

DSS Juno EHR v23 Real World Test Plan

CHPL # 15.04.04.2925.Juno.23.01.1.230620

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1. Introduction

1.1 Purpose

The purpose of this ONC test plan is to document the overall testing processes for DSS Juno EHR v23 ONC Certification Edition 2015. This test plan describes the test strategy, testing activities and methods to determine DSS Juno EHR v23 meets the "Real World Testing" requirements.

1.2 Test Objective

This ONC Test plan supports:

- Meeting the regulatory test coverage of the 15 Edition ONC requirements per DSS marketed environments.
- Execution of 100% of the test cases for each certified 15 Edition ONC test component for Juno EHR v23.
- o Identification of the functional components and ONC requirements that should be targeted by tests.
- Provision of time estimates of the testing efforts.
- Description of test data and environments per DSS target marketing environments.
- Listing of deliverable elements that are certified within Juno EHR v23 as required by §170.405 Real World Testing and included on the CHPL listing for Juno EHR v23.

1.3 Process and References

The processes and procedures that guide the implementation of this test plan are:

- o 2015 Edition Test Methods, <u>Test Procedures</u> and Conformance Method
- o 2015 Edition Cures Real World <u>Testing Regulations</u>

The references that support the implementation of this test plan are:

• Health IT Standards References and Resource Documents are listed within each criteria test case.

2. Criteria to be Tested from 2015 Edition ONC Certification

2.1 Test Inclusion

Juno EHR v23 test plan includes test scenarios for the adult inpatient setting. All data exchange and communications are secured and follow both HIPAA privacy and compliance rules. ONC technical standards have been carefully reviewed and implemented for testing.

Test cases have been created for the following criteria:

Care Coordination

<u>§170.315(b)(1) - Transitions of Care</u> <u>§170.315(b)(2) - Clinical Information and Reconciliation and Incorporation</u> <u>§170.315(b)(10) – Electronic Health Information Export</u>



Clinical Quality Measures (CQMs)

<u>§ 170.315(c)(1) - Clinical Quality Measures (CQMs) — Record and Export</u> <u>§ 170.315(c)(2) - Clinical Quality Measures (CQMs) — Import and Calculate</u> <u>170.315(c)(3) – Clinical Quality Measures (CQMs) – Report (Cures Update)</u>

Public Health Reporting

§170.315(f)(2) – Transmission to Public Health Agency – syndromic surveillance

Application Programing Interfaces

§170.315(g)(10) – Standardized API for patient and population services

Electronic Exchange

§ 170.315(h)(1) – Direct Project

2.2 Test Methodology

To demonstrate interoperability and conformance compliance during the Real World Testing the user, following written scripts that are based on application workflow, conducts System Testing and Integration Testing.

- Data is sent and/or received properly between systems.
- Interfaces between applications move data correctly and completely. Test both sending and receiving when interfaces are bi-directional.
- Connectivity with external organizations is accurate and complete as authorized (e.g., continuity of care record to referrals, personal health records for patients, disease management to/from health plan).
- System access is appropriate per assigned privileges.
- Data is processed accurately.
- Data is correctly populated in the user interfaces, reports, and clinical documents.
- All system components that share data or depend on other components work together properly.
- The workflows reflect actual new processes and workflows.
- Usage is defined in and follows policies and procedures. Reinforce training as applicable.

3. General Information for All Measures

Product and CHPL ID	Juno EHR v23 - CHPL ID 15.04.04.2925.Juno.23.01.1.230620
Care Setting	Inpatient Adult
Test Environment	Customer PROD environment or TEST environment that mirrors the organization's production environment.
Real-world networks /tools	 Juno CQMsolution[®] - for the generation of measure specific data for review, generation of the QRDA I files for consumption by CMS systems using the QualityNet Secure Portal for Hospital Quality Reporting (HQR) Juno ConnectEHR[®] – for creation of CDAs, Discharge Summaries and Referral Notes, Syndromic Surveillance messages



	 Surescripts Admin Console-for the data exchange for §170.315(b)(1) and §170.315(h)(1) if the testing is being done in a TEST environment; not needed if testing is being done in PRD environment and transmission is being done to/from other providers. Results will be captured in a pdf document through a variety of screenshots and
	extracts.
Standards Update	Standards updated to USCDI (Y/N): Not Applicable

4. Measures used in Overall Approach

4.1 §170.315(b)(1) Transitions of Care and §170.315(h)(1) Direct Project

Use Case	In order to ensure the continuity of care for any given patient, Juno EHR is able to receive transition of care documents from other providers for care prior to the patient's admission for review/reference. Likewise, Juno EHR is able to generate the transition of care documents to be transmitted to other providers in the ambulatory setting post discharge; or to other inpatient care settings upon patient discharge. To "view" or "manage" inbound CDAs that have not yet been associated with a specific patient, the administrative user who would normally handle that workflow requires additional permissions that are part of the HIM Module. Likewise, any given practitioner user with these additional permissions can generate a CDA and transmit it on demand using the external provider's direct email.
	Based on the standards for interoperability, this ensures that care can be provided based on clinical documentation that provides a complete picture and that the results of testing previously completed is available and does not necessarily need to be repeated.
Certification Criteria	§ 170.315 (b)(1) Transition of care—
	(i) Send and receive via edge protocol—
	 (A) Send transition of care/referral summaries through a method that conforms to the standard specified in § 170.202(d) and that leads to such summaries being processed by a service that has implemented the standard specified in § 170.202(a); and
	 (B) Receive transition of care/referral summaries through a method that conforms to the standard specified in § 170.202(d) from a service that has implemented the standard specified in § 170.202(a)(2). (C) XDM processing. Receive and make available the contents of a XDM package formatted in accordance with the standard adopted in § 170.205(p)(1) when
	the technology is also being certified using an SMTP-based edge protocol.
	(ii) Validate and display —
	 (A) Validate C-CDA conformance – system performance. Demonstrate the ability to detect valid and invalid transition of care/referral summaries received and



formatted in accordance with the standards specified in § 170.205(a)(3), (4), and (5) for the Continuity of Care Document, Referral Note, and (inpatient setting only) Discharge Summary document templates. This includes the ability to:
 Parse each of the document types. Detect errors in corresponding "document-templates," "section-templates," and "entry-templates," including invalid vocabulary standards and codes not specified in the standards adopted in § 170.205(a)(3), (4), and (5). Identify valid document-templates and process the data elements required in the corresponding section-templates and entry-templates from the standards adopted in § 170.205(a)(3), (4), and (5). Correctly interpret empty sections and null combinations. Record errors encountered and allow a user through at least one of the following ways to: Be notified of the errors produced. Review the errors produced.
(B) Display. Display in human readable format the data included in transition of care/referral summaries received and formatted according to the standards specified in § 170.205(a)(3), (4), and (5).
(C) <i>Display section views</i> . Allow for the individual display of each section (and the accompanying document header information) that is included in a transition of care/referral summary received and formatted in accordance with the standards adopted in § 170.205(a)(3), (4), and (5) in a manner that enables the user to:
 Directly display only the data within a particular section; Set a preference for the display order of specific sections; and Set the initial quantity of sections to be displayed.
 (iii) Create. Enable a user to create a transition of care/referral summary formatted in accordance with the standard specified in § 170.205(a)(3), (4), and (5) using the Continuity of Care Document, Referral Note, and (inpatient setting only) Discharge Summary document templates that includes, at a minimum:
 (A) (1) The data classes expressed in the standard in § 170.213 and in accordance with § 170.205(a)(4), (a)(5), and paragraphs (b)(1)(iii)(A)(3)(i) through (iii) of this section, or (2) The Common Clinical Data Set in accordance with §170.205(a)(4) and paragraph (b)(1)(iii)(A)(3)(i) through (iv) of this section for the period until
 (3) The following data classes: (i) Assessment and plan of treatment. In accordance with the "Assessment and Plan Section (V2)" of the standard specified in § 170.205(a)(4); or in accordance with the "Assessment Section (V2)"



and "Plan of Treatment Section (V2)" of the standard specified in §
$170\ 205(a)(4)$
(ii) Goals. In accordance with the "Goals Section" of the standard
specified in § 170.205(a)(4).
(iii) <i>Health concerns</i> . In accordance with the "Health Concerns Section"
of the standard specified in § 170.205(a)(4).
(iv) Unique device identifier(s) for a patient's implantable device(s). In
accordance with the "Product Instance" in the "Procedure Activity
Procedure Section" of the standard specified in § 170.205(a)(4).
(B) Encounter diagnoses. Formatted according to at least one of the following
standards:
(1) The standard specified in § $1/0.20/(1)$.
(2) At a minimum, the version of the standard specified in § 170.207(a)(4).
(C) Cognitive status.
(E) Ambulatory setting only. The reason for referral: and referring or
transitioning provider's name and office contact information.
(F) Inpatient setting only. Discharge instructions.
(G) Patient matching data. First name, last name, previous name, middle name
(including middle initial), suffix, date of birth, address, phone number, and
sex. The following constraints apply:
(1) Date of birth constraint.
(i) The year, month and day of birth must be present for a date of birth.
The technology must include a null value when the date of birth is unknown.
(ii) Optional. When the hour, minute, and second are associated with a
date of birth the technology must demonstrate that the correct time
zone offset is included.
(2) Phone number constraint. Represent phone number (home, business,
cell) in accordance with the standards adopted in § 170.207(q)(1). All
phone numbers must be included when multiple phone numbers are
present.
(3) Sex constraint. Represent sex in accordance with the standard adopted in
§ 1/0.207(I)(1).
§ 170.315 (h)(1) Direct Project—
(I) Applicability Statement for Secure Health Transport. Able to send and receive
health information in accordance with the standard specified in §
170.202(a)(2), including formatted only as a "wrapped" message.
(II) Delivery Notification in Direct. Able to send and receive nearth information in accordance with the standard specified in δ 170 202(e)(1)
References:
• §170.202(a)(2) Applicability Statement for Secure Health Transport, Version 1.2,
August 2015 (Direct)
File:Applicability Statement for Secure Health Transport v1.2.pdf - Direct Project



	• §170.202(d) ONC Implementation Guide for Direct Edge Protocols, Version 1.1, June 25, 2014
	Implementation Guide for Direct Edge Protocols, Version 1.1, June 25, 2014 (healthit.gov)
	• § 170.202(e)(1) Implementation Guide for Delivery Notification in Direct, Version 1.0, June 29, 2012
	170.202(e)(1) Interoperability Standards Advisory (ISA) (healthit.gov)
	• § 170.205(a)(4) HL7 Implementation Guide for CDA [®] Release 2: Consolidated CDA Templates for Clinical Notes (US Realm), Draft Standard for Trial Use Release 2.1, August 2015
Justification	Each of the three types of Transition of Care documents has its own specific content requirements. For the inpatient care setting, the C-CDA types to be tested include:
	Continuity of Care
	Referral Note (receive only)
	Discharge Summary
	Through use of multiple scenarios for a given patient, all components of the §170.315(b)(1) Transitions of Care and §170.315(h)(1) Direct Project will be tested i.e.,
	• Receive health information in accordance with the standard specified in §170.202(a)(2) in the form of a transition of care/referral summary for a given patient that was sent in accordance with a method that conforms to the standard specified in § 170.202(d) and process it such that it is viewable within the patient's EHR. (Note: The incorporation of some of the data elements will be tested separately as part of 170.315(b)(2).)
	• Display the transition of care/referral summary received in a manner in accordance with 170.315(b)(1)(ii)(C) which allows the user to view a specific section, set the # of sections to display and rearrange the order of display.
	 Create a transition of care /discharge summary that can be displayed in human readable format using the appropriate template that includes at a minimum: Patient matching data, i.e., First name, last name, previous name, middle name (including middle initial), suffix, date of birth, address, phone number, and sex
	Common Clinical Data Set items (see the Test Data section for details)
	Assessment and plan of treatment sections, either together or separately
	➢ Goals
	 Health Concerns Unique device identifier(s) for a nationt's implantable device(s)
	 Encounter diagnosis
	 Cognitive status
	 Functional status



	Discharge instructions
	• Transmit the transition of care/discharge summary in accordance with the standard specified in §170.202(a)(2) to an 'address' provided for use for the specific patient at a time to be specified. (Note: The address would allow the C-CDA to be transmitted to other providers in the ambulatory setting post discharge or to other inpatient care settings upon patient discharge.)
Test Methodology	 The content of transition of care documents that will be received from other EHRs will vary as the data will be unique to the specific patient. In order to evaluate each part of the criteria, the testing will be split into 4 parts, i.e., creation, receipt, view/display and transmission. Creation of documents is automatic at the time of discharge; however, the testing should include on demand generation of documents. In the event the facility has not received any inbound transition of care documents during the testing period, a reference xml file is provided by the vendor to utilize for the receipt and view/display. Patient demographic must exist to specifically match the patient for the xml files, whether received from elsewhere or provided by the vendor for import and processing. The xml files provided for use, if necessary, is "170.315b2 Turner CCD_reconciliation.xml". Transmission of documents is accomplished using Surescripts; however, the testing should include on demand generation/transmission of documents using a provider's direct email. On demand is done using the Document Builder in the HIM Module.
	Although Juno v23 is appropriate for an enterprise organization with one or more hospitals for inpatient care, the 'address' for the organization utilized for the inbound and outbound messages will be the same. The size of the organization in terms of the number of beds, the number of unique inpatient locations or the number of providers does not impact functionality.
Test Data	Given that the Continuum of Care, Referral Note and Discharge Summary CDAs are generated automatically on discharge from the inpatient setting, data included in each will be based on the content entered for a given patient. The CCD and the Discharge Summary are generated on all patients, but the Referral Note is only generated on patients who have a consult (referral) to an external provider. For testing of creation and transmission of transition of care documents, data entered in the EHR will vary by patient. At a minimum, this data will include the
	following (if entered on a given patient): Common Clinical Data Set items Patient Name Sex Date of Birth Race and Ethnicity Preferred Language Smoking Status Medication Allergies Medications Problems



	* Procedures
	 Immunizations
	 ✤ Vital Signs
	Laboratory Tests and Results
	✤ Care Team Members
	Unique Device Identifiers
	Assessment and plan of treatment sections, either together or separately
	S Goals
	* Health Concerns
	 Encounter diagnosis
	> Cognitive status
	Cognitive status
	 Functional status Discharge instructions
	Discharge instructions
	• For testing the receipt of transition of care documents from other providers and
	view/display, patient demographics must exist to specifically match the patient
	for the xml files, whether received from elsewhere or provided by the vendor for
	import and processing. The xml file provided for use, if necessary, is "170.315b2
	Turner CCD, reconciliation xml"
	See Appendix A for the details for Test Data Entry in the event data from a client
	PROD environment is not available
Expected Outcome(s)	Testing is organized according to the clinical workflow with the criteria being tested
Expected Outcome(s)	grouned according to the specific scenario
	grouped according to the specific scenario.
	Continuity of Care and Discharge Summary CDAs are automatically generated
	upon patient discharge. Referral Note CDAs are only generated if the patient has
	an External Consult.
	• Each of the CDA documents received should be accessible in the HIM module
	and should be viewable. Content will be based on data for the specific patient.
	Transmission of the CDA documents is done using Surescripts.
	Testing is organized according to the clinical workflow with the criteria being tested
	grouned according to the specific measure data. The steps for the testing and the
	Expected Outcome for each step are detailed below. In general, the Interoperability
	and electronic health information exchanged is accomplished successfully without
	and electronic health mornation exchanged is accomplished successfully without
	errors of message failures during the exchange.
	See attached Appendix A for the step by step details and Expected Outcomes.
Measure	Successful creation, display and transmission of the C-CDA for transition of
	care/referral summaries, i.e., Continuity of Care Document, Referral Note and
	Discharge Summary, in the format that conforms to the standard specified in §
	170.202(d) with no errors detected during the validation process.
	The measure includes four parts, i.e.,
	• Measure b1h1a: % C-CDAs received that can be viewed in human readable
	format as detailed in § 170.205(a)(3). (4) and (5) using the using the Continuity



of Care Document, Referral Note, and Discharge Summary document templates that allow the user to view a specific section, set the # of sections to display and rearrange the order of the display
Numerator= # C-CDAs and/or Discharge Summaries with expected results Denominator = # C-CDAs and/or Discharge Summaries received & viewed
 Measure b1h1b: % discharges that include creation of the data for Discharge Summary document type
Numerator= # Discharge Summaries created for discharges in designated period Denominator = # discharges in designated period
 Measure b1h1c: % discharges that include creation of the data for Discharge Summary document type that can be viewed in human readable format as detailed in § 170.205(a)(3), (4), and (5) using the using the Discharge Summary document templates that allow the user to view a specific section, set the # of sections to display and rearrange the order of the display.
Numerator= # Discharge Summaries with appropriate content that is viewable in human readable format Denominator = # discharges reviewed in designated period NOTE: Sample size will vary based on activity but should include a minimum of 25 encounters for a given year; however, sample size may need to be larger if issues are identified.
 Measure b1h1d: % scenarios that include creation and transmission of the data for Continuity of Care Document and Discharge Summary document types through SMTP protocol to an appropriate direct address in accordance with § 170.202(d) based on the date/time specified and that leads to such summaries being processed by a service that has implemented the standard specified in § 170.202(a)
Numerator= # C-CDAs and/or Discharge Summaries transmitted without errors Denominator = # C-CDAs and/or Discharge Summaries created and transmitted Exclusions: Error due to inappropriate direct addresses



4.2 §170.315(b)(2) Clinical Information Reconciliation and Incorporation

Use Case	In order to ensure the continuity of care for any given patient, Juno EHR is able to receive transition of care documents from other providers for care prior to the patient's admission for review/reference. These documents can then be utilized to compare the data for the patient's active medications, allergies and intolerances and problem list with those currently documented in the EHR, reconcile the discordant data, and then incorporate the reconciled list into the EHR.
	To "view" or "manage" inbound CDAs that have not yet been associated with a specific patient, the administrative user who would normally handle that workflow requires additional permissions that are part of the HIM Module.
	The provider will be able to view these external documents associated with a given patient in the Document Viewer by selecting External Records. The display will include all external documents received and associated with the specific patient and can view the specific document and reconcile the information for problems, medications and allergies with the data that is already in Juno.
	Once data is reviewed and reconciled by the provider, a Continuity of Care document can then be generated to include the data that was reconciled and incorporated for subsequent transmission to other providers in the ambulatory setting post discharge or to other inpatient care settings upon patient discharge. Any given practitioner user with the additional HIM permissions can generate a CDA and transmit it on demand using the external provider's direct email; however, this will generally be done by the HIM department.
	Based on the standards for interoperability, this ensures that care can be provided based on clinical documentation that provides a complete picture of the patient and allows the clinical staff to be aware of any discordances.
Certification Criteria	§ 170.315 (b)(2) Clinical information and reconciliation and incorporation—
	 (i) General requirements. Paragraphs (b)(2)(ii) and (iii) of this section must be completed based on the receipt of a transition of care/referral summary formatted in accordance with the standards adopted in § 170.205(a)(3) through (5) using the Continuity of Care Document, Referral Note, and (inpatient setting only) Discharge Summary document templates on and after December 31, 2022. (ii) Correct patient. Upon receipt of a transition of care/referral summary formatted according to the standards adopted § 170.205(a)(3) through (5), technology must be able to demonstrate that the transition of care/referral summary received can be properly matched to the correct patient. (iii) Reconciliation. Enable a user to reconcile the data that represent a patient's
	 active medication list, allergies and intolerance list, and problem list as follows. For each list type: (A) Simultaneously display (i.e., in a single view) the data from at least two sources in a manner that allows a user to view the data and their attributes, which must include, at a minimum, the source and last modification date. (B) Enable a user to create a single reconciled list of each of the following: Medications; Allergies and Intolerances; and problems.
	(C) Enable a user to review and validate the accuracy of a final set of data.



	 (D) Upon a user's confirmation, automatically update the list, and incorporate the following data expressed according to the specified standard(s) on and after December 31, 2022: (1) <i>Medications</i>. At a minimum, the version of the standard specified in § 170.213; (2) <i>Allergies and intolerance</i>. At a minimum, the version of the standard specified in § 170.213; and (3) <i>Problems</i>. At a minimum, the version of the standard specified in § 170.213. (iv) <i>System verification</i>. Based on the data reconciled and incorporated, the technology must be able to create a file formatted according to the standard specified in § 170.205(a)(4) using the Continuity of Care Document template and the standard specified in § 170.205(a)(5) on and after December 31, 2022.
	 References: 170.205(a)(3) HL7 Implementation Guide for CDA® Release 2: IHE Health Story Consolidation, DSTU Release 1.1 (US Realm) Draft Standard for Trial Use July 2012 170.205(a)(3) Interoperability Standards Advisory (ISA) (healthit.gov) § 170.205(a)(4) HL7 Implementation Guide for CDA® Release 2: Consolidated CDA Templates for Clinical Notes (US Realm), Draft Standard for Trial Use, Volume 1 - Introductory Material, Release 2.1, August 2015 170.205(a)(4) Interoperability Standards Advisory (ISA) (healthit.gov) § 170.205(a)(5) HL7® CDA R2 Implementation Guide: C-CDA Templates for Clinical Notes R2.1 Companion Guide, Release 2-US Realm, October 2019 170.205(a)(5) Interoperability Standards Advisory (ISA) (healthit.gov)
Justification	 Once testing for the §170.315(b)(1) Transitions of Care criteria has been completed, the xml documents will be available for importing. Each of the three types of Transition of Care documents, i.e., Continuity of Care, Referral Note and Discharge Summary, has its own specific content requirements; however, all contain medications, allergies and problems which need to be incorporated. All of the components of the §170.315(b)(1) Clinical information and reconciliation will be tested, i.e., Receipt of a transition of care/referral summary for a given patient (<i>NOTE: This was tested separately as part of</i> §170.315(b)(1).); Matching the document to the correct patient in the EHR in accordance with §170.205(a)(3) through (5). Reconciliation of the data for the patient's active medication list, allergies, and problem list, including: Display of the data from both the EHR and the C-CDA document to allow a comparison of each set of data, Creation of a single reconciled list for each set of data
	 Review and validation of the final reconciled list for each set of data Automatically update the list in the EHR based on the final reconciled list for each data set



Test Methodology	The content of transition of care documents received from other EHRs will vary as
rest methodology	the data will be unique to the specific patient. In the event the facility has not
	the data will be unique to the specific patient. In the event the facility has not
	received any inbound transition of care documents during the testing period, a
	reference xml file is provided by the vendor to utilize for the receipt and
	view/display. Patient demographic must exist to specifically match the patient for
	the xml files, whether received from elsewhere or provided by the vendor for import
	and processing. The yml files provided for use, if pecessary is "170,315b? Turper
	CCD recenciliation well
	Although Juno v23 is appropriate for an enterprise organization with one or more
	hospitals for inpatient care, the 'address' for the organization utilized for the
	inbound and outbound messages will be the same.
	The size of the organization in terms of the number of beds, the number of unique
	innationt locations or the number of providers does not impact functionality
TailData	Inpatient locations of the number of providers does not impact functionality.
lest Data	Given that the Continuum of Care, Referral Note and Discharge Summary CDAs could
	be received from other facilities/EHRs, data included in each will vary and be based
	on the data entered for a given patient.
	Data will be entered into the EHR as part of testing for §1/0.315(b)(1) for each of the
	patients to be tested in the reconciliation. The patient demographic data for these
	patients must match the data included in the xml files. At a minimum, this data will
	include the key items from the Common Clinical Data Set (CCDS) that are needed for
	testing this specific criterion, i.e.,
	A Datient Name
	Sex Sex
	◆ Date of Birth
	Race and Ethnicity
	 Preferred Language
	Medication Allergies
	✤ Medications
	* Problems
	In the event no CDAs are available for use in testing, a vendor supplied xml file is
	provided for use if peressary i.e. "170.315h2 Turner CCD reconciliation vml"
	Testing associated with b1 above can also be utilized for b2
	See Appendix B for the details for Test Data Entry.
	NOTE: Additional clinical data for the remaining items in the CCDS is not needed in
	the EUD as this reconciliation process is generally done at the beginning of the
	The Erry as this reconclination process is generally done at the beginning of the
	patient's inpatient admission, preferably before medication orders are entered so
	that the final reconciled list can be utilized in order checking.
Expected Outcomes	Testing is organized according to the clinical workflow, i.e.,
	• receipt and reconciliation of any transition of care/referral summaries. i.e.
	Continuity of Care Document, Referral Note and Discharge Summary received, and
	e creation of a Continuity of Caro C CDA that includes the data reconciled and
	• creation of a continuity of care C-CDA that includes the data reconciled and
	incorporated in accordance with § 1/0.205(a)(5).
	See attached <u>Appendix B</u> for the step by step details and Expected Outcomes.



Measure	Successful receipt and import of the C-CDAs for transition of care/referral summaries, i.e., Continuity of Care Document, Referral Note and Discharge Summary, such that the data for the active medications, allergies/intolerances and problems can be displayed for comparison, reconciliation and incorporation and a Continuity of Care document created after the incorporation contains the reconciled data.
	The measure includes two parts, i.e., Measure b2a
	• % scenarios that include receipt of the Continuity of Care Document, Referral Note, and/or Discharge Summary document templates that allow the user to view the data in a format that allows comparison of each set of data, creation of a reconciled list and updating the list in the EHR based on the final reconciled list for each data set
	Numerator= # data sets with expected results
	Denominator = # data sets reviewed and reconciled
	Data sets=active meds, allergies and intolerances, problem list
	Measure b2b
	 % Continuity of Care documents created after the incorporation that contains the final list of reconciled data for each data set Numerator= # CCD created with expected results for each of the three data sets
	Denominator = # CCDs generated
	Data sets=active meas, allergies and intolerances, problem list

4.3 §170.315(b)(10) Electronic Health Information Export

Use Case	Specific Juno EHR users with required permissions can generate an export of a folder with a series of files with patient specific data that can then be saved to a designated folder accessible to specified users. As part of the folder contents, there is a publicly accessible hyperlink
Certification Criteria	 § 170.315 (b)(10) Electronic Health Information export- 1. Single patient electronic health information export. 1. Enable a user to timely create an export file(s) with all of a single patient's electronic health information that can be stored at the time of certification by the product, of which the Health IT Module is a part. 2. A user must be able to execute this capability at any time the user chooses and without subsequent developer assistance to operate. 3. Limit the ability of users who can create such export file(s) in at least one of these two ways: To a specific set of identified users As system administrator function. 4. The export file(s) created must be electronic and in a computable format.



	5. The publicly accessible hyperlink of the export's format must be
	included with the exported file(s).
	2. Patient population electronic health information export. Create an export of
	all the electronic health information that can be stored at the time of
	certification by the product, of which the Health IT Module is a part.
	1. The export created must be electronic and in a computable format.
	2. The publicly accessible hyperlink of the export's format must be
	included with the exported file(s).
	3 Documentation The export format(s) used to support paragraphs (h)(10)(i)
	and (ii) of this section must be kept up-to-date.
Justification	Once testing for the §170.315(b)(10) has been completed, the exported folder with
	the data files will be available in the specified location. The format of the file will be
	in a computable format
Test Methodology	Must do required configuration in advance-both users and file location. Permissions
	for the FHR (b)(10) Export functionality are assigned to a specific role in
	Configuration The specific role with the required permission is then assigned to the
	users as deemed appropriate. One or more "Single Patient Export Directory
	Locations" can be configured for use. Once configured, choices will be viewable once
	the nations has been selected
	The content of the patient specific data in the exported folder will vary as the
	data/files will depend on the specific data on the specific patient.
Test Data	Data will be entered into Juno EHR as part of ongoing clinical workflow.
	At a minimum, this data will include the following (if entered on a given patient):
	Common Clinical Data Set items
	Patient Name
	✤ Sex
	✤ Date of Birth
	Race and Ethnicity
	Preferred Language
	Smoking Status
	Medication Allergies Medications
	Vital Signs
	• Laboratory resis and results
	✓ Care ream internitiers
	 Unique Device identifiers Accordent and along of the structure with a structure with a structure of the structure of
	 Assessment and plan of treatment sections, either together or separately
	✤ Health Concerns
	 Health Concerns Encounter diagnosis
	 ✤ Health Concerns ➢ Encounter diagnosis ➢ Cognitive status



	Discharge instructions
Expected Outcomes	The export file(s) created is electronic and in a computable format.
	See Appendix C for the details for Test Data Entry.
Measure	Successful generation and export of a folder with a series of files that contain all of
	the data from Juno EHR for a given patient.
	• Measure b10 % reviewed files that were created where valid content was
	confirmed by a visual inspection
	Numerator= # files reviewed with expected data based on comparison with content
	in Juno EHR
	Denominator = # files reviewed in extract folder
	Files reviewed should include (at a minimum): Problems, Meds, Allergies, Diagnoses

4.4 §170.315(c)(1) Clinical Quality Measures (CQMs) – Record and Export and §170.315 and (c)(3) Clinical Quality Measures (CQMs) – Report

Use Case	Juno EHR is appropriate for an organization with one or more hospitals for inpatient care. To ensure the capture of data in real time, configuration of the clinical documentation data is done in a manner that allows the association of codes with the data at the time of the data storage for the various data elements. This includes the demographic data as well as the data entered in the various modules within Juno including Allergies, Diagnoses, Vitals, Orders, Bar Code Medication Administration (BCMA), Clinical Note Templates (CNTs) and Laboratory Tests.
	Data required for CQM exclusions or exceptions must also be codified entries, which may include specific terms as defined by each CQM, or may include codified expressions of "patient reason," "system reason," or "medical reason." In general, the capture of the data is done as part of the usual clinical workflow utilized by physicians and nursing staff.
	Based on the scope of the facility, on an annual basis, the facility will determine which EH Clinical Quality Measures they wish to submit data. CQMsolution [®] is a user-friendly, browser-based application for calculating, displaying, and generating clinical quality measure output. Users can initiate quality measure reports, view patient-level results, analyze underlying data, and filter data on demographic variables. If the facility chooses to have a DHIT license, DHIT offers Data Submission Services (DSV) to submit to The Joint Commission (TJC) and to Quality Net for HQR. The Hospital Quality Reporting (HQR) Secure Portal is the only CMS-approved website for secure communications and health care quality data exchange between quality improvement organizations, hospitals, physician offices, nursing homes, end- stage renal disease networks and facilities, and data vendors.
Certification Criteria	§170.315(c)(1) Clinical quality measures—record and export—



	 (i) <i>Record.</i> For each and every CQM for which the technology is presented for certification, the technology must be able to record all of the data that would be necessary to calculate each CQM. Data required for CQM exclusions or exceptions must be codified entries, which may include specific terms as defined by each CQM, or may include codified expressions of "patient reason," "system reason," or "medical reason." (ii) <i>Export.</i> A user must be able to export a data file at any time the user chooses and without subsequent developer assistance to operate: a. Formatted in accordance with the standard specified in §170.205(h)(2); b. Ranging from one to multiple patients; and c. That includes all of the data captured for each and every CQM to which technology was certified under paragraph (c)(1)(i) of this section.
	§170.315(c)(3) Clinical quality measures—report—
	 Enable a user to electronically create a data file for transmission of clinical quality measurement data: (i) In accordance with the applicable implementation specifications specified by the CMS implementation guides for Quality Reporting Document Architecture (QRDA), category I, for inpatient measures in § 170.205(h)(3) and CMS implementation guide for QRDA, category III for ambulatory measures in § 170.205 (k)(3); or
	(ii) In accordance with the standards specified in § $170.205(h)(2)$ and § $170.205(k)(1)$ and (2) for the period until December 31, 2022.
	References:
	 §170.205(h)(3) CMS Implementation Guide for Quality Reporting Document Architecture: Category I; Hospital Quality Reporting Implementation Guide for 2020 <u>2020 CMS QRDA HQR IG (healthit.gov)</u> §170.205(k)(3) CMS Implementation Guide for Quality Reporting Document
	Architecture: Category III; Eligible Clinicians and Eligible Professional Programs Implementation Guide for 2020
Justification	Electronic Clinical Quality Measures (eCQMs) are specific to either the inpatient care setting or the ambulatory setting; however, the approach to data capture and calculations is the same. Although the functionality supports clinical documentation for ambulatory setting associated with the inpatient facility, eCQM testing will be limited to EH measures for inpatients and will not include any EP measures.
	An overlap exists between the CMS eCQMs and the Joint Commission ORYX Measures and in both cases, the organization selects which specific measures they report data for the subsequent year, given the expectation for quarterly reporting. Reporting is done retrospectively in accordance with established deadlines with a 4



	month delay, allowing sufficient time for the computations of the LOS after discharge to exclude patients with LOS greater than 120 days. For example, Hospital Quality Reporting data for Q2 2023 (Apr 1 – June 30, 2023) must be submitted by Nov 1, 2023.
	Through the use of a single eCQM, i.e., CMS 108v11 Venous Thromboembolism Prophylaxis/ORYX VTE-1, and standard out-of-the box clinical documentation accordance with the clinical workflow for the data capture functionality, all of the components of the §170.315(c)(1) Clinical quality measures (CQMs)-record and export will be tested as well as §170.315(c)(3) Clinical quality measures (CQMs)- report, i.e.,
	• record encounter data that includes the Taxpayer Identification Number (TIN) and National Provider Identifier (NPI), Provider Type and Practice Site Address for the provider as well as the Insurance, Age, Sex, Race and Ethnicity and Problem List for the patient;
	• record codified clinical data as defined for the specific measure utilizing the core functionality of the various modules within Computerized Patient Record System (vxCPRS) including Allergies, Diagnoses, Vitals, Orders, Bar Code Medication Administration (BCMA), Clinical Note Templates (CNTs), Laboratory Tests and Procedures;
	 utilize the DHIT CQMsolution[®] software to generate measure specific data for review; and generate data files and successfully export the QRDA I data file(s) for one or more patients and one or more patients and one or more patients.
	systems using the QualityNet Secure Portal for submissions for Hospital Quality Reporting (HQR);
Test Methodology	Juno EHR is appropriate for an organization with one or more hospitals for inpatient care such that each hospital has their own Taxpayer Identification Number (TIN), as well as their own Medicare and Medicaid numbers. In the event an organization has more than one inpatient hospital, the data included in the report would be split by TIN.
	The size of the organization in terms of the number of beds or the number of unique inpatient locations does not impact functionality.
	DHIT CQMsolution [®] is utilized for (1) calculation of the measures, (2) viewing the data in a Dashboard format or a patient drill-down screen, and (3) generation of the QRDA I files needed for submission to Joint Commission, CMS or other regulatory bodies.
Test Data	Data will be entered into Juno EHR as part of ongoing clinical workflow; however, in order to ensure that the data required for the measures is captured in a manner that it will be included in the abstract for subsequent analysis/compilation, it must be entered using specific defined processes.
	See <u>Appendix D</u> for the details for Test Data and Test Script Prerequisites.



	NOTE: The details included in Appendix D provides an example of data foe eCQM
	CMS 108v11 and usage of the prerequisite configuration for orderables and note
	templates.
Expected Outcomes	Successful compilation of the data for the selected EH eCQMs compiled by DHIT in
	CQMSolution [®] in a format that can be utilized for subsequent transmission to the
	designated site (Joint Commission or CMS).
Measure	The measure includes three parts, i.e.,
	• Measure c1a: % patients who are discharged that are included in the Measure
	108 report
	Numerator= # patients included on the Measure 108 report with data evaluated
	(1) Initial Patient Population
	(2) Numerator
	(3) Denominator
	(4) Denominator Exception (None for this measure) and
	(4) Denominator Exclusion
	Denominator = # natients who are discharged for report period
	• Measure c1b: % reports viewed that include the accurate data calculated for the
	measure and for each of the nationts detailed on the report
	incusate and for each of the patients detailed on the report
	Numerator= # natients with expected results based on data entry for the specific
	natient
	(1) Initial Patient Population
	(2) Numerator
	(2) Denominator
	(3) Denominator,
	(4) Denominator Exclusion
	Denominator – # natients reviewed prior to transmission/unloading
	benominator – # patients reviewed pror to transmission uploading
	Measure c1c: % valid ORDA L files generated using COMSolution® for
	consumption by Joint Commission or CMS systems including for Hospital Quality
	Reporting (HOR)
	Numerator- #valid ORDA L files
	Denominator - # ORDA I files requested for subsequent transmission/unloading
	benominator – # QNDA i jies requested joi subsequent transmission/uploading



4.5 §170.315(c)(2) Clinical quality measures (CQMs) – Import and Calculate

Use Case	Although it is appropriate for organizations to receive and import C-CDA documents and reconcile/integrate specific data for patients (active medications, allergies and intolerances, and problems) to be used in providing appropriate clinical care, no scenario exists where a provider would be receiving a QRDA file and incorporating data into the patient's EHR and then subsequently utilize that data documenting care provided elsewhere for the calculation of the clinical quality measures. Only clinical documentation for care provided at the organization should be included. If clinical data for care or testing provided at other facilities is appropriate for inclusion, it would be manually entered through the usual clinical documentation workflow after review by the appropriate provider and not imported from a QRDA file.
Certification Criteria	§170.315 (c)(2) Clinical quality measures—import and calculate—
	 (i) Import. Enable a user to import a data file in accordance with the standard specified in §170.205(h)(2) for one or multiple patients and use such data to perform the capability specified in paragraph (c)(2)(ii) of this section. A user must be able to execute this capability at any time the user chooses and without subsequent developer assistance to operate. (ii) Calculate each and every clinical quality measure for which it is presented for certification.
	References:
	 <u>Clinical quality measures (CQMs) — import and calculate HealthIT.gov</u> 170.205(h)(2) HL7 CDA Release 2 Implementation Guide for: Quality Reporting Document Architecture-Category 1 (QRDA I); Release 1, DTSU Release 3 (US Realm), Volume 1
Justification	QRDA I files are utilized during certification of Juno v23 contain data for patients not under the care of the organization executing the Real World Testing Plan. No use case exists for this functionality.
Test Methodology	Import QRDA I into Juno CQMsolution which can utilize that data for generation of measure specific data for review, generation of the QRDA I files for: (1) calculation of the measures, (2) viewing the data in a Dashboard format or a patient drill-down screen, and (3) generation of the QRDA I files needed for submission to regulatory bodies.
Test Data	QRDA files obtained from a vendor supplied xml file provided for use during testing, i.e., "170.315c2_QRDA 1Annie_Joseph."
Expected Outcomes	Successful ingestion of the data from the QRDA I file such that it is included in the calculation of the selected measure.
Measure	• Measure c2: % reports viewed that include the accurate data calculated for the measure and for each of the patients imported via the QRDA I file.
	Numerator= # patients with expected results based on import from the QRDA I file for the specific patient (1) Initial Patient Population, (2) Numerator (3) Denominator,



(4) Denominator Exception (None for this measure) and
(5) Denominator Exclusion,
Denominator = # patients imported from QRDA I files

4.6 §170.315(f)(2) Transmission to Public Health Agency – syndromic surveillance

Use Case	 care. In order to provide near "real-time" data to governmental public health agencies (PHAs) in support of public health surveillance functions for all inpatient admissions and discharges, clinical data are provided to PHAs for all patient encounters using the exchange of HL7 messages. Receivers may be state, regional and/or local public health authorities, or a designated third party. To ensure the capture of data in real time, configuration of the clinical documentation data is done in a manner that allows the association of codes with the data at the time of the data storage for the various data elements. This includes the data required for reporting syndromic surveillance information to public health agencies, i.e., Treatment facility information Limited personal identifiable information Demographic information about patients Visit information Diagnostic and pre-diagnostic information Kisk factor and other information Acknowledging message receipt 				
	admissions, A08s for all updates/data changes in the specific data elements and A03s for all discharges. Submission to the specific PHA is then done based on facility specific configuration in Juno ConnectEHR [®] .				
Certification Criteria	§ 170.315 (f)(2) Transmission to public health agencies – syndromic surveillance—				
	Create syndrome-based public health surveillance information for electronic transmission in accordance with the standard (and applicable implementation specifications) specified in § 170.205(d)(4).				
	References:				
	§ 170.205(d)(4) <u>Health Level 7 (HL7[*]) 2.5.1</u> . <i>Implementation specifications</i> . <u>PHIN</u> <u>Messaging Guide for Syndromic Surveillance: Emergency Department, Urgent, Care,</u> <u>Inpatient and Ambulatory Care, and Inpatient Settings, Release 2.0, April 21,</u> <u>2015</u> and <u>Erratum to the CDC PHIN 2.0 Implementation Guide, August 2015</u>				
	HL7 [®] Version 2.5.1 Implementation Guide: Syndromic Surveillance, Release 1 - US Realm Standard for Trial Use, July 2019				
Justification	Each of the three required types of HL7 messages has specific content requirements in terms of segments based on § 170.205(d)(4).				



	 Through use of multiple scenarios for a given patient, all components of the § 170.205(d)(4) are included: Admit Update Discharge
Test Methodology	Must do required configuration of Juno ConnectEHR® to generate the syndromic surveillance report. This is done as part of the implementation process and includes the ability to electronically create syndrome-based public health surveillance information for electronic transmission to PHAs.
Test Data	Given that the A01, A08 and A03 HL7 messages are generated automatically on admission, update and discharge from the inpatient setting, data included in each will be based on the content entered for a given patient.
	At a minimum, this data sent from Juno EHR to Juno ConnectEHR® each night will include the key items that are needed for testing this specific criterion, i.e., Patient Name Sex
	 Date of Birth (used for age calculation) Demographic information about patients (Address, Race & Ethnicity) Visit information for the inpatient admission Diagnostic and pre-diagnostic information
	 Vital measurement information for Height and Weight Risk factor and other information (e.g. Smoking status)
Expected Outcomes	Transmission of A01, A08 and A03 HL7 messages with the required content from JunoEHR on a nightly basis to Juno ConnectEHR [®] and used to create messages for transmission to PHAs based on facility specific configuration. See <u>Appendix F</u> for the details for Test Data Entry.
Measure	HL7 2.5.1 compliant messages based on HL7 messages- Use Smart HL7 Viewer to confirm content and code to look at messages generated in Juno ConnectEHR [®] .
	 Measure f2 % files created in DHIT for transmission to PHAs that contain the expected content based on the information provided by a specific trigger event, i.e., admission to a hospital, updating of demographic information and discharge from an inpatient setting Numerator= # files with the expected content based on the visual inspection/validation of the content when compared to the source data in Juno EHR or the data in the HL7 messages received Denominator = # files created based on incoming A01, A08 and A02, HL7 messages
	and then available for transmission to the appropriate PHA



Use Case	Third party applications allow patients to connect their medical information to an app using an API that allows data access to the patient specific data in Juno EHR.			
Certification Criteria	§ 170.315(g)(10) Standardized API for patient and population services—			
	The following technical outcomes and conditions must be met through the demonstration of application programming interface technology.			
	1. Data response.			
	 Respond to requests for a single patient's data according to the standard adopted in § 170.215(a)(1) and implementation specifications adopted in § 170.215(a) and in § 170.215(b)(1), including the mandatory capabilities described in "US Core Server CapabilityStatement," for each of the data included in the standards adopted in § 170.213. All data elements indicated as "mandatory" and "must support" by the standards and implementation specifications must be supported. 			
	 Respond to requests for multiple patients' data as a group according to the standards and implementation specifications adopted in § 170.215(a), (b)(1), and (d), for each of the data included in the standards adopted in § 170.213. All data elements indicated as "mandatory" and "must support" by the standards and implementation specifications must be supported. 			
	2. Supported search operations.			
	 Respond to search requests for a single patient's data consistent with the search criteria included in the implementation specifications adopted in § 170.215(b)(1), specifically the mandatory capabilities described in "US Core Server CapabilityStatement." 			
	 Respond to search requests for multiple patients' data consistent with the search criteria included in the implementation specification adopted in § 170.215(d). 			
	3. <i>Application registration</i> . Enable an application to register with the Health IT Module's "authorization server."			
	4. Secure connection.			
	 Establish a secure and trusted connection with an application that requests data for patient and user scopes in accordance with the implementation specifications adopted in § 170.215(b)(1) and (c). 			
	 Establish a secure and trusted connection with an application that requests data for system scopes in accordance with the implementation specification adopted in § 170.215(d). 			
	5. Authentication and authorization.			

4.7 §170.315(g)(10) Standardized API for Patient and Population Services



	1. Authentication ar	nd authorization for patient and user scopes.
	1. First time	connections.
	1. A tł a a 1	Authentication and authorization must occur during he process of granting access to patient data in accordance with the implementation specification adopted in § 170.215(c) and standard adopted in § .70.215(e).
	2. A a tł a sj	Health IT Module's authorization server must issue refresh token valid for a period of no less than hree months to applications using the "confidential pp" profile according to an implementation pecification adopted in § 170.215(c).
	3. A a m	Health IT Module's authorization server must issue refresh token for a period of no less than three nonths to native applications capable of securing a efresh token.
	2. Subseque	ent connections.
	1. A a a a	access must be granted to patient data in accordance with the implementation specification adopted in § 170.215(c) without requiring re- authorization and re-authentication when a valid efresh token is supplied by the application.
	2. A a tł a sj	Health IT Module's authorization server must issue refresh token valid for a new period of no less than hree months to applications using the "confidential pp" profile according to an implementation pecification adopted in § 170.215(c).
	2. Authentication ar and authorization application access Backend Services implementation s application must	nd authorization for system scopes. Authentication in must occur during the process of granting an is to patient data in accordance with the "SMART :: Authorization Guide" section of the specification adopted in § 170.215(d) and the be issued a valid access token.
6.	Patient authorization revo must be able to revoke an at a patient's direction wi	<i>ocation.</i> A Health IT Module's authorization server nd must revoke an authorized application's access ithin 1 hour of the request.
7.	<i>Token introspection.</i> A He to receive and validate to implementation specification	ealth IT Module's authorization server must be able okens it has issued in accordance with an otion in § 170.215(c).
8.	Documentation.	
	 The API(s) must in that contains, at a 	nclude complete accompanying documentation a minimum:



	 API syntax, function names, required and optional parameters supported and their data types, return variables and their types/structures, exceptions and exception handling methods and their returns. The software components and configurations that would be necessary for an application to implement in order to be able to successfully interact with the API and process its response(s) 				
	 All applicable technical requirements and attributes necessary for an application to be registered with a Health IT Module's authorization server. 				
	 The documentation used to meet paragraph (g)(10)(viii)(A) of this section must be available via a publicly accessible hyperlink without any preconditions or additional steps. 				
	References:				
	§ 170.215(a)(1) <u>Health Level 7 (HL7[°]) Version 4.0.1 Fast Healthcare Interoperability</u> <u>Resources Specification (FHIR[°]) Release 4, October 30, 2019</u>				
	§ 170.215(b)(1)(i) <u>HL7[°]FHIR[°] US Core Implementation Guide STU V3.1.1</u> (Adoption of this standard expires on January 1, 2026)				
	§ 170.213(a) <u>United States Core Data for Interoperability (USCDI)</u> , Version 1 (Adoption of this standard expires on January 1, 2026)				
	§ 170.215(d)(1) HL7° FHIR° Bulk Data Access (Flat FHIR°) (V1.0.0:STU 1)				
	§ 170.215(c)(1) <u>HL7[°] SMART Application Launch Framework Implementation Guide</u> <u>Release 1.0.0</u> (Adoption of this standard expires on January 1, 2026)				
	§ 170.215(c)(2) <u>HL7[*] SMART App Launch Implementation Guide Release 2.0.0</u> , including mandatory support for the "Capability Sets" of "Patient Access for Standalone Apps" and "Clinician Access for EHR Launch"; all "Capabilities" as defined in "8.1.2 Capabilities," excepting the "permission-online" capability; "Token Introspection" as defined in "7 Token Introspection" (This standard is required by December 31, 2025)				
Justification	Patient specific data in Juno EHR will be available to third party applications via the API if the required configuration has been completed for the external application.				
Test Methodology	Must do required configuration in advance for the external application configuration, e.g. Apple Health [®] or Inferno [®] .				
	The content of the patient specific data that appears in the external application will vary based on both the functionality of the specific application and the specific data on the specific patient.				
Test Data	 No data entry is routinely needed if the necessary configuration exists for utilization of an app. Data entered in the EHR will vary by patient. At a minimum, this data will include the following (if entered on a given patient): ➢ Common Clinical Data Set items ❖ Patient Name 				



	✤ Sex				
	✤ Date of Birth				
	Race and Ethnicity				
	 Preferred Language 				
	 Smoking Status 				
	Medication Allergies				
	Medications				
	✤ Problems				
	 Procedures 				
	Immunizations				
	 Vital Signs 				
	Laboratory Tests and Results				
	✤ Care Team Members				
	Unique Device Identifiers				
	Assessment and plan of treatment sections, either together or separately				
	✤ Goals				
	✤ Health Concerns				
	Encounter diagnosis				
	Cognitive status				
	Functional status				
	Discharge instructions				
	Test data entry would apply ONLY to the event the facility is not utilizing any				
	external applications such as Apple Health® If the Inferno test tool is to be used the				
	data for the 2 patients required is detailed below				
	data for the 2 patients required is detailed below.				
	Jane Annie Clarkson				
	Susan Marie Smith				
Expected Outcomes	Storage of data with the required codes in JunoEHR for retrieval by the standardized				
•	API used by external applications, including but not limited to the Inferno test tool.				
	See <u>Appendix G</u> for the details for Test Data Entry.				
Measure	Measure g10 % patients evaluated that are validated/do not generate any errors				
	Numerator= # patients evaluated using the Inferno test tool/external application that				
	passed validation/did not generate any errors				
	Denominator = # patients evaluated using the Inferno test tool/external application				



5. Schedule of Key Milestones

Key Milestone	Date/Timeframe
Release of documentation for the Real World Testing to be provided to authorized representatives and providers running Juno EHR v23. This includes surveys, specific instructions on what to look for, how to record issues encountered, and Customer Agreements.	December 1, 2024
Begin collection of information as laid out by the plan.	March 1, 2025
Meet with previously identified providers and authorized representatives to ensure that Real World Testing protocols are effective.	March 31, 2025
Follow-up with providers and authorized representatives to understand any issues arising with the data collection.	Quarterly, 2025
Data collection and review.	Quarterly, 2025
End of Real World Testing period/final collection of all data for analysis.	January 2026
Analysis and report creation.	January 15, 2026
Submit Real World Testing report to ACB (per their instructions).	February 1, 2026

6. Attestation

This Real World Test plan is complete with all required elements, including measures that address all certification criteria and care settings. All information in this plan is up to date and fully addresses the Health IT developer's Real World Testing Requirements.

Authorized	Hilary Kloska	Authorized	561-402-9621
Representative Name	Diverter Clinical Content	Representative	
and Title:	Director, Clinical Content	Phone:	
Authorized	YPA YSAA	Date Signed:	10/31/2024
Representative	Milary Moska		
Signature:			



7. Appendices

7.1 Appendix A §170.315(b)(1) Transitions of Care and §170.315(h)(1) Direct Project

7.1.1 Criteria

- §170.315(b)(1) Transitions of Care and
- §170.315(h)(1) Direct Project

7.1.2 Test Data Entry

• Test data entry would apply ONLY to the event the facility has not received any inbound transition of care documents during the testing period and a reference xml file provided by the vendor is to be utilized for the receipt and view/display. Patient demographic must exist to specifically match the patient for the xml files, whether received from elsewhere or provided by the vendor for import and processing. The xml files provided for use, if necessary, is "170.315b2 Turner CCD_reconciliation.xml". It is available at

- <u>https://ett.healthit.gov/ett/#/validators/ccdar3</u> which can be accessed using the following steps:
 - Navigate to <u>https://ett.healthit.gov/ett/#/direct/certdiscovery/dcdt2</u>
 - \circ ~ Select ETT Home from the top navigation bar ~
 - o Select Message Validators
 - Select C-CDA R2.1 from the top navigation bar
 - \circ $\;$ Then navigate to the middle section "To validate your C-CDA with ONC 2015 Edition".
 - Under step 1, select "Receiver"
 - Under step 2, select 170.315_b2_CIRI_Inp from the drop down
 - Under step 3, select 170.315_b2_ciri_r21_sample1_ccd_v11.xml from the drop down
 - \circ $\;$ Then select "Download" and your file will be downloaded.
- No data entry is routinely needed for creation and transmission functionality as Discharge Summary CDAs are generated on all discharges.

7.1.3 Test Script Prerequisites

- (1) The patient demographic data for these patients must match the data included in the xml file. In addition, this data can then be used for the §170.315(b)(2) scenarios for reconciliation in Appendix B if necessary. Manual data entry for Scenario #1 if using the vendor supplied xml for:
 - Susan Turner

NOTE: No data is required for Problems, Allergies or Medications. If entered, it will be utilized as part of the testing for b2.

Patient Name:	First Name: Susan	Middle Name: Jones	Last Name: Turner
			Alias: Susy Turner
Sex:	Female (F)	DOB:	08/01/1970
Race:	White (2106-3)	Ethnicity:	Not Hispanic or Latino (2186-5)
More Granular Race Code:	2108-9 (White European)	Preferred Language	



Home Address:	1011 Amber Drive Beaverton, OR 97006					
Telephone Number:	Mobile: 555-335-1234 Home: 555-33			55-336	5-1544	
Problems	Problem Name	SNOMED CT	Code Healt Statu:		th Concern s	Start Date End Date
	Fever	386661006		Activ	е	6/22/2015
Medication Allergies:	Allergy Substance	Reaction	Severity		Status	Date
	Penicillin G benzathine RxNorm 7980	Hives SNOMED -CT 247472004	Moderat	e	Active	5/01/1980
Medications:	Medication	Dose	Frequ	ency	Route	Start Date End Date
	Tylenol 500 mg RxNorm 209459		PR	N	Oral	6/22/2015
	Ceftriaxone 100 mg/ml RxNorm 309090		BI)	Intravenou	s 6/22/2015

- (2) The tester must be assigned to the appropriate role, with prescriptive authority, and the environment must be configured appropriately to execute the scenario.
- (3) Vendor supplied xml files to be used for the import in b1 to receive and validate and then for the reconciliation in b2
 - 170.315b2 Turner CCD_reconciliation.xml

NOTE: If vendor supplied xml files need to be utilized, an email with the attached file will need to be sent to the direct address of the tester.



7.1.4 Measure Data

Measure b1h1a:

% C-CDAs received that can be viewed in human readable format as detailed in § 170.205(a)(3), (4), and (5) using the using the Continuity of Care Document, Referral Note, and Discharge Summary document templates that allow the user to view a specific section, set the # of sections to display and rearrange the order of the display
 Numerator= # C-CDAs and/or Discharge Summaries with expected results
 Denominator = # C-CDAs and/or Discharge Summaries received & viewed

Report:

- Numerator data is based on the results of the visual inspections/validation that the expected functionality can be confirmed.
- Denominator data is based on the number of documents that were received and viewed before being associated with a specific patient

Testing is focused on the receipt via Direct message and validation of the Continuity of Care C-CDAs and/or Discharge Summary CDAs such that the document can be viewed in human readable format and the user can view a specific section, set the # of sections to display and rearrange the order of the display.

Step	Steps to View, Parse and Import incoming CCD.xml	Expected Outcome
Step 1	Select External Records from the Left Nav in the HIM Module	5 Tabs will appear under the
		External Mail, i.e., Inbox,
		Outbox, Drafts, Sent and
		Deleted with Inbox showing
		as the default.
Step 2	Review the listing of emails received with attachments to determine	Subject, Body of email and
	which ones might have CDAs that need to be associated with a patient.	attachment will be displayed.
	Click on the email to open it.	
Step 3	In the dropdown for the attachment, select View to open the human	The selection in the Left Nav
	readable html and view the contents of the document. User can Expand	will switch to Document
	All of Collapse All to do a quick review of the content of the document if	Builder and the html will
	so desired before associating it with the patient for the provider's	open with the patient header
	review/action.	info in the body of the
		document and the content
		reflected in the Table of
		Contents on the right nav.
Step 4	Click on a specific item in the Table of Contents to navigate to that item	Specific section selected will
	or drag and drop a specific item within the Table of Contents to rearrange	be displayed. Content will be
	the content prior to associating it with the specific patient.	rearranged based on actions
		taken within the TOC.



Step 5	Record the # C-CDAs received/reviewed with the expected content as the	
	numerator and the # C-CDAs received/reviewed as the denominator	

Measure b1h1b:

• % discharges that include creation of the data for Discharge Summary document type Numerator= # Discharge Summaries created for discharges in designated period Denominator = # discharges in designated period

Report: Patients by Discharge Date

Power I	BI Report Server Home $ ightarrow$ ON	NC_Testing $>$	Patients_by_Disc	hargeDate		✓ Search	\$ ⊉
Favorites	Browse						
DC Begin Date:	1/1/2023	LOS I	ess than or equal to:	120			
DC End Date:	8/30/2023	Pat T	уре	Inpatient] ~	
I4 <	1 of 2 ? > >	Ö	€ 100%	✓	ß	$\wp \mid \rightarrow$	
	Pa	tients by	Discharge Dat	e	Last Refresh	Date: Aug 30 2023 11:50AM	
Encounter #	Patient	Birth Date	Admission Date	Discharge Date	LOS	DC Diagnosis	Dx Code
902015215	CQM, Test Oneeleven	07/01/1956	06/21/2023	06/22/2023	1	Essential (primary) hypertension	110
902015228	Mauve, TW OneElevenOneNow	05/20/1973	06/23/2023	06/23/2023	0	Unspecified acute appendicitis	K35.80
902015247	Purple, TW OneElevenOneNow	05/05/1955	06/26/2023	06/26/2023	0	Unspecified acute appendicitis	K35.80
902015249	Mango, TW OneElevenOneNow	05/05/1955	06/26/2023	06/26/2023	0	Unspecified acute appendicitis	K35.80

Step	Steps to verify CDA creation	Expected Outcome
Step 1	Select Document Builder from the Left Nav in the HIM Module	Patient Search window opens
Step 2	Enter the name of the patient to be verified/select the patient.	Display of listing of Historical
	(NOTE: This has to be done one patient at a time.)	Documents
Step 3	Set filter to Discharge Summary	Display of Discharge
		Summary CDAs that were
		created upon patient
		discharge to include the
		Encounter date range and
		the date/time the CDA was
		created.



Step 4	Compare the patients in the report of Patients by Discharge Date to those	Record the # created/verified
	reviewed/verified to validate a Discharge Summary CCD exists for each	as the numerator for the
	discharge.	measure-> should be a 1:1
		match with the # discharges

Measure b1h1c:

 % discharges that include creation of the data for Discharge Summary document type that can be viewed in human readable format as detailed in § 170.205(a)(3), (4), and (5) using the using the Discharge Summary document templates that allow the user to view a specific section, set the # of sections to display and rearrange the order of the display

Numerator= # Discharge Summaries with appropriate content that is viewable in human readable format Denominator = # discharges reviewed in designated period NOTE: Sample size will vary based on activity, but should include a minimum of 25 encounters for a given year; however, sample size may need to be larger if issues are identified.

Report:

• Denominator data can be obtained directly from the report for Measure b1h1b to randomly select specific cases for review.

NOTE: if not all of the documents received in the period specified for the data sample for the quarter are reviewed, the denominator should be adjusted to reflect the # reviewed. If issues are identified during the review, a larger sample size may be needed to increase the confidence level of the results.

• Numerator data is based on the results of the visual inspections/validation that the expected functionality can be confirmed.

Step	Steps to verify CDA creation	Expected Outcome
Step 1	Select Document Builder from the Left Nav in the HIM Module	Patient Search window opens
Step 2	Enter the name of the patient to be verified/select the patient.	Display of listing of Historical
	(NOTE: This has to be done one patient at a time.)	Documents
Step 3	Set filter to Discharge Summary (or Referral Note, etc. as deemed	Display of Discharge
	appropriate for data needed)	Summary CDAs (or Referral
		Notes based on the filter
		applied) that were created
		upon patient discharge to
		include the Encounter date



		range and the date/time the
		CDA was created.
Step 4	Click on the name of the document, i.e., Discharge Summary to open the	Human readable html opens
	human readable html.	and includes a Table of
		Contents that allows user to
		navigate to a specific section.
Step 5	Click on an item in the Table of Contents to view that specific entry.	The view of the html will
		change to display the specific
		element selected.
Step 6	Compare the data in the human readable html to the content in the	Record the # as the
	patient's EHR for that specific encounter.	numerator for the measure->
		should be a 1:1 match

Measure b1h1d:

 % scenarios that include creation and transmission of the data for Continuity of Care Document and Discharge Summary document types through SMTP protocol to an appropriate direct address in accordance with § 170.202(d) based on the date/time specified and that leads to such summaries being processed by a service that has implemented the standard specified in § 170.202(a)

Numerator= # C-CDAs and/or Discharge Summaries transmitted without errors Denominator = # C-CDAs and/or Discharge Summaries created and transmitted Exclusions: Error due to inappropriate direct addresses

Referral Note CDAs and Discharge Summary CDAs are created automatically upon discharge and are viewable within Juno in the HIM Module. The CDAs can then be selected for transmission using Secure Mail and a direct address. Data can be obtained using either of two options depending on available data for the facility.

Option 1: Review of data for Discharge Summaries – to be used in the event no external consults/referrals are generated during the reporting period.

Step	Steps to transmit CDAs created	Expected Outcome
Step 1	Select Document Builder from the Left Nav in the HIM Module	Patient Search window opens
Step 2	Enter the name of the patient to be verified/select the patient.	Display of listing of Historical
	(NOTE: This has to be done one patient at a time.)	Documents
Step 3	Set filter to Discharge Summary	Display of Discharge
		Summary CDAs that were
		created upon patient
		discharge to include the
		Encounter date range and
		the date/time the CDA was
		created.



Step 4 Step 5	Select document to be transmitted. Click on the mail envelope (Compose with attachment) in the top right corner. Once the email is created with the attached CDA file, complete	Html will open for content review. Content will be based on data available for that patient encounter. Email will appear in the Outbox and then in the Sent
	 the required pieces: To: uses direct address from the Surescripts Directory Subject: Should reflect content but no PHI Body: Should include name of patient Checkbox for Protected Health Information Click on Send 	items mailbox.
Step 6	Click on the Sent items mailbox to view the email with the attached CDA.	Emails will be color coded based on the status. See details below.
Step 7	Review the status to determine the # emails that are sent and are either Successful (Green) or Partially Successful (Yellow)	Numerator= Green + Yellow Denominator = # emails

In reviewing the Sent emails, the color of the email is based on the status:

- Grey: Pending
 - \circ $\;$ Hover over recipient name in To will show as Code 000 Description: Pending
- Green: Successful
 - Hover over recipient name in To will show as Code 010 Description: Successfully accepted by the ultimate receiver.
- Yellow: Partially Successful
 - Hover over @ least one recipient name in To will show as Code 010 Description: Successfully accepted by the ultimate receiver.
 - Hover over @ least one recipient name in To will show as Code 601 Description: The address that you sent to does not appear to be a Direct Project address
- Red: Failed to send to all recipients
 - Hover over recipient name in To will show as Code 601 Description: The address that you sent to does not appear to be a Direct Project address



Summers, Allan 4/16/1982 M

Option 2: Report for Electronic Referral Loops (External Consultations)

Within

Power BI Report Server Home > ONC_Testing > g2_Rt7								
Favorites Browse								
Discharge Start Date: 9/1/2023								
Discharge End Date: 9/11/2023								
⊲ < 1	of 1 >	⊳i Č) (6)	100%		~ 🛱		,2 →
RT 7 - Electronic Referral Loops								
	Denominat	tor Critera		Nun	nerator Crit	teria		
Patient Transition of Care or Referr Within Reporting Period		are or Referral rting Period	Summary o Transmitted Du	f Care Record Crea I / Exchange Electr ring Calander Year	ronically	Receipt of Su Record Con Caler	ummary of Care firmed During Idar Year	Results Total

Patients who have a Referral CDA created upon discharge will be included if the discharge date is within the period.
If the Referral Note was sent using the same process detailed in Option 1 for Discharge Summaries, the Numerator 1 wil
be Yes. If the direct email address was valid, the status of the email will be "green" and Numerator 2 will be Yes.

Yes

0/1

0/2

No

Cumulative Total:

Step	Steps to review Referral Note CDAs	Expected Outcome
Step 1	Use the data for the Cumulative Total for the Numerator 1 (Created and Transmitted) and Denominator.	Data should be 100%
Step 2	In the event that the data for Numerator 2 is not the same as Numerator #1, it is likely due to an incorrect direct address and the message status in Sent emails will be red (or possibly yellow if there were multiple email addresses on the email). This can be verified by checking the email address for the Practitioner in Juno configuration against the Surescripts Directory.	Data can be adjusted to exclude any with incorrect email addresses.



7.2 Appendix B §170.315(b)(2) Clinical Information Reconciliation and Incorporation

7.2.1 Criteria

§170.315(b)(2) Clinical information and Reconciliation and Incorporation

7.2.2 Test Data Entry

• Test data entry would apply ONLY to "receipt" in the event the facility has not received any inbound transition of care documents during the testing period and a reference xml file provided by the vendor is to be utilized for the receipt and view/display. Patient demographic must exist to specifically match the patient for the xml files, whether received from elsewhere or provided by the vendor for import and processing. The xml files provided for use, if necessary, is "170.315b2 Turner CCD_reconciliation.xml". It is available at

https://ett.healthit.gov/ett/#/validators/ccdar3 which can be accessed using the following steps:

- Navigate to https://ett.healthit.gov/ett/#/direct/certdiscovery/dcdt2
- Select ETT Home from the top navigation bar
- Select Message Validators
- o Select C-CDA R2.1 from the top navigation bar
- \circ Then navigate to the middle section "To validate your C-CDA with ONC 2015 Edition".
- Under step 1, select "Receiver"
- Under step 2, select 170.315_b2_CIRI_Inp from the drop down
- Under step 3, select 170.315_b2_ciri_r21_sample1_ccd_v11.xml from the drop down
- Then select "Download" and your file will be downloaded.

7.2.3 Test Script Prerequisites

NOTE: Information in this section is the same as 7.1.3 above for the b1 criteria.

- (1) The patient demographic data for these patients must match the data included in the xml file. In addition, this data can then be used for the §170.315(b)(2) scenarios for reconciliation in Appendix B if necessary. Manual data entry for "receipt" scenario if using the vendor supplied xml for:
 - Susan Turner

Telephone	Beaverton, OR 97006 Mobile: 555-335-1234		Home: <mark>55</mark>	5-336-1544		
Home Address:	1011 Amber Drive				•	
More Granular Race Code:	2108-9 (White Europe	an)	Preferred	Language		
Race:	White (2106-3)		Ethnicity: No La		Not F Latino	lispanic or o (2186-5)
Sex:	Female (F)		DOB:		Allas: 08/01	L/1970
Patient Name:	First Name: Susan		Middle Name: Jones		Last Name: Turner	



	Fever	386661006	Acti	ve	6/22/2015
Medication Allergies:	Allergy Substance	Reaction	Severity	Status	Date
	Penicillin G benzathine RxNorm 7980	Hives SNOMED -CT 247472004	Moderate	Active	5/01/1980
Medications:	Medication	Dose	Dose Frequency		Start Date End Date
	Tylenol 500 mg RxNorm 209459		PRN	Oral	6/22/2015
	Ceftriaxone 100 mg/ml RxNorm 309090		BID	Intravenous	6/22/2015

- (2) The tester must be assigned to the appropriate role, with prescriptive authority, and the environment must be configured appropriately to execute the scenario.
- (3) Vendor supplied xml files to be used for the import in b1 to receive and validate and then for the reconciliation in b2
 - 170.315b2 Turner CCD_reconciliation.xml

NOTES:

- In the event there is no data in the report detailed below in Section 7.2.4, and no CDAs have been received for the testing period, the vendor supplied document can be utilized, i.e., the Continuity of Care C-CDA (Susan Turner). In this case, the CDA would be available for reconciliation based on the steps for 7.1.4 for Measure b1h1a above.
- If CDA documents have been received and associated with patients, but not yet reconciled, the steps shown below can be utilized.

Step	Steps to associate CDA with patient for reconciliation (if vendor document being utilized)	Expected Outcome
Step 5	Click on the Import to EHR icon in the right nav above the Table of	Listing of Potential Patient
	Contents.	Matches will open with the
		Confidence Level Score.
Step 6	Select the patient with who the document should be associated for	The document will be
	review by the provider in the Reconciliation.	associated with the patient
		and will be displayed in
		ProDash in the left Nav under
		External Documents for the
		specific patient.
Step	Steps to verify CDA creation after reconciliation	Expected Outcome



Step 1	In ProDash, enter the name of the patient to be verified/select the	Display of patient's record for
	patient.	the selected encounter.
	(NOTE: This has to be done one patient at a time and may require	
	(NOTE. This has to be done one patient at a time and may require	
	selection of the specific encounter if the patient has more than one	
	encounter. The Admission Date is included in the report generated.)	
Step 2	On the Left Nav, select Document Viewer and External Records.	Display of listing of External Documents that includes:
		Document Source
		Document Date
		Date Received
		• Type
		 Red Button "Needs
		reconciliation"
Step 3	Single click to select the document for review based on the report that	Document will open in a
	includes the Admission Date. Clicking on the individual items in the Table	format that the user can view
	of Contents will take the user directly to that section in the document.	
Step 4	After a quick review of the document received, click on the red button	Side by side comparison of
	"Needs reconciliation"	the External Record (yellow)
		and the data from JUNO
		(green)are displayed for
		action.
Sep 5	For each of the items, i.e. Allergies, Medications and Problems,	The listing of reconciled data
	determine what action should be taken, i.e. Add, Consolidate, No Action,	will be displayed for
	based on the data. Click Next	review/capture of a
		screenshot for reference
		when new C-CDA is created
		in Scenario 2.
Step 6	Click Import	Display of Toaster Message
		that CDA Reconciliation was
		successful.
Step 7	Repeat process for additional patients	



7.2.4 Measure Data

• **Measure b2a**: % scenarios that include receipt of the Continuity of Care Document, Referral Note, and/or Discharge Summary document templates that allow the user to view the data in a format that allows comparison of each set of data, creation of a reconciled list and updating the list in the EHR based on the final reconciled list for each data set

Numerator= # data sets with expected results Denominator = # data sets reviewed and reconciled Data sets=active meds, allergies and intolerances, problem list

Report:

- Denominator data can be obtained from RT15 report that provides a listing of patients with # CDAs received/associated with patients as well as the # CDAs with Meds, Allergies and Problems reconciled
- Numerator data is based on the results of the visual inspections/validation that the expected functionality can be confirmed.

The RT15 Receive and Reconcile Report can be generated on demand. It is based on activity that is part of normal clinical flow that includes receipt of the CDA from an external source via direct messaging and review/import/association of that document with the appropriate patient in the HIM module and subsequent review/reconciliation of the document by the provider. Based on that report, the following steps can be utilized to compare the data to review the activity for a given patient.



1	Power BI Report Server Home > ONC_Testing > g2_Rt15									
Fa	Favorites Browse									
Choose Report Begin Date: 8/29/2023										
Cho	ose Report End [Date: 8/29/2023								
ŀ	⊲ <	1 of 1 >	⊳i Ö	© 100% V	₽ [→]					
RT 15 - Receive & Reconcile										
		Denomin	ator Critera		Numerator Criteria	I	Results			
	Patient	Patient Admitted Within Reporting Period	Number of Electronic Summary of Care Records Received	Number of Electronic Summary of Care Records Where Medication Reconcillation Occurred During the Perfomance Period	Number of Electronic Summary of Care Records Where Medication Allergy Reconcilation Occurred During the Perfomance Period	Number of Electronic Summary of Care Records Where Current Problem List Reconsiliation Occured During the Performance Period	Results Total			
	Daniels, Tracy 5/2/2000 Sex: F	Admitted Community Health Hospitals	4	3	3	3	3/4			
	Potter, Dianne 4/13/1963 Sex: F	Admitted Community Health Hospitals	2	1	2	2	1/2			
	Owens, Katrina 10/18/1994 Sex: F	Admitted Community Health Hospitals	5	2	2	2	2/5			
	Summers, Allan 4/16/1982 Sex: M	Admitted Community Health 1 Hospitals		1	1	1	1/1			
	Mack, Ada 5/18/1968 Sex: F	Admitted Community Health Hospitals	2	0	0	0	0/2			
			•			Cumulative Total:	7/14			

• Measure b2b: % Continuity of Care documents created after the incorporation that contains the final list of reconciled data for each data set

Numerator= # CCD created with expected results for each of the three data sets Denominator = # CCDs generated Data sets=active meds, allergies and intolerances, problem list

Report:

- Denominator data is based on the number of CCDs generated as part of testing since this is not necessarily a process implemented in actual clinical workflow
- Numerator data is based on the results of the visual inspections/validation that the expected functionality can be confirmed.

The steps detailed below delineate the process to create a Continuity of Care C-CDA that includes the data reconciled and incorporated in accordance with § 170.205(a)(5). NOTE: If the same patient(s) reviewed for Measure b2a are utilized, the denominator would be the same; however, the data in the report should reflect the specifics of the CDAs actually reconciled. Any of the patients in the report used for Measure b2a can be utilized to generate the C-CDAs



based on the reconciled data. If CDAs are generated for only a portion of the patients in the report, that # would be used as the denominator.

Step	Steps to verify CDA creation after reconciliation	Expected Outcome
Step 1	Select Document Builder from the Left Nav in the HIM Module	Patient Search window opens
Step 2	Enter the name of the patient to be verified/select the patient. (NOTE: This has to be done one patient at a time.)	Display of listing of Historical Documents
Step 3	Click on the green Generate Document button Select the appropriate encounter Select Document Type, i.e., CCD-Transition of Care	New document is added to the listing.
Step 4	Click on the name of the document that was created, i.e., Transition of Care to open the human readable html.	Human readable html opens and includes a Table of Contents that allows user to navigate to a specific section.
Step 5	Click on an item in the Table of Contents to view that specific entry.	The view of the html will change to display the specific element selected.
Step 6 Step 7	Compare the data in the human readable html to the content in the patient's EHR for that specific encounter for the items reconciled, i.e., Problems Allergies Medications for a given patient. Repeat process for additional patients	Record the # C-CDAs reviewed with the expected content as the numerator and the # C-CDAs reviewed as the denominator



7.3 Appendix C §170.315(b)(10) Electronic Health Information Report

7.3.1 Criteria

• §170.315(b)(10) – Electronic Health Information Export

7.3.2 Test Data Entry

Actual patient data should be utilized for the testing. No specific data is required; however, clinical documentation should include, at a minimum, allergies, medications, problems, vital signs, smoking status and diagnoses.

7.3.3 Test Script Prerequisites

- Configuration of permissions based on role
- Configuration of users with specific role
- Configuration of the Single Patient Export Directory Location such that it is accessible by the user who will be exporting the data.

7.3.4 Measure Data

Successful generation and export of a folder with a series of files that contain all the data from Juno EHR for a given patient.

• **Measure b10** % reviewed files that were created where valid content was confirmed by a visual inspection Numerator= # files reviewed with expected data based on comparison with content in Juno EHR Denominator = # files reviewed in extract folder Files reviewed should include (at a minimum): Problems, Meds, Allergies, Diagnoses

Testing is focused on the files created on the designated directory location.

Step	Steps to Access/View files	Expected Outcome
Step 1	Navigate to the designated directory location.	Location will include a D: API
		Data location
Step 2	Change the formatting of the display to include the listing of all of the	Alphabetical display of all of
	files in the folder to make it easier to find/access the desired data file.	the files available for the
		specific patient based on the
		data in JunoEHR at the time
		the export was created.
Step 3	Determine which data will be reviewed. At a minimum, review the	Majority of patients should
	following: Problems, Meds, Allergies, Smoking status, Diagnoses	have data for these, though
		Allergies may just be No
		known allergies
Step 4	Open the specific file noted above in Notepad++ to view the data	Data should match



Comp	are the data shown in the file to that in Juno EHR for the specific
patien	t encounter. Examples:
Proble	ems: Use files
•	Conditions.csv
•	ConditionCoding.csv
•	ConditionCategory.csv
Medic	ations: Use files
•	Order.csv
•	Orderable.csv
Allergi	es: Use files
•	Allergy.csv
•	AllergyCoding.csv
Diagno	oses: Use files
•	Conditions.csv
•	ConditionCoding.csv
•	ConditionCategory.csv

7.4 Appendix D §170.315(c)(1) Clinical Quality Measures (CQMs) – Record and Export and §170.315(c)(3) Clinical Quality Measures (CQMs) – Report

7.4.1 Criteria

- §170.315(c)(1) Clinical quality measures (CQMs) record and export
- §170.315(c)(3) Clinical quality measures (CQMs) report

7.4.2 Test Data Entry

Actual patient data should be utilized for the testing, but it needs to be entered as detailed below so that the data is captured in a manner for each of the eCQM specific data elements such that it can then be saved to the SQL tables utilized by the DHIT CQMsolution[®] for (1) calculation of the measures, (2) viewing the data in a Dashboard format or a patient drill-down screen, and (3) generation of the QRDA I files needed for submission to Joint Commission, CMS or other regulatory bodies.

The following QDM Data Elements will be captured as part of the patient registration and admitting process based on standard configuration validated during implementation:

- ✓ Encounter, Performed: Non-elective Inpatient Encounter
- ✓ Encounter, Principal Diagnosis
- ✓ Patient Characteristic: Ethnicity
- ✓ Patient Characteristic: Payer
- ✓ Patient Characteristic: Race
- ✓ Patient Characteristic: ONC Administrative Sex
- ✓ Patient Characteristic: Birthdate
- ✓ Discharge Disposition



In addition to the common QDM Data Elements listed above, data will also be captured for the following QDM Data Elements for eCQM CMS 108v11 Venous Thromboembolism as part of the clinical documentation process:

- ✓ Assessment, Performed (used for VTE Risk Assessment for 108v11)
- ✓ Device, Not Ordered (used for VTE Prophylaxis Devices for 108v11)
- ✓ Intervention, Order (used for Comfort Measures for 108v11)
- ✓ Intervention, Performed (used for Comfort Measures for 108v11)
- ✓ Laboratory Test, Performed (used for INR for 108v11)
- ✓ Medication, Administered (used for Antithrombotics for 108v11)
- ✓ Medication, Not Administered (used for Antithrombotics ordered but not administered for 108v11)
- ✓ Medication, Ordered (used for Antithrombotics for 108v11)
- ✓ Medication, Not Ordered (used for Antithrombotics not ordered for 108v11)
- ✓ Procedure, Not Performed (used for VTE Prophylaxis Device not applied for 108v11)
- ✓ Procedure, Performed (used for Application of VTE Prophylaxis Device & Surgical Procedure for 108v11)

7.4.3 Test Script Prerequisites

- (1) Configuration of DHIT CQMsolution[®] to generate the report. This is done by the facility's CQM Solution administrator and includes the selection of the specific EH Measures that the facility wishes to include in the evaluation/data submission to CMS and/or Joint Commission.
- (2) Usage of specific orders and templates with configuration required for the capture of coded data utilized for a given eCQM, i.e., eCQM CMS 108v11.

Orderable items with specific associated codes defined in eCQM Value Sets:

- VTE Device (GCS, IPC, FP)
- Comfort Care
 - Intervention, Order (used for Comfort Measures for 108v11)
- Antithrombotic medications
 - Medication, Ordered (used for Antithrombotics for 108v11)
 - Medication, Administered (used for Antithrombotics for 108v11)
 - Medication, Not Administered (used for Antithrombotics ordered but not administered for 108v11)
- Laboratory Tests-INR
 - Laboratory Test, Performed (used for INR for 108v11)

Templates with items with specific associated codes defined in eCQM Value Sets:

- VTE/PE Assessment (T264)
 - Assessment, Performed (used for VTE Risk Assessment for 108v11)
- History & Physical (T104)
 - Assessment, Performed (used for VTE Risk Assessment for 108v11)
- Nursing Shift Assessment (T242)
 - Intervention, Performed (used for Comfort Measures for 108v11)
 - Procedure, Not Performed (used for VTE Prophylaxis Device not applied for 108v11)
 - Procedure, Performed (used for Application of VTE Prophylaxis Device & Surgical Procedure for 108v11)
- Discharge Summary-Physician (T081)



- Device, Not Ordered (used for VTE Prophylaxis Devices for 108v11)
- Medication, Not Ordered (used for Antithrombotics not ordered for 108v11)
- (3) Data capture of the data for eCQM CMS 108v11 as part of routine clinical workflow is available in the staging table utilized by DHIT CQMsolution[®] to generate the report that contains the measure specific data and the QRDA I data files. Based on the patient specific data, the report will evaluate the patient to determine whether the patient is in the Initial Population, Denominator, Numerator, Exclusion or Exception. NOTE: Not all EH eCQMs have Exceptions.

Me	Measure 108												
Patient Name	Age/ DOB	Device Ordered	Device Applied	Med Order	Med Admin	Comfort Care	Proc- SCIP	DX: OB or VTE	IPP	Den	Excl	Num	Ехср
Example 1	>18	~	~	~	~				~	~		~	
Example 2	>18	~	✓ pt refused	~	~				~	~		~	
Example 3	>18	~	✓	✓	✓ Not admin- med reason				~	✓		~	
Example 4	>18					~			~	~	~		
Example 5	>18						~		~	~	~		
Example 6	>18							~					

7.4.4 Measure Data

The measure includes three parts, i.e.,

• Measure c1a: % patients who are discharged that are included in the Measure 108 report

Numerator= # patients included on the Measure 108 report with data evaluated

- (1) Initial Patient Population,
- (2) Numerator
- (3) Denominator,
- (4) Denominator Exception (None for this measure) and
- (5) Denominator Exclusion,

Denominator = # patients who are discharged for report period



Report: Patients by Discharge Date

Power	BI Report Server Home > ON	IC_Testing $>$	Patients_by_Disc	hargeDate			\$\$ ↓		
Favorites	Favorites Browse								
DC Begin Date:	1/1/2023	LOS I	ess than or equal to:	120					
DC End Date:	8/30/2023	Pat Ty	ype	Inpatient] ~			
⊲ <	1 of 2 ? > >	Ŭ	€ 100%	✓	ß	\wp \rightarrow			
	Ра	tients by	Discharge Dat	e	Last Refresh	Date: Aug 30 2023 11:50AM			
Encounter #	Patient	Birth Date	Admission Date	Discharge Date	LOS	DC Diagnosis	Dx Code		
902015215	CQM, Test Oneeleven	07/01/1956	06/21/2023	06/22/2023	1	Essential (primary) hypertension	110		
902015228	Mauve, TW OneElevenOneNow	05/20/1973	06/23/2023	06/23/2023	0	Unspecified acute appendicitis	K35.80		
902015247	Purple, TW OneElevenOneNow	05/05/1955	06/26/2023	06/26/2023	0	Unspecified acute appendicitis	K35.80		
902015249	Mango, TW OneElevenOneNow	05/05/1955	06/26/2023	06/26/2023	0	Unspecified acute appendicitis	K35.80		

Report: CQMsolution[®] Measure 108

Step	Steps to verify eCQM data in CQMsolution®	Expected Outcome
Step 1	In the HIM Module, select the link for CQMsolution [®] at the top.	User will see the CQM
		Solution log in screen.
Step 2	Enter the User credentials for CQMsolution [®] .	Listing of Queued Reports
		that were previously
		configured by the facility's
		CQM Solution Administrator
Step 3	Select the report that includes EH Measure 108v11 Venous	Report is displayed and you
	Thromboembolism Prophylaxis.	can view aggregated data in the report as well as line by line listing of the patients discharged in the period specified
Step 4	On the Queued Reports Page, Click the View Results Button, then Click	
	the View Detail Link	
Step 5	Count the # patients displayed for Measure 108	Numerator = # pts on report
		Denominator = # patients on
		"Patients by Discharge Date"
		tor same period



• Measure c1b: % reports viewed that include the accurate data calculated for the measure and for each of the patients on the report

Numerator= # patients with expected results based on data entry for the specific patient

(1) Initial Patient Population,

(2) Numerator

(3) Denominator,

(4) Denominator Exception (None for this measure) and

(5) Denominator Exclusion,

Denominator = # patients reviewed prior to transmission/uploading

NOTE: See Venous Thromboembolism Prophylaxis 11.1.000 (healthit.gov) for details on expected results

Report: CQMsolution® Measure 108

Step	Steps to verify eCQM data in CQMsolution®	Expected Outcome
Steps 1-5	See above for Measure c1b	
Step 6	 On the Queued Reports Page, Click the View Results Button, then Click the View Detail Link IPP: Pts >18 with LOS <120 days Den: Same as IPP Excl: LOS <2 days Admits/Transfers to ICU Principal Dx = MH disorder or stroke Principal procedure =SCIP VTE Comfort measures Num: VTE prophylaxis (device & med) Docum of negation rationale Excp: None for this measure 	Report is displayed and you can view aggregated data in the report as well as line by line listing of the patients discharged in the period specified for each of the "bubbles". See the measure specifications for full details/specifications.
Step 7	On a patient row, click on the patient name to display the full details of data captured for that patient to determine whether data was captured and analyzed as expected.	

• Measure c1c: % valid QRDA I files generated using CQMSolution[®] for consumption by Joint Commission or CMS systems, including for Hospital Quality Reporting (HQR)

Numerator= # valid QRDA I files generated

Denominator = # QRDA I files requested for subsequent transmission/uploading



Report: Measure 108

Step	Steps to validate the QRDA 1 files	Expected Outcome
Step 1	Using the Reports generated in for	QRDA I zip file is downloaded
	Measure 108	
	Click the View Results Button, then Click the Detail Tab.	
	Click on the Downloads Button, then Select "Download QRDA I"	
Step 2	Locate QRDA I zip file in designated storage location	File can be uploaded based on facility
		process and agreement with DHIT as
		per license

7.5 Appendix E §170.315(c)(2) Import and Calculate

7.5.1 Criteria

• §170.315(c)(2) Clinical quality measures (CQMs) – import and calculate

7.5.3 Test Data Entry

QRDA files obtained from Cypress Test Tool.

7.5.3 Test Script Prerequisites

Configuration of DHIT CQMsolution[®] to generate the report.

7.5.4 Measure Data

The measure includes three parts, i.e.,

Measure c2: % reports viewed that include the accurate data calculated for the measure and for each of the patients imported via the QRDA I file.

Numerator= # patients with expected results based on import from the QRDA I file for the specific patient

- (1) Initial Patient Population,
- (2) Numerator
- (3) Denominator,
- (4) Denominator Exception (None for this measure) and
- (5) Denominator Exclusion,

Denominator = # patients imported from QRDA I files

See 7.4.4 Measure Data for steps to generate the report.



7.6 Appendix F §170.315(f)(2) Transmission to Public Health Agency - syndromic surveillance

7.6.1 Criteria

• §170.315(f)(2) – Transmission to Public Health Agency – syndromic surveillance

7.6.2 Test Data Entry

Actual patient data should be utilized for the testing. No specific data is required; however, clinical documentation should include, at a minimum:

- Home Address -in PID-11
- Age-calculated based on Date of Birth (LOINC Code 21612-7) in PID-7
- Race (LOINC code as appropriate) in PID-10
- Ethnicity Height (LOINC Code as appropriate) in PID-22
- Height (in) (LOINC Code 8302-2)
- Weight (lbs) (LOINC Code 29463-7)
- Smoking
- Diagnosis(es) -Coded as either W (Working) of F (Final)

7.6.3 Test Script Prerequisites

Configuration of Juno ConnectEHR[®] to generate the syndromic surveillance report. This is done as part of the implementation process and includes the ability to view/access data files that are created based on incoming HL7 messages and are then used to transmit the messages to the appropriate PHA as configured in the database.

7.6.4 Measure Data

• **Measure f2** % files created in DHIT for transmission to PHAs that contain the expected content based on the information provided by a specific trigger event, i.e., admission to a hospital, updating of demographic information and discharge from an inpatient setting

Numerator= # files with the expected content based on the visual inspection/validation of the content when compared to the source data in Juno EHR <u>or</u> the data in the HL7 messages received

Denominator = # files created based on incoming A01, A08 and A03 HL7 messages and then available for transmission to the appropriate PHA

Testing is focused on theHL7 files on the ConnectEHR server such that the document can be viewed in human readable format and used for comparison of the data using the Smart HL7 Viewer.

Step	Steps to Access/View files	Expected Outcome
Step 1	Navigate to the Remote Desktop for the D drive (DATA) which allows access to the ConnectEHR database.	File folder for Program Data



Step 2	Navigate to the ConnecEHR DATA folder. Select/open the DATA folder	Series of folders that includes
Stop 2	Onen the Dublic Health folder to see the display of messages that have	One or more in a series with
step s	been generated	11:50 time stamps as the
	been generated.	11.59 time stamps as the
		lupoEHB opeo por day
Stop 1	Salact (apon the first in a given series with Netanad (Netanad L. This	The contents of the HL7
Step 4	select/open the first in a given series with Notepad/Notepad++ . This	massage will be displayed
	ancounter	and can be conied and placed
		in the Smart HIZ viewer
Stop 5	View the HIT message in Smart HIT viewer	Locate the Encounter # found
Step 5	view the HL7 message in smart HL7 viewer.	in $PV/1_1Q$
Stop 6	Open June EUR Registration module and enter the Encounter # in the	Access to the patient data
stepo	Detion to source to identify the nation	within JunoEHP for side by
	Patient Search to identify the patient.	side comparison of the data
Stop 7	Compare the data shown in the Smart UI 7 viewer to that in June EUP	Data should match
Step 7	for the energific nations ancounter. Data to be reviewed for A01 and A09	
	includes:	
	includes.	
	• PID-5 Name or "S"	
	• PID-8 Sex	
	• PID-10 Race	
	PID-11 Address	
	PID-22 Ethnicity	
	PV1-2 Patient Class	
	• PV1-19 Visit Number (Encounter Number)	
	• PV1-44 Admit Date/Time	
	• PV2-3 Admit Reason (Optional)	
	OBX3-3 Observation Identifier (Age at Time Patient Reported)-code	
	=LOINC 21612-7	
	OBX3-5 Observation Value (Age)	
	• OBX4-3 Observation Identifier (Height)-code =LOINC 8302-2	
	OBX4-5 Observation Value (Height)	
	OBX4-6 Observation Units (Inches)	
	• OBX5-3 Observation Identifier (Weight)-code =LOINC 3141-9	
	OBX4-5 Observation Value (Weight)	
	OBX4-6 Observation Units (lbs)	
	 DG1-3 Diagnosis Code (Code & Cide Description) 	
	• DG1-6 Diagnosis Type (A for Admitting, W for Working or F for Final)	



Step 8	Compare the data shown in the Smart HL7 viewer to that in Juno EHR	Data should match
	for the specific patient encounter. Data to be reviewed for A03	
	includes:	
	• PID-5 Name or "S"	
	PID-8 Sex	
	PID-10 Bace	
	PID-11 Address	
	PID-22 Ethnicity	
	PV1-2 Patient Class	
	 PV1-19 Visit Number (Encounter Number) 	
	PV1-44 Admit Date/Time	
	PV1-45 Discharge Date/Time	
	PV2-3 Admit Reason (Optional)	
	DG1-3 Diagnosis Code (Code & Cide Description)	
	DG1-6 Diagnosis Type (F for Final)	
	OBX3-3 Observation Identifier (Age at Time Patient Reported)-	
	code =LOINC 21612-7	
	OBX3-5 Observation Value (Age)	
	 OBX4-3 Observation Identifier (Height)-code =LOINC 8302-2 	
	OBX4-5 Observation Value (Height)	
	OBX4-6 Observation Units (Inches)	
	• OBX5-3 Observation Identifier (Weight)-code =LOINC 3141-9	
	OBX4-5 Observation Value (Weight)	
	OBX4-6 Observation Units (lbs)	
	 OBX6-3 Observation Identifier (Smoking Status)-code =LOINC 	
	72166-2	
	OBX6-5 Observation Value (Smoking Status)	

7.7 Appendix G §170.315(g)(10) Standardized API for Patient and Population Services

7.7.1 Criteria

• §170.315(g)(10) Standardized API for Patient and Population Services

7.7.2 Test Data Entry

- Test data entry would apply ONLY to the event the facility is not utilizing any external applications such as Apple
- Health[®]. If the Inferno test tool is to be used, the data for the 2 patients required is detailed below.
 - o Jane Annie Clarkson
 - o Susan Marie Smith



• No data entry is routinely needed if the necessary configuration exists for utilization of an app.

Facility	Community Health	& Ho	spitals	Site		USA (Community 1	
Patient Name:	First Name: Jane			Middle Name	e:	Last I	Name: As	
				Annie		appro	opriate	
	Previous Name: Jan	ie An	nie Clarkson	Suffix: Jr				
Sex:	Female (F)			DOB:		8/6/2	2020	
Race:	White (2106-3)			Ethnicity:		Not H	lispanic or Latino	
	European (2108-9)					(2186	5-5)	
More Granular	2108-9 (White Euro	pear	ו)	Preferred La	nguage	Engli	sh	
Race Code:								
Home Address:	1357 Amber Drive			Email		jawill	iams@gmail.com	
	Beaverton, OR 9700	06					-	
Telephone	Mobile: 555-335-12	234		Insurance/Pa	ayer	Com	oany: Blue Cross	
Number:	Home: 555-336-154	14			-	Blue	Shield; Plan 61-	
						Mana	aged Care	
Contact	Name: Marie Willia	ms		Contact		Name	e: Donald Williams	
	Relationship: Moth	Relationship: Mother				Relationship: Father		
	Address: 1357 Amb	er Di	rive			Addr	ess: 1357 Amber	
	Beaverton, OR 9700	06				Drive		
	Phone: 555-723-123	34				Beaverton, OR 97006		
					Phon	e: 555-723-1544		
Encounter/	Admit Type	Adn	nitting	Attending		Admi	tting Diagnosis	
Admit		Phy	sician	Physician	ysician		5 5	
Date/Time								
Current date	Emergency	Dr. I	Henry Seven	Dr. Henry Se	ven	D63.1	L Anemia in chronic	
(don't backdate)						kidne	y disease	
Problems	SNOMED CT Code		Proble	m Name	Healt	n	Start Date	
					Concer	'n	End Date	
					Status			
	87522002		Iron deficien	cy anemia	Active		2022	
	64667001		Interstitial p	neumonia	Active		January 2023	
Medication Allergies:	Allergy Substance		Reaction	Severity	Status		Date	
	Penicillin G benzath	ine	Hives	Moderate	Active		1/15/2021	
	RxNorm 7980		SNOMED -CT	r			,,	
			247472004					
	Ampicillin		Hives	Moderate	Active		1/20/2023	
	RxNorm 733		SNOMED -C	г			, , , , , ,	
			247472004					
Medications:			Dava	Erequency	Ro	ute	Start Date	
	Medication		Dose	Trequency			End Date	
Home Med &	Medication Tylenol		500 mg	Every 4 hrs	01	al	End Date As appropriate	
Home Med & Continue on	Medication Tylenol RxNorm 209459		500 mg	Every 4 hrs PRN	01	al	End Date As appropriate based on	



Home Med &	Torsemide		20 mg	BI	D	(Dral	As app	oropriate
Continue on	RxNorm 209	9459						based	on
admission	Indication -	Edema						admis	sion date
Immunizations	Vaccine			Da	te	Origi	n of info		
(Historical)									
	Hep B, unspecified form(45)		ulation (CVX	4/1,	/23	Parei	nt/Guaro	dian/Pa	atient Recall
	Hep B, unsp 45)	ecified form	ulation (CVX	5/1,	/23	Parei	nt/Guaro	dian/Pa	atient Recall
Not Given-	DTAP, 5, Pe	rtussis Antige	ens	6/5,	/23	Parei	nt/Guard	dian/Pa	atient Recall
Patient Refusa	(CVX=106)								
Lab Tests:	Test ordere	d	Test re	esults		LOIN	IC Codes	5	
Date/time	Urinalysis N	lacro	Color-Yellow			Color	LOINC	5778-6	
specimen	dipstick		Appearance-0	Clear		Appe	arance l	OINC	5767-9
collected: Day of			Specific gravit	y- 1.01	15	Speci	fic gravi	ty LOI	NC 5811-5
Admission			pH- 5.0			pH L	OINC 58	03-2	
			Glucose- 50 m	ng/dL		Gluce	ose LOIN	IC 5792	2-7
			Ketones- Neg	ative		Keto	nes LOIN	IC 579	7-6
			Protein- 100 r	ng/dL		Prote	in LOIN	C 5804	-0
Date/time	Prothrombi	n time	Lab Interface	results	s with	LOIN	C 34714	-6	
specimen	w/INR		INR=1.5 and Comment :		ent =				
collected: Day of			MD notified						
Admission									
Consults:	Consult to C	Cardiology-Dr	. Anderson						
Care Team	Mary McDo	nald	Adminis Healthca	trative re Staf	e ff				
Vitals							BMI		
	Height	85 cm	Temperature		38C		Percent	ile	56%
							FHIR On	ly -	
	Weight	12 kg	Respiratory R	ate	18/mi	in	HEIGHT	•	UNK
							Weight	for	
	Blood	145/88					Length		
	Pressure	mmHg	02		95%		Percent	ile	51%
							Head Oo	ccipital	
							Frontal		
	Heart						Circumf	erence	2
	Rate/Pulse	80/min	Flow Rate		4 L/M	in	Percent	ile	18%
							BMI		
	– – .	AL 1 /	Inhaled Oxyge	en	36%		Percent	ile	56%
Notes:	Exam Date	NOTE/Tab	and Duc to d	- T - I	conte	nt			0047004
		Surgical Hist	ory-procedure	es lab	Bronc	nosco		IVIED 1	U84/UU1
	Aumission				culture	ient:	sputum	collect	ed for
	Day of	Surgical Hist	on-Procedure	as Tab	Chest	e. Xrav	SNOME	D. 160	731000
	Admission		.ory rioceuult	23 100	Chest	лау	SINCIVIE	J. 100	, 51005
		1							



	Surgical History-Procedures Tab Introduction of cardiac pacemaker
	system via vein SNOMED ICD10PCS:
	175135009
	Procedure Narrative: Pacemaker is
	operating properly and clinical
	symptoms are unrelated.
	Check Implantable Device
	Implantable Device Pop-up
	(01)00643169007222(11)151201(17)160128(10)A213B2(21)BLC200461H
	(will need to change a digit in the red portion to get a unique device;
	Device Code = 704708004
Day of	ONC Clinical Notes
Admission	Procedure Narrative: None as already entered
(after adm)	Consult Narrative: Text as appropriate (content does not matter)
	Imaging Narrative: Bilateral Chest Xray: Inflammation and patchy
	consolidations, mainly in the right upper lobe.
	Pathology Narrative: Sputum from Bronchoscopy showed evidence of
	ongoing issues with pneumonia. Culture results pending.
Day of	Progress Note Physician
Admission	Clinical Impression: Patient admitted due to developing high fever and
(after adm)	since has shown considerable improvement and can be discharged
	shortly
Day of	History & Physical-Patient/History Tab
Admission	Current Smoking Status: Current every day smoker
(after adm)	
Day of	ECG/EKG Reports:
Admission	EKG-Interpretation/Impression: Normal EKG
(after adm)	
Day of	Health Concerns & Goals
Admission	Health Concerns:
(after adm)	Condition: Health concerns related to documented anemia problem
	Status: Active
	Onset: 2022
	Goals-Patient Goal: Need to gain more energy to do regular activities
Day of	Discharge Summary-Physician
Admission	Hospitalization Summary-Chief Complaint: Shortness of breath
(after adm)	Hospitalization Summary-Hospital Course: Text of your choice
	Labs/Imaging-Laboratory Results Review: Normal results
	Condition/Disposition-Diagnoses: No additional Dx needed
	Condition/Disposition-Functional Status Level: Level: Requires
	assistance
	Impairment: Dependence on walking stick
	Condition/Disposition-Cognitive Status Level: Level: Alert/disoriented
	Impairment: Amnesia
	Instructions:
	Activities ordered: Activities as tolerated



	Discharge S	ge Summary Narrative: Text of your choice					
	General Ins	structions: Schedule follow-up appt with Dr. Seven					
	Follow-up-A	-up-Assessment: Text of your choice					
	Follow-up-F	ollow-up-Plan: Text of your choice					
Discharge	Discharge Date/Time	Discharge Disposition:	Discharge Diagnosis:				
	After all clinical	Discharge to Home or Self	D63.1 Anemia in chronic kidney				
	documentation	Care	disease				

	Community	0 11-	a natital a	C:+-			Community of	
	Community Health	& HC	ospitais	Site		USA		
Patient Name:	First Name: Susan			Middle Nam	ie:	Last N	Name: As	
				Marie		appro	opriate	
	Previous Name: Sus	an N	Aarie Smith	Suffix: Jr				
Sex:	Female (F)			DOB:		8/1/2	020	
Race:	White (2106-3)			Ethnicity:		Not ⊦	lispanic or Latino	
	European (2108-9)					(2186	i-5)	
More Granular	2108-9 (White Euro	pear	า)	Preferred La	anguage	Englis	sh	
Race Code:								
Home Address:	1357 Amber Drive			Email		jawill	iams@gmail.com	
	Beaverton, OR 9700)6						
Telephone	Mobile: 555-335-12	. <mark>34</mark> ⊦	lome: <mark>555-</mark>	Insurance/F	Payer	Comp	any: Blue Cross	
Number:	336-1544					Blue S	Shield; Plan 61-	
						Mana	ged Care	
Contact	Name: Marie Smith			Contact		Name: Donald Williams		
	Relationship: Mothe	er				Relat	ionship: Father	
	Address: 1357 Amb	er D	rive			Addre	ess: 1357 Amber	
	Beaverton, OR 97006					Drive		
	Phone: 555-723-123	34				Beave	erton, OR 97006	
						Phon	e: 555-723-1544	
Encounter/	Admit Type	Adn	nitting	Attending		Admi	tting Diagnosis	
Admit		Phy	sician	Physician				
Date/Time								
Current date	Emergency	Dr.	Henry Seven	Dr. Henry S	even	D63.1 Anemia in chronic		
(don't backdate)	U ,					kidney disease		
Problems	SNOMED CT Code		Proble	m Name	Healt	5	Start Date	
					Concer	'n	End Date	
					Status			
	87522002		Iron deficier	icv anemia	Active		2022	
	64667001		Interstitial p	neumonia	Active		January 2023	
Medication	Alleray Substance		Reaction	Severity	Status		Date	
	Ancigy Substance		neaction	Sevency	Status		Duic	
	Penicillin G benzath	ino	Hives	Moderate	Active		1/15/2021	
	RyNorm 7980	inte	SNOMED -C	T	Active		1/13/2021	
	171101111 / 300		2/17/172004	'	1			
	Ampicillin		Livos	Moderate	Activo		1/20/2022	
	RyNorm 722			T	Active		1/20/2023	
	171101111/22		3110IVIED -C	'	1			
			247472004					



Medications:	Medication		Dose	Frequency	Ro	oute	Start L	Date
Home Med 8	Tulonol		E00 mg	Even Abre		ral	End Do	ate
	RyNorm 200	9459	500 mg	EVELY 4 HIS	0	Idi	As app hased	on
Admission	Indication -	Pain		T IXIN			admise	sion date
Home Med &	Torsemide	- uni	20 mg	BID	0	ral	As app	ropriate
Continue on	RxNorm 209	9459	20118	515		. ui	based	on
admission	Indication -	Edema					admiss	sion date
Immunizations	Vaccine		L	Date	Origin	of info		
(Historical)					- 0			
	Hep B, unsp 45)	ecified form	ulation (CVX	4/1/23	Paren	t/Guaro	dian/Pa	atient Recall
	Hep B, unsp 45)	ecified form	ulation (CVX	5/1/23	Paren	t/Guaro	dian/Pa	atient Recall
Not Given-	DTAP, 5, Pe	rtussis Antig	ens	6/5/23	Paren	t/Guaro	dian/Pa	tient Recall
Patient Refusa	(CVX=106)							
Lab Tests:	Test ordere	d	Test r	esults	LOIN	C Codes	5	
Date/time	Urinalysis M	lacro	Color-Yellow		Color	LOINC	5778-6	
specimen	dipstick		Appearance-	Clear	Appearance LOIN		LOINC	5767-9
collected: Day of			Specific gravi	ty- 1.015	Specific gravity LOINC 5811-5			NC 5811-5
Admission			рН- 5.0		pH_LOINC 5803-2			
			Glucose- 50 r	ng/dL	g/dL Glucose LOINC 5792-7		2-7	
			Ketones- Neg	gative	Keton	Ketones LOINC 5797-6		
			Protein- 100	mg/dL	Protein LOINC 5804-0			
Date/time	Prothrombi	n time	Lab Interface results with		LOINC	34714	-6	
specimen	w/INR		INR=1.5 and Comment					
collected: Day of			MD notified					
Admission								
Consults:	Consult to C	Cardiology-Di	r. Anderson					
Care Team	Mary McDo	nald	Admini	strative				
			Healthca	are Staff				
Vitals					E	BMI		
	Height	85 cm	Temperature	38C	F	Percent	ile	56%
					F	HIROn	ıly -	
	Weight	12 kg	Respiratory R	late 18/m	nin F	IEIGHT		UNK
					N	Neight	for	
	Blood	145/88			L	ength		
	Pressure	mmHg	02	95%	F	ercent	ile	51%
					ŀ	iead Oo	ccipital	
	lloart				ŀ		0.0000-	
		leo PO/min Flow Pote		Circumference			1.8%	
	nate/Pulse			4 L/N	/1111 F		IIE	10/0
			Inhaled Ovyg	en 36%) Percent	ile	56%
Notes	Exam Date	Note/Tab		Cont	r ≏nt	creent		
10103.			Cont	CIIC				



Admissio	Surgical History-Procedures Tab Bronchoscopy SNOMED 10847001
	n Comment: Sputum collected for
	culture.
Day of Admissio	Surgical History-Procedures Tab Chest Xray SNOMED: 168731009
	Surgical History-Procedures Tab Introduction of cardiac pacemaker
	system via vein SNOMED ICD10PCS:
	175135009
	Procedure Narrative: Pacemaker is
	operating properly and clinical
	symptoms are unrelated.
	Check Implantable Device
	Implantable Device Pop-up
	(01)00643169007222(11)151201(17)160128(10)A213B2(21)BLC200461H
	(will need to change a digit in the red portion to get a unique device;
	Device Code = 704708004
Day of	ONC Clinical Notes
Admissio	Procedure Narrative: None as already entered
(after adr	m) Consult Narrative: Text as appropriate (content does not matter)
	Imaging Narrative: Bilateral Chest Xray: Inflammation and patchy
	consolidations, mainly in the right upper lobe.
	Pathology Narrative: Sputum from Bronchoscopy showed evidence of
	ongoing issues with pneumonia. Culture results pending.
Day of	Progress Note Physician
Admissio	n Clinical Impression: Patient admitted due to developing high fever and
(after adr	m) since has shown considerable improvement and can be discharged
	shortly
Day of	History & Physical-Patient/History Tab
Admissio	Current Smoking Status: Current every day smoker
(after adr	m)
Day of	ECG/EKG Reports:
Admissio	n EKG-Interpretation/Impression: Normal EKG
(after adr	(II) Health Concorns & Coals
Day of Admissio	Health Concerns & Goals
Aumissio (after adv	 medium concerns: related to documented enemia problem
(arter au	Status: Active
	Onset: 2022
	Goals-Patient Goal: Need to gain more energy to do regular activities
Day of	Discharge Summary-Physician
Day Of	Hospitalization Summary-Chief Complaint: Shortness of breath
Admissio	m) Hospitalization Summary-Hospital Course: Text of your choice
Admissio (after adr	Labs/Imaging-Laboratory Results Review: Normal results
Admissio (after adi	
Admissio (after adi	Condition/Disposition-Diagnoses: No additional Dx needed
Admissio (after adi	Condition/Disposition-Diagnoses: No additional Dx needed Condition/Disposition-Functional Status Level: Level: Requires
Admissio (after adi	Condition/Disposition-Diagnoses: No additional Dx needed Condition/Disposition-Functional Status Level: Level: Requires assistance
Day of Admissio (after adr Day of Admissio (after adr Day of	ECG/EKG Reports: n EKG-Interpretation/Impression: Normal EKG m) Health Concerns & Goals Health Concerns: m) Condition: Health concerns related to documented anemia problem Status: Active Onset: 2022 Goals-Patient Goal: Need to gain more energy to do regular activities Discharge Summary-Physician m Hospitalization Summary-Chief Complaint: Shortness of breath m) Hospitalization Summary-Hospital Course: Text of your choice Labs/Imaging-Laboratory Results Review: Normal results



	Condition/D	Condition/Disposition-Cognitive Status Level: Level: Alert/disoriented				
	Impairment	Impairment: Amnesia				
	Instructions	Instructions:				
	Activities or	dered: Activities as tolerat	ed			
	Dietary insti	ructions: Renal				
	Discharge Si	Discharge Summary Narrative: Text of your choice				
	General Inst	ructions: Schedule follow-	up appt with Dr. Seven			
	Follow-up-A	ssessment: Text of your ch	noice			
	Follow-up-P	Follow-up-Plan: Text of your choice				
Discharge	Discharge Date/Time	Discharge Disposition:	Discharge Diagnosis:			
	After all clinical	Discharge to Home or Self	D63.1 Anemia in chronic kidney			
	documentation	Care	disease			

7.7.3 Test Script Prerequisites

• Configuration of external application such as Apple Health[®] or the Inferno test tool if no external applications are configured/utilized

7.7.4 Measure Data

• Measure g10 % patients evaluated that are validated/do not generate any errors

Numerator= # patients evaluated using the Inferno test tool/external application that passed validation/did not generate any errors

Denominator = # patients evaluated using the Inferno test tool/external application

Testing must be application specific. Testing using the Inferno Test Tool is detailed below.

Step	Steps to Validate Inferno Test Tool	Expected Outcome
Step 1	Navigate to Inferno Test Tool and select Create Test Session.	https://inferno.healthit.gov/t
		est-kits/g10-certification/
Step 2	Select test 1: Standalone Patient App and click Run Tests and fill in required fields.	Inferno Tool Runs
Step 3	Click Submit for Test Case 1.	Inferno Tool brings up patient
Ston /	Login as nations, select nations and click ves, allow at bottom of screen	Inferno Runs test and receive
Step 4	Login as patient, select patient and energys, anow at bottom of selecti.	all green check marks.
Step 5	Select test 2: Limited Access App and click Run Tests and then Submit.	Inferno Tool brings up pop
		up.
Step 6	Click the link provided in pop-up and then select your patient and then	Inferno Tool runs test and
	select only Patient, Condition and Observation.	receive all green check
		marks.



Step 7	Select test 3: EHR Practitioner App and fill in required fields and select	Inferno Tool brings up a pop-
	Run Tests.	up EHR Practitioner App.
Sten 8	Navigate to Juno FHB and select Inferno button	luno FHR brings you back to
otep o		the ConnectEHB server to
		select patient
Step 9	Select patient from list and click ves, allow at bottom of page.	Inferno runs test and receive
otep 5		all green check marks
Sten 10	Select test 4 [.] Single Patient API and click Run Tests	Inferno runs all tests and
5100 10		receive all green check
		marks.
Step 11	Select test 7: Multi-Patient API and select Run Tests and fill in required	All fields are filled in properly
	fields.	
Step 12	Click Submit.	Inferno runs test and receive
		all green check marks.
Step 13	Select test 9.1 SMART Public Client Launch and fill in required fields and	Inferno will run test and you
	click Submit.	will receive a pop-up.
Step 14	Select the link in the pop-up and login to the patient, select patient and	Inferno will run test and
	click ves. allow at bottom of the screen.	receive all green check
		marks.
Step 15	Select test 9.3 Token Revocation and fill in required fields.	All required fields are
		populated.
Step 16	Access Postman and enter the bearer token into Postman and select	Receive an active status in
	Send.	Postman.
Stop 17	Login to ConnectEHP as nations and select Povoke Access	Access has been revoked for
Step 17	Login to connecterin as patient and select nevoke Access.	the nationt
Ston 18	Click Submit for test 9.3	Inferno will run test and
5100 10		receive all green check
		marks
Step 19	Select test 9.4: SMART Invalid AUD Launch and click Submit	Inferno brings up a pop-up.
0100 10		
Step 20	Select the "perform invalid launch" link.	Directed to ConnectEHR
		Identity Server and receive
		an error.
Step 21	Select the back button and click "Attest Launch Failed".	Inferno runs test and receive
		all green check marks.
Step 22	Select 9.5 SMART Invalid Token Request and click Run Test and click	Inferno brings up a pop-up.
	submit.	



Step 23	Select the "follow this link to authorize the SMART server".	Directed to the ConnectEHR
		Identity Server.
Step 24	Select patient and confirm scopes and select yes, allow at bottom of the	Inferno runs test and receive
	screen.	all green check marks.
Step 25	Select test 9.8 EHR Launch with Patient Scopes and select Run Tests and	Inferno Tool brings up a pop-
	fill in required fields. Click Submit.	up EHR Practitioner App.
Step 26	Navigate to Juno EHR and select the Inferno button.	Juno EHR brings you back to
		the ConnectEHR server to
		select patient.
Step 27	Click the follow this link button on pop-up and select patient. Click yes,	Inferno runs test and receive
	allow at bottom of screen.	all green check marks.
Step 28	Select test 9.10 Visual Inspection and select Run Tests and fill in required	All fields are filled in properly.
	fields.	
Step 29	Click Submit.	Inferno runs test and receive
		all green check marks.
Step 30	Select Report at bottom of test list.	Report is exported and
		available.