging study (radiation treatment) is--what are its elements and how 304 are they related [19]. Thus, regardless of the details of the implementation of a procedure, all 305 partners in a communication exchange agree on the essential elements. The definition of such standards in other areas of medicine is rapidly increasing. For example, a relatively commonly 306 used standard for data exchanges between EHR's is the standard Health Level 7 (HL7). HL7 307 308 version 2 standardized types of data and the allowed values and permitted organizations and 309 vendors to develop software for the reliable interchange of certain data. However, it was considered to be quite limited, and version 3 was built around the Reference Information 310 311 Model which was a much more robust view of healthcare in general [20]. Even more recently, they have started developing HL7-FHIR which instantiates an even more up-to-date view of 312 medical practice, but also highlights the importance of appropriate technology. HL7-FHIR is 313 built upon the REST specification that is the current industry standard for web-based 314 applications [21]. Other data standards, such as the NCI thesaurus [22], provide additional 315 resources that facilitate the development of software for access and extraction of data. 316

With rare exception, major vendors of ROIS, TPS and EHR systems, store information in relational databases. A few types of large volume objects (e.g. DICOM images) are stored in files that are referenced in the relational databases. Custom extractions from databases are carried out using structured query language (SQL). SQL queries may have dialectical variation among

relational database systems (e.g. Oracle, Microsoft SQL). Ideally, relational databases are designed with categories of data grouped into tables and views (stored SQL query results) reflecting an overall view of the procedure itself. They also use normalization strategies to prevent redundant information, reduce complexity in SQL queries and increase performance in retrieving data. Secure data retrieval requires granting read access to specific authenticated network accounts. Access may be controlled at the level of the database, table or views. Skill with SQL is essential to any staff constructing or extracting data for a data repository.

Application programming interfaces (APIs) are provided by vendors of many TPSs. These may be used to gather subsets of information stored in the ROIS database or elements only calculated at run time in the TPS (e.g. DVH curves for some systems). APIs allow custom software applications to be constructed by users that interface with the TPS. Access is controlled by end user system administrators, subject to constraints of the commercial system. Clinical staff members with coding skills are necessary for effective use of API's.

Legacy issues with vendor changes to both database and API structures are an issue for groups automating extraction from electronic records systems. Effort to re-write queries and scripts when systems are upgraded can be substantial.

337 Patient reported outcomes (PROs) are important outcome measures and their routine monitoring during cancer therapy has been demonstrated to improve survival [23]. However, 338 339 use of paper based rather than electronic systems are more common. Electronic systems are 340 significantly better for making the data accessible, but require substantial effort in setting up 341 systems and arranging for staffing resources to assist patients with completing electronic surveys is required. In addition, lack of standardization in instruments to be used, redundant 342 343 questions between surveys, excessive length diminishing patient willingness to participate, and 344 question formats and logic that translate poorly to electronic systems already used in patient work flows are issues for generalized use of PROs. 345

Diagnostic images are stored on Picture Archive and Communication Systems (PACS) in Digital Imaging and Communication in Medicine (DICOM) format and accessed with DICOM servers. Graphic user interfaces for clinical use are not well suited to large volume, batch access of sets

of patient images. The objective in utilizing these resources in connection with BDARS is not creation of a parallel PACS. Instead, when large sets of images are identified for utilization in a study, e.g. developing predictive radiomics measures for a disease type, downloading a large specific set of images for batch processing is needed. Negotiating access is the primary barrier.

353 Finally, it is important to discuss the role that legal and commercial considerations play in 354 limiting access to data. The Health Insurance Portability and Accountability Act of 1996 355 requires certain standards to be met when exchanging private health information. The standards depend on the intended use of the data, for example, clinical decisions, insurance 356 357 coverage, quality improvement and research. They also depend on the entities exchanging the 358 information. These standards add time, effort and new procedures to any effort to obtain data 359 access. Intra-institutional exchange, for example between a departmental data repository and the hospital EHR, is in general easier than between institutions, but even that type of 360 361 transaction usually requires some level of administrative oversight and/or procedure. In 362 addition, storing data in a clinical data repository for possible future research can be viewed as problematic under national ethics guidelines for human research [24]. Overall, it is difficult to 363 364 make any broad statements or recommendations regarding these issues since they are, to 365 some degree, institution- and use-specific. In addition, how the regulations are interpreted is 366 evolving, particularly in response to some of the national healthcare programmatic initiatives such as the Affordable Care Act. 367

368 Selecting technologies

369 The objective is to use the treatment data, rather than to utilize a novel database technology.

370 Selecting database technologies which minimize investment overhead and risk while

371 maximizing productivity and interoperability for addressing particular tasks requires careful

372 consideration [25,26].

At a high level, four process steps can be considered and technology choices should be madefit-for-purpose for these steps.

375 1. Capture of treatment data

The primary use for health care data is delivery of patient care. Health care database

technology is often vendor dependent and under regulatory oversight. For structured data

elements (e.g. record and verify, electronic health records, outcome) relational databases are

379 the most common technologies. Images and related objects such as treatment plans and record

are generally object stores (e.g. PACS) with a relational schema containing object pointers.

381 2. Extraction

Since the primary use sources have to be taken as-is, the extraction technologies providing
connectors to these primary sources should be able to handle many different sources and
formats including all common relational sources. They should be able to handle non-relational
sources including "databases" that researchers and physicians often use (e.g. Excel, SPSS) and
include JSON and XML support as these are common export format for more technical users.
Ideally, the technology can be extendible to support common medical standards (HL7v2, HL7v3,
HL7 FHIR and DICOM) as needed.

A wide range of programming languages and standard database import tools are frequently used. These have the advantage of hiding very little from the user. There are also commercial and open source software systems intended to reduce the technical skill requirements for users with the trade-off of obscuring details about the extraction, cleaning and loading processes. Since primary sources change and extraction tools generally expand and change over time, a crucial requirement is versioning. Users of technology should be able to store different versions of the extraction scripts and configurations so that subsequent users can re-use their solutions.

396 3. Transformation, integration and storage

For successful secondary use, the primary use sources need to be combined, integrated and common data elements mapped on each other. An example is the combination of ROIS/EHR data (diagnosis, comorbidities, prescriptions, treatments, follow-up), Record and Verify data (radiotherapy treatment) and DICOM data (imaging/plan). This transformation and integration is generally the most time consuming task of the process. Knowledge of the primary sources and of the secondary use data model is a requirement for staff using the tool. Again, versioning and manageability is crucial as sources change and sharing transformation scripts with others is

needed for work to not be duplicated. Defining distinctions between data element categories
and relationships means mapping the raw values onto a schema. For example, a schema needs
to be applied so that we can inform our analytics programs if an extracted value "30"
corresponds to a dose, an age, a day of the month, etc. and how that value relates to other
information e.g. toxicity, survival, PROs, treatment dates, etc.

409 From a technology standpoint two main approaches exist.

410 Schema-On-Aggregate (aka schema-on-write): Upon extraction each data element from each source is considered more or less separately, transformed 411 and mapped to the secondary use data model and then written in the secondary 412 use data store. Schema-on-aggregate has as its main benefit that it often re-uses 413 the knowledge contained in the primary use schema and forces one to decide up 414 front how to map data items and think about transformation for each data 415 416 element. The end-result is often a data store with a structured schema. Relational databases are widely used for this approach owing to their speed, 417 ease of integration with other systems and large pool of talent for use. Non-418 relational databases (e.g. object stores, graph databases and triple stores) have 419 also been used in some research settings. 420

421 Schema-On-Query (aka schema-on-read): The secondary use data model is 422 applied when the secondary user requests, or queries, the data from the 423 secondary source. In a schema-on-query system the data is stored from the primary source "as-is" and by necessity this is a non-relational store (e.g. a data 424 425 lake). An example is Apache Hive which can be used for SQL-like schema-on-426 query for Apache Hadoop. NoSQL databases, such as MongoDB or CouchDB, are 427 another example. The main benefit of this approach is that the transformation and secondary use data model can be defined fit-for-purpose, and different for 428 different use cases. Also all primary use data can be stored immediately for later 429 430 secondary use. The main drawback is that knowledge of original schema is often 431 not available by the time the data is used and that data is stored without de432 identification. Variability in nomenclature for key data elements, relationships

element are internally consistent and stable.

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custom code for each to enable programmatic extraction. Care must be taken to

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Note that many solutions allow a combination of the above approaches, with some data
elements stored in a schema generation upon aggregate and some stored "as-is" for schema at
a later time point. In that case, key data elements are often duplicated into the secondary use
storage.

and formats among the various "as-is" sources requires creating and maintaining

ensure consistent meaning at the time of data entry so that contents of an

442

443 Secondary use application

Secondary use of subsets of data extracted from BDARS to address specific research or clinical 444 questions is a common use case. The secondary user usually has defined their own data model, 445 store and the application to analyze the data. The technology choices made by secondary users 446 vary widely and limited influence exists especially if the secondary user is external to the 447 primary use institution. The main job of technology here is to provide the secondary end-user 448 449 with a dataset and format which he or she can use (often called a data mart). Typical requested 450 formats include SQL database dumps, Microsoft Excel, comma (or tab) separated values (CSV), DICOM, HL7 FHIR, HL7v3, HL7v2, XML and JSON. Additionally, data visualization and allowing 451 the end-user to navigate the data store established in the previous step increase the efficiency 452 453 and effectiveness of secondary use. The tools mentioned above generally allow such export to a variety of data formats. Figure 1 illustrates one such use case, a semantic triple store database 454 (a.k.a. Resource Description Framework) was applied for the purpose of combining datasets 455 from several clinical trials. Semantic triples can be used to define a range of relationships 456 457 between objects (e.g. PTV \rightarrow is a type of \rightarrow target structure).

458

Specific recommendations for work flows and standardizations 459 460 1) Diagnosis and staging data should be entered into quantified fields in accessible, electronic systems that 461 have quantified fields for staging elements and overall staging, and staging guideline 462 0 463 system used (e.g. American Joint Committee on Cancer (AJCC)) 464 0 ensure correct selection of staging from component elements provide explicit linkage to treatment courses and plans used to treat 465 0 link metastatic diagnosis (e.g. C79.51, Secondary malignant neoplasm of bone) to 466 0 diagnosis for originating sites (e.g. C34.1, Malignant neoplasm of upper lobe, 467 bronchus or lung) 468 469 470 In the current vendor landscape, the ROIS is frequently the only system in the clinical 471 process workflow meeting these objectives. 472 2) Nomenclature standardizations recommended by AAPM Task Group 263 should be adopted 473 into routine practice. These define standardized nomenclature for structure, target and 474 DVH metric naming to promote ability to automate aggregation [27]. 475 476 477 3) Course cumulative as-treated plan sums should be constructed as part of routine practice. Since more than one image set may be used in the construction of the ATPS's, and relative 478 positioning of structures may vary between sets, using the image set providing the best 479 representation for the clinical evaluation carried out for treatment is currently the most 480 viable approach. 481

482

4) Toxicities, recurrence and PRO outcomes need to be routinely collected as quantified fields 484 (instead of free text fields) in accessible electronic systems. Standardizations for specific 485 items and values are needed. This includes, for example, definition of recurrence 486 nomenclature. Ability to automatically recover these values from the electronic record is 487 important.

488

489 5) Detailing of key data elements and relationships (i.e. an ontology) is needed for a broad 490 range of practice quality improvement and translational research efforts. An initial set, drawn 491 from experience in constructing BDARS, is presented as an appendix to this paper. Success in 492 gathering this information requires that clinical systems should be utilized to ensure ability to accurately aggregate these elements and relationships from the electronic record (ROIS, TPS, 493 494 EHR). Ideally, professional societies such as ASTRO, AAPM, ESTRO and CARO would combine 495 efforts to eventually take the role of maintaining standardized ontologies to promote interoperability among institutions and commercial systems. Combining the ontology presented 496 497 in the appendix with related ontologies would be a valuable step toward a common standard 498 [28,29].

499

500 6) In addition to demonstrating adherence to standard quality metrics, clinical entities will face 501 increasing demands for demonstration of the value of the care they deliver as medicine in the 502 transitions from fee for service to value based payments. Success in the value based payment 503 environment will require the ability to conduct on-demand analysis of patient and tumor 504 characteristics, all aspects of treatment delivery, outcomes, and cost of care.

505

506 We note that the task of creating ATPSs (item 3) needs to begin as soon as possible, guided by 507 clinical judgment, in order to replace complete lack of data with reasonable data. In addition, 508 further refinement is needed. Collaborations between professional societies, vendors and 509 clinical trials groups for defining standards for the end-of-treatment dose composite are 510 needed. Issues include means to quantify quality of the composite, identifying source images, 511 identifying trade-off decisions in image registrations, uncertainties in structure dosimetric 512 measures when multiple image sets are used, and realistic appraisal of the role of image 513 deformation.

514 **Examples of Clinical Data Repositories**

515 Several groups have been actively engaged in construction of clinical data repositories (CDR), 516 also known as data lakes and Big Data Analytic Resource Systems (BDARSs). These systems 517 become important components for both research and clinical practice efforts in their clinics. 518 Practical recommendations from this group have been grounded in the experience of 519 constructing, using and sharing these systems. Brief summaries of several are highlighted to 520 convey the scope and volume of these resources.

The University of Michigan Radiation Oncology Analytics Resource (M-ROAR) automates 521 • aggregation of electronic data from the Treatment Planning System (TPS), Radiation 522 523 Oncology Information System (ROIS), Electronic Health Record (EHR) and other databases for all patients treated. Data types include demographics, treatment and 524 dosimetric data, chemotherapy, toxicities, comorbidities, labs, medications, encounters 525 and patient reported outcomes (PROs). The system contains records for over 20,000 526 patients. Key data elements are extracted utilizing a combination of SQL queries, TPS 527 application programming interface (API) based scripts and custom code to extract and 528 process data from multiple source systems [25]. 529

The UCLA Clinical Informatics Management System (CIMS) consists of three major 530 modules: a physician interaction module that interacts closely with EHR, a physics 531 532 parameter module that handshakes with PACS systems, treatment planning and delivery stations for quantitative value collection and exchange, and a patient reported outcome 533 management system (Patient Reported Outcomes Measurement Information System, 534 PROMIS) with a web/mobile portal. The physician interaction module supports 535 comprehensive query for collection and integration of radiotherapy relevant 536 537 information from other departments. The patient reported outcome management

538 module consists of a front-end with site-specific patient-oriented Common Terminology 539 Criteria for Adverse Events (CTCAE) questionnaires tailored to patients. As of now, the 540 registry contains records for 1790 definitive prostate treatment, 209 post-operative 541 prostate treatment, 1950 breast, 663 lung, 531 brain metastasis, 484 GYN, 424 glioma, 542 409 meningioma, 209 rectum, 151 metastatic bone, 164 trigeminal, 111 pancreas, and 543 over 3000 general cases [30].

- The Ohio State University Radiation Oncology Department's "Quality Database" has 544 been designed to serve as a data aggregation platform to capture clinical, technical, and 545 546 health outcome data on all patients who receive radiation treatments. All data are stored in a REDCap database. Smart texts have been implemented in EHR to enable 547 automated capture and extraction of discrete data elements such as adverse events 548 549 from provider notes. The dosimetry data for radiation therapy are extracted via TPS's API. Demographics, diagnosis, tumor biomarkers, surgery, systemic therapy, radiation 550 therapy, and adverse events constitute the collected data and provide means for 551 determining effectiveness of treatment modality. The Quality Database currently 552 contains 3385 patients and is being populated prospectively with new patient data. 553
- Oncospace: Johns Hopkins University developed a comprehensive data collection and 554 data repository system [31]. The system consists of a network of data collection 555 systems (ROIS, clinic computer terminals, mobile devices, hospital EHR) that provides 556 data that is transformed and loaded into a SQL database. Using a federated database 557 approach (including University of Washington, University of Virginia, Odette Cancer 558 Center-Sunnybrook), each institution has implemented compatible schemas so 559 560 federation-wide gueries will succeed. This approach has the advantages of "crowdsourcing" ideas and technology and allowing each institution to keep control of 561 their data while still permitting individual flexibility. 562
- The Veterans Health Administration (VHA) developed a pilot Radiation Oncology
 Practice Assessment (ROPA) program to assess the quality of radiotherapy across the
 entire VHA network with 40 institutions participating [32]. Data types include quality

566 metrics targeted at workup, diagnosis, treatment planning, delivery and follow-up. The 567 gathered quality metrics were developed by the VHA in partnership with ASTRO for 568 locally advanced non-small cell lung cancer, limited stage small cell lung cancer, and 569 intermediate and high-risk prostate cancer. Data extraction for the initial pilot project 570 will be completed in 2018. At that time ROPA is anticipated to contain 45,000 scores for 571 49 metrics aggregated from approximately 2,000 patients.

Large data sets from sources outside of radiation oncology are now available for 573 analysis. Waddle et al. recently published utilization data derived from insurance 574 records from a commercial warehouse (Optum Labs) to examine treatment technologies 575 used (proton, stereotactic body radiotherapy, IMRT, 3D, other) by diagnosis code used 576 in billing records. The data base contains utilization data on a subset of 474,533 577 radiation oncology patients from a larger database of over 100 million insured lives. 578 However, connection of this data to clinical outcomes and other cofactors was pending 579 at the time of that analysis [33]. 580

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586 **Recommendations for next steps needed to improve data availability.**

587 Adopting national standards

As discussed above, an important aspect of data exchange is employing a generally recognized view of the medical process. HL7 FHIR is an emerging standard and one that has the crucial elements of (a) flexibility, (b) state-of-the-art technologically, and (c) widespread support [34]. As this standard is just not being formalized, this is an excellent time for the radiation oncology community to support efforts to develop radiation oncology-specific resources for this standard [35].

594 Increasing multi-institutional collaborative efforts

595 Real, effective standards emerge from being actively engaged in exchanging data with outside 596 groups as part of more frequent collaborations. Professional and government grant support for 597 research efforts that develop and proof these standards as by-products are important to their 598 emergence.

Included in this effort is need to facilitate information exchanges that support re-treatment. As patients are able to survive longer with cancer, likelihood of visiting more than one center for subsequent treatments increases. Clinical process and data exchange standardizations needed to facilitate these exchanges also support collaborative efforts.

603

604 Links to institutional registries

Institutions which are members of the National Comprehensive Cancer Network (NCCN) are
required to have access to a registry which carries out longitudinal follow-up on a few key data
elements (e.g. survival, cause of death) for treated patients. EHR database records may be
substantially different from registry database records. Providing electronic access registry
databases provides opportunities to synchronize data sources in constructing big data analytics
resource systems.

611

612 Support for Public Data Sets

The value of producing data sets that can be publicly shared (without compromising PHI) has been heralded by several authors. [36-38]. There is growing interest from funding agencies for publicly funded research to produce publically available datasets. Similarly, an increasing number of journals require publication of datasets accompanying findings. Recently Medical Physics has introduced a special publication category just for data sets. Principles for ensuring that data are findable, accessible, interoperable, and reusable (FAIR) for public access of data sets have been set out by Wilkinson *et al.* [39] and others [40].

The National Cancer Institute has recently begun to implement a Cancer Research Data Commons which meet the standards of FAIR. In their announcement, they echo a number of the themes that we have set forth in this article. This is clearly a propitious time for radiation oncology to join with others in the oncology fields to make these sorts of community-wide efforts more productive [41].

625 Informatics Training

Clinical staff bring great value to informatics efforts because of the depth of their clinical
domain knowledge with respect to key data elements, their inter-relationships, clinical
processes by which data is entered, end user expectations for meaning, etc. The set of clinical
staff that take on expanding their informatics skills to include database, programming,
statistical analysis and machine learning also improve ability to develop practical solutions
bridging needs between the larger number of specialists entirely focused in either the clinical or
informatics domains.

633 Conclusions

We have laid out an argument for why it is important for the radiation oncology community to improve the means by which we can collect, share and use the data that we encounter every day. However, for various reasons, much of this data remains inaccessible to us in a format that makes it easy for us to transform data to knowledge.

The technological challenges to implementing a community-wide system of data collection, 638 sharing and usage are formidable but the tools have been or are currently being developed. 639 640 More difficult is developing the collective will to make it happen. Such a change in our clinical 641 behavior and workflow requires buy-in from everyone, including clinic staff, physicians, and 642 vendors. It is our hope and expectation that this sea change has already started to occur as diffuse networks grow in size and analytic power. It is necessary to do so if we are to continue 643 to be at the forefront of harnessing technological advances to improve the treatments that we 644 provide our patients. 645

646

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- 651
- 652 Acronyms:

- 653 AAPM: American Association of Physicists in Medicine
- 654 AJCC: American Joint Committee on Cancer
- 655 API: Application Programing Interface
- 656 ASTP: As Treated Plan Sums
- 657 ASTRO: American Society for Radiation Oncology
- 658 BDAR: Big Data Analytic Resource Systems
- 659 CARO: Canadian Association of Radiation Oncology
- 660 CDR: Clinical Data Repository
- 661 CER: Comparative Effectiveness Research
- 662 CTCAE: Common Terminology Criteria for Adverse Events
- 663 DB: Database
- 664 DICOM: Digital Imaging and Communications in Medicine
- 665 DVH: Dose Volume Histogram
- 666 ESTRO: European Society for Therapeutic Radiation Oncology
- 667 EHR: Electronic Health Record
- 668 FAIR: Findable, Accessible, Interoperable, and Reusable
- 669 FHIR: Fast Healthcare Interoperability Standards

- 670 FIGO: International Federation of Gynecology and Obstetrics
- 671 HIPAA: Health Insurance Portability and Accountability Act
- 672 HL7: Health Level 7
- 673 ICD-O: International Classification of Diseases for Oncology
- 674 ICD9: International Classification of Diseases, Ninth Revision
- 675 ICD10: International Classification of Diseases, Tenth Revision
- 676 JSON: JavaScript Object Notation
- 677 NCCN: National Comprehensive Cancer Network
- 678 NIH: National Institutes of Health
- 679 OIS: Oncology Information System
- 680 PACS: Picture Archive and Communication Systems
- 681 PHI: Protected Health Information
- 682 PQI: Patient Quality and Improvement
- 683 PRO: Patient Reported Outcome
- 684 PROMIS :Patient-Reported Outcomes Measurement Information System
- 685 REDCap: Research Electronic Data Capture
- 686 ROIS: Radiation Oncology Information System
- 687 RCT: Randomized Controlled Trial
- 688 ROILS: Radiation Oncology Incident Learning System
- 689 RTOG: Radiation Therapy Oncology Group
- 690 SQL: Structured Query Language
- 691 TPS: Treatment Planning System
- 692 XML: Extensible Markup Language

693 VHA: Veterans Health Administration

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- 788
- 789
- 790 Appendix

791 Key Data Elements and Relationships: A Radiation Oncology Translational Research Ontology

792

793 We have defined of a common set of key data elements and relationships important to a broad range of 794 patient quality improvement and translational research efforts. Ranking treatment information for effectiveness requires a broad scope of information types: Radiation Treatments, Surgery, Outcomes, 795 796 etc. While it is desirable to have all the data readily available, that is not a practical starting point. Our 797 objective here is to define a minimal set of information needed to handle frequently encountered 798 questions as a common use starting point. With that, technical and procedural efforts attempting to 799 automate electronic aggregation supporting Big Data efforts can use these recommendations as a guide. 800 Optimally professional organizations (e.g. AAPM, ASTRO, ESTRO, CARO) would establish an official listing

801 of key data elements and relationships. Our intention here is to provide a practical starting point from 802 our experience in aggregations from multiple source systems.

The listing of key data elements and relationships define an explicit conceptualization of a body of formally represented knowledge about Radiation Oncology, i.e. an ontology [42] The listing provided here was based on the ontology developed for M-ROAR [25] and expanded as an outgrowth of discussions at the Practical Big Data Workshop. Incorporation of the ontology into a programmatic form using Ontology Web Language (OWL) is underway.

Classes (⊕) of information, list key data elements (aka properties) denoted by one of three symbols (●,
 ⊙, ○). Most elements (●) do not require special consideration for protection of patient health
 information (PHI). Elements that contain PHI (⊙), are problematic for data sharing or storage in cloud
 based systems. Alternatives (○), containing, reduced information, may be sufficient for a wide range of
 collaborative efforts or cloud based storage.

For example, dates are a type of patient health information (PHI) that institutional review boards (IRB) will not allow for many applications. For a wide range of investigations, detailing temporal relationships between events is important. Recording the patient's age at the event, rather than the date for the event is an alternative. For example, if the date of an event is 3/2/2013, and the patient's date of birth is 8/17/1967, then the patient's age at the time of the event, to three decimal places (Decimal F3), is 45.541. This is sufficient resolution to differentiate day on a timeline and meets requirements for protecting PHI.

Several key data elements typically are not present as distinct values in source data systems but have to
be programmatically derived (X) from other elements. For example, the age of the patient at the time

of an event is derived from date of birth and date of the event. Starred (*) items indicate particular need
 for recommendations of standardized values recommendations from professional societies.

824 When elements have only one instance they are indicated by the name of the class or element (e.g.

825 DateOfBirth, Patient). When there may be more than one instance of an element, this is indicated by

specifying a list of elements of this class (e.g. List<Course>).

827 Relationships among classes are categorized as $Parent(\Leftrightarrow)$, $Child(\Rightarrow)$, $Sibling(\Leftrightarrow)$ or $Property(\blacksquare)$. Parent-828 Child are dependent relationships: a parent class object is referenced in each instance of a child class 829 object. Sibling relationships are tracked if elements exist but do not imply dependence. Sibling 830 relationships rather than parent-child relationships may be selected when the current state of the data 831 will not practically support the dependent relationship. For example, Prescriptions are used in sibling 832 relationships with respect to TreatedPlans because the current state of electronic data is inadequate to 833 assure consistent mapping. Property relationships are used when class incorporates a set of elements 834 grouped under a single concept.

835 Patient - \oplus 836 O PatientMRN (String) -: Medical Record Number • PatientGUID (String): Generalized Universal Identifier that can be used in cloud 837 based storage, when PatientMR is not. 838 839 \odot DateOfBirth (Date) 840 O YearOfBirth (Int?) ₩ 841 DateLastSurvivalCheck (Date?) 842 ● AgeAtLastSurvivalCheck (Decimal F3) 爰 843 DateOfDeath (Date?) ● AgeAtDateOfDeath (Decimal F3) 署 844 845 IsAlive (Bool) – Status at last at Last Survival Check Date *CauseOfDeath (String) – Need for standardized list 846 847 Gender (String) 848 Race (String) Ethnicity (String) 849 850 851 Child class relationships

852	List <radiation course="" therapy=""></radiation>
853	⇔ List <prescription></prescription>
854	⇒ List <diagnosisandstaging></diagnosisandstaging>
855	➡ List <treatedplan></treatedplan>
856	List <patienttreatmentoutcome></patienttreatmentoutcome>
857	➡ List <patientreportedoutcome></patientreportedoutcome>
858	➡ List <planningstructureset></planningstructureset>
859	➡ List <healthinformation></healthinformation>
860	⇔ List <lab></lab>
861	➡ List <medication></medication>
862	➡ List <image/>
863	List <chemotherapy course=""></chemotherapy>
864	⇒ List <surgical procedure=""></surgical>
865	➡ List <pathology></pathology>
866	➡ List <charge></charge>
867	\mathbf{O}
868	
000	
869	
870	courses
8/1	 CourseName (String) NT CourseName (String)
8/2	• NTXSessionsInCourse (Int) # – Each treatment episode is a session, sessions used for
873	imaging only are exclude from the count
874	 DateFirstTreatment (Date)
875	O AgeAtFirstTreatment (Decimal F3) ₩
876	 DateLastTreatment (Date)
877	● AgeAtLastTreatment (Decimal F3) 器
878	
879	Sibling Class Relationships
880	⇔ List <prescription></prescription>
881	List <chemotherapy course=""></chemotherapy>

882	⇔ List <surgical procedure=""></surgical>
883	
884	Child class relationships
885	➡ List <treatedplan></treatedplan>
886	List <diagnosisandstaging> - Typically only one per Course</diagnosisandstaging>
887	List <patienttreatmentoutcome> - Typically only one per Course</patienttreatmentoutcome>
888	⇔ List <charge></charge>
889	\mathbf{O}
890	Parent Class Relationships
891	← Patient
892	
893	Prescription : The prescription needs to fully convey the intent of the physician for the treatment
894	plan. The Course contains a list of prescriptions
895	Name (String)
896	NTxSessions (Int)
897	 NTxPerDay (Int)
898	 DaysBetweenTxSessions (Decimal)
899	 StartOnNthDayFromCourseStart (Int) 光
900	 StartOnNthSessionInCourse (Int) 発
901	 RxDoseUnits (String) – "cGy" or "Gy" or "CGE"
902	 IsCourseCummulativePrescription (Bool) 第 – Only one value of True per Course
903	
904	
905	Class Property Relationships
906	List <prescriptiondoselevel></prescriptiondoselevel>
907	List <prescriptiondvhobjectives></prescriptiondvhobjectives>
908	
909	Sibling Class Relationships

910		⇔ List <treatedplan></treatedplan>
911		
912		Parent Class Relationships
913		🗢 DiagnosisAndStaging
914		🗢 Patient
915		⇐ Course
916		
917		C
918	\oplus	PrescriptionDoseLevel
919		 RxDose (Decimal F3)
920		 RxStructure (String) – AAPM TG263 compliant name
921		 RxPointName (String)
922		
923		Parent Class Relationships
924		← Prescription
925		
926	\oplus	PrescriptionDVHObjectives
927		 Structure (String) – AAPM TG263 compliant name
928		 DVHMetric (String) – AAPM TG263 compliant name e.g. Max[Gy], V20Gy[%]
929		• Constraint (String) - allowed values are =,<, \leq ,>, \geq , ALARA
930		• Value (Decimal F3) – null if constraint is ALARA
931		
932		Class Property Relationships
933		Prescription
934		
935	\oplus	DiagnosisAndStaging
936		 StagingSystem (String) - e.g. AJCC 7, FIGO
937		 ICD9Or10 (String)

938		 ICD0 (String) – Defines location of disease
939		 Laterality (String) – Left, Right, Bilateral
940		• Overall Staging (String): e.g. IIa, X,
941		 T (String)
942		• N (String)
943		• M (String)
944		• P (String)
945		• G (String)
946		 OtherStagingComponents (String)-Staging components other than T,N,M,P,G
947		 PrimaryOrMetastatic (String)
948		0)
949		Child Class Relationships
950		PatientTreatmentOutcome
951		DiseaseSiteStatus
952		M
953		Parent Class relationships
954		PrimaryICD9Or10? – If Metastatic, indicate Primary DiagnosisAndStaging element
955		⇐ Course
956		Patient
957		\mathbf{O}
958		Ĕ
959	\oplus	DiseaseSiteStatus
960		 DateOfStatus (Date)
961		 AgeAtDateOfStatus (Decimal F3) 発
962		 *Status(String) – Need standardized list e.g. (No Evidence of Disease, Local Recurrence,
963		Regional Recurrence, Distant Recurrence)
964		
965		
966		

967	\oplus	TreatedPlan : Every course has a list of treated plan objects. One table for all types of plans defining
968		key elements to track. This simplifies mixed modality tracking e.g. External + Brachy and handling of
969		individual plans vs plan sums. Only plans actually treated are tracked. Details of actual vs number of
970		fractions delivered are tracked.
971		 PlanName (String): Corresponds to PlanID in ARIA
972		 *TreatmentAreaClassifier (String) : e.g. Head and Neck, Lung_L, Breast_R+SC
973		• TPSSourceSystem (String) 発
974		IsCourseCummulativePlan (Bool): The plan or plan sum(ATPS) represents all plans treated in
975		the course
976		• IsPlanSum (Bool): The dose associated with the plan is created by summing dose from other
977		plans
978		
979		 DateOfFirstPlanTreatment (DateTime)
980		● AgeAtFirstPlanTreatment 器
981		 DateOfLastPlanTreatment (DateTime)
982		• AgeAtLastPlanTreatment 発
983		
984		 PrimaryTxDeliveryFacility (String) – Facility where most of plan fractions were delivered
985		 PrimaryTxDeliveryMachine (String) – Machine on which most of the plan fractions were
986		delivered
987		 NFractions_Planned (Int)
988		• NFractions_Delivered (Int)
989		 TotalDose_Planned (Decimal) – Dose planed for highest dose structure e.g. PTV_High
990		 TotalDose_Delivered (Decimal) – Dose delivered for highest dose structure e.g. PTV_High
991		 TotalDose_Units (String) – Gy, cGy, CGE
992		
993		● UsedFiducials (Bool) 発
994		 FiducialType (String) – Gold, Calypso, Carbon
995		 UsedBreathMotionControl (Bool) 発
996		 BreathMotionControlType (String): SDX, ABC, Compression
997		

998	 MeanSessionTimeMinutes(Int) 発
999	 MeanSessionBeamOnTimeMinutes (Int) 発
1000	 MeanSessionImagingTimeMinutes (Int) 光
1001	
1002	● NImages_MV (Int) 発 - Total number of MV images for all sessions treating this plan
1003	● NImages_kV (Int) 発 - Total number of kV images for all sessions treating this plan
1004	● NImages_CBCT (Int) 器 :Total number of CBCT for all sessions treating this plan
1005	• NImages_MR (Int)
1006	0
1007	List <supplementaltreatmentdetail></supplementaltreatmentdetail>
1008	
1009	Sibling Class Relationships
1010	⇔ Prescription
1011	List <images> - Image Class Objects related to the TreatedPlan e.g. CBCT, kV</images>
1012	
1013	Child Class Relationships
1014	⇒ PlanningStructureSet
1015	➡ List <dvhcurve></dvhcurve>
1016	⇒ List <dvhmetric></dvhmetric>
1017	➡ List <patientpositioningdevice></patientpositioningdevice>
1018	TreatmentPlanDetails_XRT
1019	TreatmentPlanDetails_Brachy
1020	TreatmentPlanDetails_Particles
1021	⇒ PlanningStructureSet
1022	
1023	Parent Class Relationships
1024	← Patient
1025	← Course
1026	□ ComponentOfATPS (TreatedPlan) - Plans that are components of ATPS link back to the ATPS

1027		
1028		
1029	\oplus	PlanningStructureSet
1030		 StructureSetName (String)
1031		● ImageModality (String)
1032		 DateOfImageAcquisition (Date)
1033		• AgeAtImageAcquisition (Decimal F3)器
1034		• DICOMImage_UID (String) DICOM_UID of image use for the plan. In the Image list attached
1035		to the patient.
1036		 DICOMPlan_UID (String)
1037		O DICOMStructure_UID (String)
1038		O DICOMDose_UID (String)
1039		 PatientPosition (String)
1040		
1041		Parent Class Relationships
1042		← Patient
1043		🗢 TreatedPlan
1044		
1045	\oplus	PatientPositioningDevice
1046		 *DeviceCategory (String) – Need standardized list
1047		 DeviceName(String)
1048		 SetupDetails (String)
1049		
1050		
1051	\oplus	TreatmentPlanDetails_XRT
1052		 List<energymodality></energymodality>
1053		• TotalPlanMU (Decimal)
1054		● UsedIMRT (Bool) 光
1055		● UsedVMAT (Bool) 業

1056	● UsedFIF (Bool) 発
1057	● UsedWedges (Bool) 発
1058	● UsedBolus (Bool) 光
1059	 UsedNonCoplanarBeams (Bool) 発
1060	● NBeams (Int) 発
1061	 NFractionsPlanned (Int)
1062	 NFractionsDelivered (Int)
1063	 List<supplementaltreatmentdetail></supplementaltreatmentdetail>
1064	Parent Class Relationship
1065	TreatedPlan
1066	
1067	
1068	TreatmentPlanDetails_Brachy
1069	 List<energymodality></energymodality>
1070	 NSourcesTotal (Int)
1071	TotalActivity (Decimal)
1072	 *TotalActivityUnits (String)- Need standardized list e.g. MBq, Ci, mCi, GBq
1073	 UsedRadiopharm (Bool)
1074	UsedApplicator (Bool)
1071	
1075	 TotalHDRDwellTimeMin (Decimal)
1075 1076	 TotalHDRDwellTimeMin (Decimal) TotalPDRDwellTimeMin (Decimal)
1075 1076 1077	 TotalHDRDwellTimeMin (Decimal) TotalPDRDwellTimeMin (Decimal) TotalLDRImplantTimeMin (Decimal)
1075 1076 1077 1078	 TotalHDRDwellTimeMin (Decimal) TotalPDRDwellTimeMin (Decimal) TotalLDRImplantTimeMin (Decimal) List<supplementaltreatmentdetail></supplementaltreatmentdetail>
1075 1076 1077 1078 1079	 TotalHDRDwellTimeMin (Decimal) TotalPDRDwellTimeMin (Decimal) TotalLDRImplantTimeMin (Decimal) List<supplementaltreatmentdetail></supplementaltreatmentdetail>
1075 1076 1077 1078 1079 1080	 TotalHDRDwellTimeMin (Decimal) TotalPDRDwellTimeMin (Decimal) TotalLDRImplantTimeMin (Decimal) List<supplementaltreatmentdetail></supplementaltreatmentdetail>
1075 1076 1077 1078 1079 1080 1081	 TotalHDRDwellTimeMin (Decimal) TotalPDRDwellTimeMin (Decimal) TotalLDRImplantTimeMin (Decimal) List<supplementaltreatmentdetail></supplementaltreatmentdetail> Child Class Relationships List<applicator></applicator>
1075 1076 1077 1078 1079 1080 1081 1082	 ■ Jotal#DRDwellTimeMin (Decimal) ■ TotalPDRDwellTimeMin (Decimal) ■ TotalLDRImplantTimeMin (Decimal) ■ List<supplementaltreatmentdetail></supplementaltreatmentdetail> Child Class Relationships ⇒ List<applicator></applicator>
1075 1076 1077 1078 1079 1080 1081 1082 1083	 ■ TotalHDRDwellTimeMin (Decimal) ■ TotalPDRDwellTimeMin (Decimal) ■ TotalLDRImplantTimeMin (Decimal) ■ List<supplementaltreatmentdetail></supplementaltreatmentdetail> Child Class Relationships ⇒ List<applicator></applicator>

1085		
1086		
1087	\oplus	Applicator
1088		 *ApplicatorType (String) Need standardized list e.g. Needle, BrachyCath, TandemAndOvoid,
1089		Cylinder, Mamosite, Savi
1090		 NApplicatorsInserted (Int) 発
1091		● NApplicatorsUsedInTx (Int) 光
1092		\mathbf{O}
1093		Parent Class Relationships
1094		TreatmentPlanDetails_Brachy
1095	\oplus	TreatmentPlanDetails_Particles
1096		 List<energymodality></energymodality>
1097		 UsedPassiveScattering (Bool)
1098		 UsedSpotScanning (Bool)
1099		 UsedEndOfRangeToSpareCriticalOAR (Bool)
1100		List <supplementaltreatmentdetail?></supplementaltreatmentdetail?>
1101		Parent Class Relationships
1102		TreatedPlan
1103		
1104	\oplus	EnergyModality
1105		Energy (String) – Need standardized list e.g. X06, X06FFF, X10, X10FFF, E06, E09, E12, E16,
1106		E20, Ir192, I125, P70, C250
1107		 *Modality (String) – Need standardized list e.g. XRT, HDR, LDR, Proton, CyberKnife,
1108		GammaKnife
1109		
1110		Parent Class Relationship
1111		← TreatedPlanDetails_XRT
1112		TreatedPlanDetails_Brachy

1113		TreatedPlanDetails_Particles
1114		
1115		
1116	\oplus	SupplementalTreatmentDetail
1117		Name (String)
1118		• Value (String)
1119		• ValueType (String)
1120		\mathbf{O}
1121		Parent Class Relationships
1122		TreatedPlanDetails_XRT
1123		TreatedPlanDetails_Brachy
1124		TreatedPlanDetails_Particles
1125		← TreatedPlan
1126		N
1127		
1128	\oplus	Image : Information about image objects relevant to patient's treatment
1129		 ImageName (String)
1130		 DICOM_UID (String)
1131		 ImageModality (String) e.g. CT, kV, CBCT, MR-T1w, MR-T2w, PET, etc
1132		● SourceSystem (String) 策 Where to find the image and how to get it e.g. ARIA, Velocity,
1133		Hospital PACS, etc
1134		 AccessionNumber (String)
1135		 StudySeries (String)
1136		 BodySite (String)
1137		 DateOfImageAcquisition (Date)
1120		• AgeAtImageAcquisition (Decimal F3) 器
1120		
1138		 RelevanceComment (String?) e.g. TumorResponse
1138 1139 1140		 RelevanceComment (String?) e.g. TumorResponse

1142	⇔ List <imagedatafeature></imagedatafeature>
1143	⇔ TreatedPlan
1144	⇔ Course
1145	
1146	Parent Class Relationships
1147	➡ Patient
1148	
1149	DVHCurve : Store the DVH curve for as treated (i.e. number of fractions delivered) plans and plan
1150	sums. Every Treated Plan has a list of DVH curves
1151	 StructureName (String) – Use TG263 Standardization
1152	 Volume[cc] (Decimal)
1153	 Min[Gy] (Decimal)
1154	• Max[Gy] (Decimal)
1155	 Mean[Gy] (Decimal)
1156	 Median[Gy] (Decimal)
1157	 Stdev[Gy] (Decimal)
1158	● DVHCurve (String) 発 – Dose, Volume tuples separated by semi colons. Dose is in units of Gy,
1159	Volume is in units of percent of structure volume e.g. 0,100; 50,100;50.5,99.5;
1160	
1161	Sibling Class Relationships
1162	⇔ List <dvhmetric></dvhmetric>
1163	
1164	Parent Class Relationships
1165	← TreatedPlan
1166	
1167	DVHMetric : Metrics provide quick look up of most important values. Sibling relationship to DVH
1168	curves is maintained so that they can be reported separately if needed.
1169	 StructureName (String) - Use standard nomenclature from TG263
1170	 MetricName (String) - Use standard nomenclature from TG263

1171	Value
1172	
1173	Sibling Class Relationships
1174	⇔ List <dvhcurve></dvhcurve>
1175	
1176	Parent Class Relationships
1177	← TreatedPlan
1178	
1179	ImageDataFeature : specific values associated with the image that e.g Radiomics values.
1180	Every Image has a list of image data features
1181	 *FeatureName(String) – Need for a standardized list of defined feature names and
1182	acceptable values
1183	 Data Type (String): text, number, datetime, bool
1184	• Value (String) 米
1185	 DateOfImageDataFeature (Date)
1186	● AgeAtImageDataFeature (Decimal F3) 発
1187	
1188	
1189	Parent Class Relationships
1190	← Image
1191	← Patient
1192	PatientTreatmentOutcome
1193	 *DiseaseStatus (String) – Need standardized list e.g. Local Recurrence, NED,
1194	BiochemicalFailure
1195	 DateOfStatus (Date)
1196	• AgeAtStatus (Decimal F3) 発
1197	Class Property Relationship

1198		DiagnosisAndStaging
1199		
1200		Parent Class Relationships
1201		수 Patient
1202		⇔ Course
1203		
1204	\oplus	PatientReportedOutcome
1205		 *SurveyInstrumentName (String) – Need for standardized list
1206		 *ElementName (String) – Need for standardized list
1207		 DateOfPRO (Date)
1208		• AgeAtPRO (Decimal F3) 光
1209		• Value (String)
1210		 ValueType (String) – e.g. Bool, Date, Number
1211		Sibling Class Relationship
1212		⇔ - Course
1213		\geq
1214		Parent Class Relationship
1215		⇐ Patient
1216		0
1217		
1218	\oplus	ProviderReportedToxicity
1219		 *ToxicityName – Use standard names from CTCAE or other standards
1220		 ToxicityStandard (String) e.g. CTCAE
1221		 DateOfReportedToxicty (Date)
1222		• AgeAtReportedToxicity(Decimal F3) 器
1223		• Value (String)
1224		 ValueType (String) – e.g. Bool, Date, Number
1225		• Attribution (String)

1226	Sibling Class Relationship
1227	⇔ Course
1228	
1229	Parent Class Relationship
1230	← Patient
1231	
1232	$\overline{\mathbf{O}}$
1233	HealthInformation: Used to record data elements relevant to patient status e.g. smoker, rock
1234	climber, diabetes, etc.
1235	 *HealthInformationItemName (String) –Need for standardized list e.g. HasDiabetes,
1236	IsCurrentSmoker, SmokingPackYears
1237	Oate (Date)
1238	● AgeDate (Decimal F3) 発
1239	 Value (String) – e.g. True, 20
1240	 ValueType (String) – Decimal, Bool, Date, String
1241	
1242	Sibling Class Relationships
1243	
1245	
1211	
1245	Parent Class Relationships
1246	← Patient
1247	
1248	① Lab
1249	 LabName (String)
1250	 LOINCShortName (String)
1251	 LOINCCodeName (String)
1252	 Date (Date)
1253	● AgeAtDate (Decimal F3) ¥

1254	• Value (String)
1255	 Units (String)
1256	 ValueType (String) – Decimal, Bool, Date, String
1257	
1258	Sibling Class Relationships
1259	⇔ Course
1260	
1261	Parent Class Relationships
1262	← Patient
1263	
1264	① Medication
1265	 MedicationType (String)
1266	 MedicationName (String)
1267	 DosageValue (Decimal)
1268	 DosageUnit (String)
1269	 Frequency (String)
1270	 DateOfMedicationRecord
1271	• AgeAtMedicationRecord (Decimal F3) 発
1272	
1273	Sibling Class Relationships
1274	⇔ Course
1275	
1276	Parent Class Relationships
1277	← Patient
1278	
1279	ChemotherapyCourse: Set of Chemotherapy administrations
1280	 *Protocol (String) – Need standardized list
1281	• Agent (String)

1282		• Facility (String)
1283		 IsNeoAdjuvant (Bool)
1284		 IsConcurrent (Bool)
1285		 IsAdjuvant (Bool)
1286		 DateFirstTreatment (Date)
1287		● AgeAtFirstTreatment (Decimal F3) 器
1288		DateLastTreatment (Date)
1289		● AgeAtLastTreatment (Decimal F3) 策
1290		0
1291		Sibling Class Relationships
1292		Radiation Therapy Course
1293		⇔ Surgical Procedure
1294		
1295		Child Class Relationships
1296		⇒ List <chemotherapy administration=""></chemotherapy>
1297		5
1298		Parent Class Relationships
1299		⇐ Patient
1300		DiagnosisAndStaging
1301		0
1302	\oplus	ChemotherapyAdministration
1303		 Agent (String)
1304		 Dosage (String)
1305		 DateOfAdministration (Date)
1306		O AgeAtAdministration (Decimal F3) ₩
1307		
1308		
1309	\oplus	SurgicalProcedure

1310	• Facility (String)
1311	 *Purpose (String) – Need for standardized list
1312	 *Margins (String) – Need for standardized values
1313	 *BiopsyStatus (String) – Need for standardized values
1314	 Is PreIrradiation (Bool)
1315	 DateOfSurgery (Date)
1316	O AgeAtSurgery (Decimal F3) 米
1317	
1318	Sibling Class Relationships
1319	⇔ Radiation Therapy Course
1320	⇔ ChemoTherapy Course
1321	
1322	Parent Class Relationships
1323	🗢 Patient
1324	DiagnosisAndStaging
1325	
1326	① Pathology
1327	 *ElementName(String) – Need standardized list
1328	 *ElementValue (String)
1329	 *ElementType (String)
1330	 DateOfPathology (Date)
1331	O AgeAtPathology (Decimal F3) 米
1332	
1333	Sibling Class Relationships
1334	⇔ DiagnosisAndStaging
1335	
1336	Parent Class Relationships
1337	⇐ Patient

1338	
1339	
1340	Charge
1341	• CPTCode (String)
1342	NCodeInstances(Int)
1343	 DateStartRange (Date)
1344	• AgeAtStartRange (Decimal F3) 器
1345	 DateEndRange (Date)
1346	● AgeAtEndRange (Decimal F3) 発
1347	S S S S S S S S S S S S S S S S S S S
1348	Parent Class Relationships
1349	← Patient
1350	⇐ Course
1351	ð
1352	Figure Legend
1353	Figure 1: The data from RTOG 0012, RTOG 0247, and RTOG 0822 were converted into Resource
1354	Description Framework (RDF) specifications and were uploaded onto the NRG/IROC/ACR node

1355 of the Varian learning portal. The mapping was performed according to the diagram shown

above. Distributed learning is enabled for contracted institutions. The distributed learning 1356

between this node and another node on the Varian learning portal (MAASTRO Clinic, 1357

Netherlands) was tested successfully. 1358

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