RESOURCE AND PATIENT MANAGEMENT SYSTEM

Consolidated Clinical Document Architecture (CCDA)

(BCCD)

User Manual

Version 2.0 Patch 1
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Office of Information Technology
Division of Information Technology
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Preface

This manual provides information for site managers regarding operations of the Consolidated Clinical Document Architecture (CCDA) v2.0 (BCCD) package, including patch 1 updates. This manual is not intended for end users of RPMS since there is no end user functionality in this package.

The BCCD package is designed to generate industry standard Continuity of Care Documents (CCD) in Health Level 7 (HL7) CCDA format, following the August 2015 Draft Standard for Trial Use (DSTU) standard, further restricted by 2015 Certified Health Information Technology (CHIT) requirements, the 2015 Edition Cures Update (CURES) and the U.S. Core Data for Interoperability version (USCDI) 2.0 requirements. These documents can be transmitted to Indian Health Service (IHS) Health Information Exchange (HIE) repositories, retrieved by the Electronic Health Record (EHR) Graphical User Interface (GUI) using web services, or viewed as export summary documents.
1.0 **Introduction**

The Consolidated Clinical Document Architecture (namespace BCCD) software is a component of the IHS Resource and Patient Management System (RPMS). The software provides the ability to generate industry standard Clinical Summary, Transitions of Care, Export Summary, Health Information Exchange (HIE) and Case Reporting documents that meet the Health Level Seven (HL7) August 2015 DSTU standard, the 2015 CHIT specifications, the 2015 CURES specifications and the USCDI version 2.0 requirements.

Clinical Summary and Transitions of Care documents are retrieved by the EHR GUI using web services. Data Export summary documents are requested via an RPMS Application Programming Interface (API). HIE documents are generated by a nightly background job or via an RPMS menu option and are transmitted to IHS HIE repositories via web services. Case Reporting documents are requested by the iCare (BQI) application, which retrieves completed documents for transmission.

1.1 **Purpose**

CCDA documents can serve a variety of purposes, including enabling clinician access to patient data in an emergency scenario, quality reporting, bio surveillance, patient access to the patient's own data via a Personal Health Record (PHR) system, and medication or allergy reconciliation.

Each CCDA document consists of two components: a human readable part known as a narrative block, which can be displayed by a web browser; and a machine-readable part intended for automated data processing. The machine-readable component may contain more detailed information than the human-readable part.

1.2 **Scope**

A CCDA document is an Extensible Markup Language (XML) document summarizing current and pertinent historical information about an individual patient’s health care record at a given facility. Although not every document type contains every section, the current IHS implementation of the CCDA standard supports the following sections:

- Allergies and Intolerances
- Assessment and Plan
- Care Team
- Case Reporting
- Consultation Notes
- Discharge Diagnoses (inpatient visits only)
- Discharge Summary Notes
- Functional Status
- Goals
- Health Concerns
- History and Physical Notes
- Hospital Course (inpatient visits only)
- Hospital Discharge Instructions (displayed for observation (“O”) and day surgery (“S”) visits, in addition to inpatient visits)
- Immunizations
- Medical Equipment (Implantable Devices)
- Medications
- Mental/Cognitive Status
- Patient
- Problems/Encounter Diagnoses
- Procedures and Interventions
- Progress Notes
- Reason for Hospitalization (inpatient visits only)
- Reason for Referral
- Reason for Visit (ambulatory visits only)
- Recent Encounters
- Recent Test Results
- Recent Vital Signs
- Security and Privacy Prohibitions
- Social History (Smoking Status)
2.0 **System Navigation**

2.1 **Setup — CCDA**

To set up CCDA at an RPMS site:

1. Calculate the additional disk space required for the BCCD application and allocate disk space.

2. Perform pre-install steps, including enabling Long Strings, verifying access to OS configuration file, designating a directory where the new CCDA database will reside, mapping BMW globals, and stopping CCDA.

3. Install the provided Kernel Installation and Distribution System (KIDS) build.

4. Perform post-install steps, including setting up optional encryption, optional e-mail notifications, disabling journaling in the CCDA database, configuring the CCDA CSP Application, setting up site-specific BCCD parameters, and starting CCDA Messaging.

See the CCDA Installation Manual for further details on the installation and configuration steps.

The BCCD package has one RPMS menu used by site managers, CCDA MENU (BCCD MENU), which presents the following options:

- BCCD Notes Definition Report (BCCD NOTES DEFINITION REPORT)
- Edit CCDA Site Parameters (BCCD EDIT SITE PARAMETERS)
- Edit CCDA Clinical Site Parameters (BCCD EDIT CLINICAL PARAMETERS)
- Generate CCDA for a single patient (BCCD CCDGENONE)
- Generate CCDA documents for all patients in RPMS (BCCD CCDALLGEN)
- Generate CCDA documents for a specified date range (BCCD GEN CCDA DATE RANGE)
- Manage CCDA transmissions (BCCD CCDA MGR)
- View and Purge Error Documents (BCCD ERROR PURGE)
- Display the CCDA application status report (BCCD APP STATUS)

The BCCD package contains no menu options accessible by end users.
Site managers are responsible for performing the following two tasks to ensure that CCDA documents are successfully generated at their site:

1. Ensure that site-specific CCDA parameters are set up appropriately. Section 3.1 provides more information about the set up.

2. Ensure that CCDA Messaging is running. Section 3.2 provides more information about managing CCDA Messaging.

### 2.2 Setup — Hospital Course Information

BCCD v2.0 added a new section to inpatient CCDA documents called Hospital Course. The information for the Hospital Course section comes the signed Discharge Summary TIU note associated with the Inpatient stay. However, the Discharge Summary note may include additional information not related to the Hospital Course. To extract only the Hospital Course information only in the CCDA from the Discharge Summary note, it is recommended that any Discharge Summary template have a Hospital Course header and footer. The header should be fifty (50) hyphens on their own line followed by the text "HOSPITAL COURSE" (without quotes) on its own line. The footer should follow the Hospital Course information and be fifty (50) hyphens on its own line.

Later, when the provider creates the Discharge Summary note for the patient, the Hospital Course information must be entered by the provider between the header and footer; otherwise, the CCDA will include all the information entered in the Discharge Summary note. To make it easier for the provider to identify what is the header and footer, you may want to consider entering some blank lines between the header and footer.

Please work with your Clinical Informaticists on entering the Hospital Course header and footer. An example of how to add the Hospital Course header and footer to a Discharge Summary template is shown in the figure below.
Figure 2-1: Adding Hospital Course Header and Footer to a Discharge Summary template
2.3 Architecture

CCDA processing can be initiated in four ways:

- If EHR version 1.1, patch 13 has been installed and configured at the site, then authorized EHR users are able to request Clinical Summary and Transitions of Care documents for display within the EHR GUI client.

- If the site is a part of the IHS HIE, then HIE documents are generated nightly for all patients whose data has changed since the previous nightly upload.

- Data Export summary documents can be generated for specific patients or for all patients via an RPMS API.

- Case Reporting documents are requested by the iCare (BQI) application, which retrieves completed documents for transmission.

To accommodate requests from the EHR GUI, Ensemble has a web service process listening for requests for Clinical Summary and Transitions of Care documents.

If the site is part of the IHS HIE, a nightly TaskMan task called BCCD NHIE PUSH JOB is automatically scheduled to run every night at the time specified by the site manager in option Edit CCDA Site Parameters. When BCCD NHIE PUSH JOB runs for the first time, it creates new CCDA requests for all patients in the RPMS database. On subsequent executions, BCCD NHIE PUSH JOB finds all patients whose data has changed since the last time it ran and creates new requests in the CCDA queue.

To request Data Export summary documents, the site manager accesses an RPMS API. In order to prevent unintended initiation of Data Export documents, the API is not included as an RPMS option in the menu system. Instead, the site manager must first access Programmer Mode, then enter the necessary command. Because it is expected that this Data Export functionality will rarely, if ever, be used, the IHS Software Quality Assurance (SQA) team will distribute instructions for generating Data Export documents on an “as needed” basis.

Case Reporting documents are requested and processed by the iCare (BQI) application. Details on this process are outside the scope of this document. For further information, consult the appropriate iCare documentation.

The only difference between CCDA requests created by these four mechanisms is that the DocType property is set to the requested document type and several flags are set to control where the documents are sent. For all CCDA requests, a new entry is placed in the CCDA queue, recording the IDs of the requested patient and visit(s), and sets the record’s status to R (for Request).
The CCDA queue is monitored by BCCD BACKGROUND JOB, the main CCDA generator. The BCCD BACKGROUND JOB is a persistent TaskMan job started at TaskMan startup time, or manually from menu option Manage CCDA transmissions which constantly runs in the background until the site manager chooses to stop CCDA. When the BCCD BACKGROUND JOB finds a new CCDA request with the status of R in the CCDA queue, it changes the request’s status to CS (for Compile Started) and extracts all relevant RPMS data for the specified patient and visit and adds the data to the body of the CCDA request. Depending on the amount of eligible data in the patient’s records, it may take anywhere from under a second to over 10 seconds to extract all needed data from RPMS.

Once the extraction process for the patient is finished, BCCD BACKGROUND JOB changes the status of the request to CE (for Compile Ended). The BCCD BACKGROUND JOB then checks the CCDA queue for other outstanding requests with the status of R. If no outstanding requests are found, the BCCD BACKGROUND JOB goes into hibernation for a specific amount of time, usually one second.

The CCDA queue is also monitored by the CCDA Ensemble production running in the CCDA namespace. When the CCDA production finds a new request whose status is set to CE, it retrieves the CCDA data from the body of the request, changes the status to TS (for Transmission Started), and transforms it into a valid CCDA document. For EHR Clinical Summary and Transitions of Care requests, the document is sent back to the requesting EHR. For Data Export and Case Reporting, the production saves this document to a file. For HIE documents, the production transmits it to an IHS HIE. Finally, the production changes the status of the request to T (for Transmitted).

If a site is a part of the IHS HIE, then the site manager is provided with the URL of the associated CCDA repository. Ensemble uses this URL to send CCDA documents to the repository. HIE CCDA processing typically occurs at night to minimize the impact on RPMS performance.
3.0 Package Management

3.1 Manage Site-Specific CCDA Parameters

The RPMS option Edit CCDA Site Parameters (BCCD EDIT SITE PARAMETERS) has been divided into two options with different security keys.

3.1.1 Edit CCDA Site Parameters

The option Edit CCDA Site Parameters (BCCD EDIT SITE PARAMETERS) allows site managers to configure site-specific CCDA parameters at installation time and modify them if necessary.

**Note:** This menu option is locked by security key XUMGR, which site managers already have.

The following four parameters can be edited using this option:

- **TIME TO RUN NIGHTLY TASK**
- **DAYS KEEP TRANSMISSION ENTRIES**
- **REPOSITORY LOCATION**
- **ENABLED**

The first prompt is “TIME TO RUN NIGHTLY TASK” and there is no default value. If the RPMS site is not a part of the IHS HIE at this time, then skip the “TIME TO RUN NIGHTLY TASK” prompt. If the RPMS site is a part of the IHS HIE and the site manager has been provided with the location of the off-site CCDA repository, then enter the time when the site wants CCDA documents to be generated and transmitted on a nightly basis. It is recommended to enter a time outside of the regular business hours to minimize impact on the end users. The “TIME TO RUN NIGHTLY TASK” prompt accepts time values in all valid FileMan formats, e.g., 15:34 or 03:34PM.

The second prompt is “DAYS KEEP TRANSMISSION ENTRIES” and it is set to 30 days by default. The value may be set to any value between 7 and 999 days. The higher the number, the more disk space the intermediate compile structures created by the CCDA application will consume as described in Section 3.3 of this installation guide. On the other hand, if the default value is set too low, the compile data may be purged too soon and will not be available to facilitate debugging should any problems arise. If the site manager is not sure what value to enter, accept the default value of 30 days.
The third prompt is “REPOSITORY LOCATION” and there is no default value. If the RPMS site is not a part of the IHS HIE at this time, then skip the REPOSITORY LOCATION prompt. If the RPMS site is a part of the IHS HIE and the site manager has been provided with the location of the off-site CCDA repository, then enter that location at the REPOSITORY LOCATION prompt exactly as it was provided to the site manager. When a value is entered for both the “REPOSITORY LOCATION” and the “TIME TO RUN NIGHTLY TASK” prompts, the system will check whether the CCDA database has enough disk space to accommodate the process of generating CCDA documents for all patients in the system. An error message will be displayed if there is not enough disk space. If not enough disk space error message displays, allocate more disk space and try again.

The fourth prompt is “ENABLED”, and the default value is NO. This NO value controls whether CCDA documents are generated when CCDA is running. Change this value to YES.

Figure 3-1 contains a screen capture of a typical setup session.

<table>
<thead>
<tr>
<th>Select CCDA Menu Option: EDIT CCDA Site Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Now editing CCDA parameters:</td>
</tr>
<tr>
<td>TIME TO RUN NIGHTLY TASK: 01:00</td>
</tr>
<tr>
<td>Now editing CCD (Summarization of Episode Note)-specific parameters:</td>
</tr>
<tr>
<td>DAYS KEEP TRANSMISSION ENTRIES: 30</td>
</tr>
<tr>
<td>REPOSITORY LOCATION:</td>
</tr>
<tr>
<td><a href="http://sample.ihs.gov:19090/PatientRecordReceiverService/PatientRecordReceiverService">http://sample.ihs.gov:19090/PatientRecordReceiverService/PatientRecordReceiverService</a></td>
</tr>
<tr>
<td>ENABLED: YES</td>
</tr>
</tbody>
</table>

Figure 3-1: Setting up CCDA Site Parameters

### 3.1.2 Edit CCDA Clinical Site Parameters

The option Edit CCDA Clinical Site Parameters (BCCD EDIT CLINICAL PARAMETERS) allows clinicians to configure site-specific clinical CCDA parameters at installation time and modify them if necessary. Note that this menu option is locked by security key BCCDZCLIN, which must be assigned before the option can be used.

The following four parameters can be edited using this option:

- ADD SOCIAL ENVIRONMENT PROB
- ADD INACTIVE PERS HISTORY PROB
- DISCHARGE PLANNER PROV TYPE
- CLINICAL NOTE
The first prompt is “ADD SOCIAL ENVIRONMENT PROB”, which is set to NO by default. This NO value determines whether social environment problems will be included in Clinical Summary documents for the site. All other document types will always display the social environment problems.

The second prompt is “ADD INACTIVE PERS HISTORY PROB”, which is set to NO by default. This NO value specifies whether inactive personal problems are included in Clinical Summary documents for site. All other document types will always display the inactive personal history problem.

The third prompt is “DISCHARGE PLANNER PROV TYPE”, which has no default value. This parameter specifies the BDP DESG SPECIALTY PROVIDER type to be extracted and displayed as the discharge planner on an inpatient CCDA document.

The fourth prompt is “Select CLINICAL NOTE”, which is a multiple value parameter with subsequent additional prompts. This parameter allows sites to designate what signed TIU documents should appear in the new Consultation Notes, Discharge Summary Notes, History and Physical Notes, Hospital Discharge Instructions, Progress Notes and Procedure Notes (found in the Procedures and Interventions) sections of the CCDA documents. To update this information, users will first be prompted to select one of the note types to enter or edit. Figure 3-2 shows the list of note choices to select from.

Select CLINICAL NOTE: ?
You may enter a new CLINICAL NOTE, if you wish
Enter the note type to set up a custom definition for.
Choose from:
C    CONSULTATION
H    H&P
D    DISCHARGE SUMMARY/HOSPITAL COURSE
P    PROCEDURE
I    DISCHARGE INSTRUCTIONS

Select CLINICAL NOTE:

Figure 3-2: Clinical Note Type Definition Choices

Once a CLINICAL NOTE type has been selected, the user is prompted to enter one or more INCLUDE TITLE/DOCUMENT CLASS items. For this prompt, the user can select from available TIU document classes and titles. Any patient note created with a title entered here or, if a document class is selected, all titles in that document class will be included in the chosen section of the CCDA document. After entering all of the INCLUDE TITLE/DOCUMENT CLASS selections, the user will be prompted to enter any EXCLUDE NOTE TITLE items which are limited to document titles. This prompt is useful if a site wishes to include a specified document class in a CCDA notes section but does not wish to have one or more titles in that class included in the specified section.
After entering information for all of the CLINICAL NOTE types, the user is given the opportunity to enter SITE NOTE EXCLUSION values. Document classes or titles entered in this section will be excluded from all CLINICAL NOTE document types. This field is particularly useful for controlling what appears in the Progress Note section of the CCDA document. Progress Notes are defined as any patient note whose document title is not listed in one of the CLINICAL NOTE definitions described above. It should be noted that if one of the above sections excludes a particular document title from be displayed in a specified section, then that document title, unless included in another CLINICAL NOTE section, will be included as a Progress Note. If a site wishes to have any document class or title excluded from every notes section (including the Progress Notes section) of the CCDA, then they can enter that document class or title in the SITE NOTE EXCLUSION prompt.

Figure 3-3 contains a screen capture of a typical setup session.

```plaintext
Select CCDA Menu Option: CLIN  Edit CCDA Clinical Site Parameters

Edit CCDA Clinical Site Parameters

NOW EDITING CCDA CLINICAL PARAMETERS:

ADD SOCIAL ENVIRONMENT PROB: NO//
ADD INACTIVE PERS HISTORY PROB: NO//
DISCHARGE PLANNER PROV TYPE:
Select CLINICAL NOTE: CON (C  CONSULTATION)
  Are you adding 'CONSULTATION' as a new CLINICAL NOTE (the 1ST for this
  BCCD MESSAGE TYPE)? No// Y (Yes)
  Select INCLUDE TITLE/DOCUMENT CLASS: CONSULTS DOCUMENT CLASS
  DOCUMENT CLASS
  Are you adding 'CONSULTS' as
  a new INCLUDE TITLE/DOCUMENT CLASS (the 1ST for this CLINICAL NOTE)?
  No// Y (Yes)
  Select INCLUDE TITLE/DOCUMENT CLASS:
  Select EXCLUDE NOTE TITLE: CHS TITLE TITLE
  Are you adding 'CHS' as a new EXCLUDE NOTE TITLE (the 1ST for this
  CLINICAL NOTE)? No// Y (Yes)
  Select EXCLUDE NOTE TITLE:
  Select CLINICAL NOTE: DISCHARGE I (I  DISCHARGE INSTRUCTIONS)
  Are you adding 'DISCHARGE INSTRUCTIONS' as a new CLINICAL NOTE (the 2ND
  for this BCCD MESSAGE TYPE)? No// Y (Yes)
  Select INCLUDE TITLE/DOCUMENT CLASS: DISCHARGE INS
  1  DISCHARGE INSTRUCTIONS DOCUMENT CLASS DOCUMENT CLASS
  2  DISCHARGE INSTRUCTIONS TITLE TITLE
  CHOOSE 1-2: 1  DISCHARGE INSTRUCTIONS DOCUMENT CLASS DOCUMENT CLASS
  Are you adding 'DISCHARGE INSTRUCTIONS' as
  a new INCLUDE TITLE/DOCUMENT CLASS (the 1ST for this CLINICAL NOTE)?
  No// Y (Yes)
  Select INCLUDE TITLE/DOCUMENT CLASS:
  Select EXCLUDE NOTE TITLE:
  Select CLINICAL NOTE: DISCHARGE SU (D  DISCHARGE SUMMARY/HOSPITAL COURSE)
  Are you adding 'DISCHARGE SUMMARY/HOSPITAL COURSE' as
  a new CLINICAL NOTE (the 3RD for this BCCD MESSAGE TYPE)? No// Y (Yes)
  Select INCLUDE TITLE/DOCUMENT CLASS: DISCHARGE SU
  1  DISCHARGE SUMMARIES DOCUMENT CLASS DOCUMENT CLASS
  2  DISCHARGE SUMMARY TITLE TITLE
  3  DISCHARGE SUMMARY DOCUMENT CLASS DOCUMENT CLASS
  4  DISCHARGE SUMMARY TITLE TITLE
```
### 3.1.3 BCCD Notes Definition Report

CLINICAL NOTE information entered in Section 3.1.2 can be displayed in the BCCD Notes Definition Report (BCCD NOTES DEFINITION REPORT). A sample display of the report is found in Figure 3-4.

```plaintext
| Select OPTION NAME: BCCD NOTES DEFINITION REPORT | BCCD Notes Definition Report |
```

Figure 3-3: Setting up CCDA Clinical Site Parameters
This report will display the 'Edit CCDA Clinical Site Parameters' option current definition for the clinical notes sections on the CCDA documents. These settings determine what TIU note titles show up in each section.

Print report on which device: HOME// 0;80;999  VIRTUAL

BCCD Notes Definition Report  APR 28, 2022  16:04  PAGE 1

Discharge Summary/Hospital Course
(in Discharge Summary/Hospital Course Notes sections)

Included Documents:
  DISCHARGE SUMMARY which is in DISCHARGE SUMMARIES (class) which is in DISCHARGE SUMMARY (class)

Excluded Documents:
  TELEPHONE

Consultation Note (in Consultation Notes section)

Included Documents:
  CHS which is in CONSULTS (class)
  REFERRALCONSULT which is in CONSULTS (class)
  RADIOLGYCONSULT which is in CONSULTS (class)
  DIABETESCONSULT which is in CONSULTS (class)
  DIABETESENDOCRINOLOGYCONSULT which is in CONSULTS (class)

Excluded Documents:
  CHS

History and Physical Note (in History and Physical Notes section)

Included Documents:
  ADVERSEREACTION/ALLERGY

Excluded Documents:
  No Excluded Documents defined for this note type

Procedures Note (in Procedures section)

Included Documents:
  GENERAL

Excluded Documents:
  TELEPHONE

Hospital Discharge Instructions (in Hospital Discharge Instructions section)

Included Documents:
  DISCHARGEINSTRUCTIONS which is in DISCHARGE INSTRUCTIONS (class)

Excluded Documents:
  No Excluded Documents defined for this note type
Progress Note (in Progress Notes Section)

All TIU Note Title(s) except:

- ADVERSE REACTION/ALLERGY which is in History and Physical Notes
- DIABETESCONSULT which is in Consultation Notes
- DIABETES_ENDOCRINOLOGY_CONSULT which is in Consultation Notes
- DISCHARGE_INSTRUCTIONS which is in Hospital Discharge Instructions
- DISCHARGE_SUMMARY which is in Discharge Summary Notes
- GENERAL which is in Procedures Notes
- RADIOLGY_CONSULT which is in Consultation Notes
- REFERRAL_CONSULT which is in Consultation Notes

General Exclusions:

-------------------
TELEPHONE

Imaging Narrative (in Recent Test Results section)

The imaging impression is a short description or summary of a patient’s exam results report. For legal purposes, it is strongly recommended that an impression always be entered for every patient imaging report.

Laboratory Report Narrative (in Recent Test Results section)

The lab comment is a short description or summary of the reference laboratory results and/or comments from the laboratory department.

<End of Report>

Press RETURN to continue or '^' to exit:

Figure 3-4: Sample CCDA Notes Definition Report

3.2 Manage CCDA Messaging

CCDA Messaging is automatically started whenever TaskMan starts, which typically occurs when Ensemble is started. CCDA Messaging will not run if TaskMan is not running.

CCDA Messaging should be running at all times and does not require a manual start or stop once it has been installed. However, if the need arises, CCDA Messaging can be brought up or down using the Manage CCDA transmissions option, which is available in the CCDA Menu. The CCDA Menu can be accessed from the regular “OPTION NAME” prompt in RPMS.
If CCDA Messaging is not running, type **YES** at the prompt as shown in Figure 3-5:

<table>
<thead>
<tr>
<th>Select CCDA Menu Option: MANG Manage CCDA transmissions</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCDA status:</td>
</tr>
<tr>
<td>No configuration problems found</td>
</tr>
<tr>
<td>CCDA processing task is not running</td>
</tr>
<tr>
<td>Start CCDA? No// <strong>YES</strong> (Yes)</td>
</tr>
<tr>
<td>Attempting to start CCDA  (JUL 16, 2013@16:09:54)...CCDA started</td>
</tr>
</tbody>
</table>

Figure 3-5: Starting CCDA Messaging

If CCDA Messaging is running, type **NO** at the prompt as shown in Figure 3-6:

<table>
<thead>
<tr>
<th>Select CCDA Menu Option: MANG Manage CCDA transmissions</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCDA Messaging status:</td>
</tr>
<tr>
<td>No configuration problems found</td>
</tr>
<tr>
<td>CCDA processing task is running</td>
</tr>
<tr>
<td>Stop CCDA? No// <strong>Y</strong> (Yes)</td>
</tr>
<tr>
<td>Attempting to stop CCDA....CCDA stopped</td>
</tr>
</tbody>
</table>

Figure 3-6: Stopping CCDA Messaging

### 3.3 Manage CCDA Ensemble Production

As described in Section 2.2, if the RPMS site is a part of the IHS HIE, CCDA documents are sent to outside repositories using the CCDA Ensemble production. In addition, if EHR version 1.1, patch 13 is installed and configured, authorized EHR users can send requests for CCDA documents to the CCDA Ensemble production.

The CCDA Ensemble production is automatically started when HealthShare starts up. During normal business operations, the CCDA Ensemble production will remain running and will not require maintenance. Refer to the Managing CCDA Ensemble Production section in the CCDA Technical Manual to bring the CCDA Ensemble production up or down manually, if required.

### 3.4 CCDA Application Status Report

The STAT (Display the CCDA application status report) option may be used to display a report summarizing various information about the state of the CCDA application, as well as information about the most recent pull, push, and error messages. The intent of the report is to provide support personnel with a snapshot of the application. It is not intended to be used by site managers, and no information is provided to assist site managers with interpreting the report.
When the user runs the report, it will ask whether to display the report with page breaks. If the user intends to view the report on the screen, it is recommended that page breaks be used. If the user is running the report in order to copy and paste information for the support team, then not using page breaks may be preferred.

A sample report is shown in Figure 3-7:

```
When the user runs the report, it will ask whether to display the report with page breaks. If the user intends to view the report on the screen, it is recommended that page breaks be used. If the user is running the report in order to copy and paste information for the support team, then not using page breaks may be preferred.

A sample report is shown in Figure 3-7:
```

```
<table>
<thead>
<tr>
<th>QID</th>
<th>DFN</th>
<th>Type</th>
<th>Status</th>
<th>RequestTS</th>
<th>TransmissionEndTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>5422</td>
<td>7207</td>
<td>CS</td>
<td>T</td>
<td>2018-04-03 15:06:06</td>
<td>2018-04-03 15:06:06</td>
</tr>
<tr>
<td>658</td>
<td>174</td>
<td>TC</td>
<td>E</td>
<td>2018-03-19 14:47:55</td>
<td></td>
</tr>
<tr>
<td>638</td>
<td>174</td>
<td>TC</td>
<td>E</td>
<td>2018-03-19 14:35:31</td>
<td></td>
</tr>
<tr>
<td>628</td>
<td>174</td>
<td>TC</td>
<td>T</td>
<td>2018-03-19 14:34:22</td>
<td>2018-03-19 14:34:36</td>
</tr>
</tbody>
</table>
```

```
<table>
<thead>
<tr>
<th>QID</th>
<th>DFN</th>
<th>Type</th>
<th>Status</th>
<th>RequestTS</th>
<th>TransmissionEndTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>5477</td>
<td>7207</td>
<td>HIE</td>
<td>T</td>
<td>2018-04-19 12:34:45</td>
<td></td>
</tr>
<tr>
<td>5476</td>
<td>7207</td>
<td>HIE</td>
<td>T</td>
<td>2018-04-19 12:30:57</td>
<td></td>
</tr>
<tr>
<td>5475</td>
<td>7207</td>
<td>HIE</td>
<td>T</td>
<td>2018-04-19 11:40:00</td>
<td></td>
</tr>
</tbody>
</table>
```
Due to different rules surrounding patient DFNs, some sites may want to prevent DFNs from displaying on the report. To suppress DFNs site-wide, go to RPMS programmer mode and run the following command:

```
DO DFNOFF^BCCDSTAT
```

To clear the setting and re-enable DFNs on the report, go to RPMS programmer mode and run the following command:

```
DO DFNCLR^BCCDSTAT
```
4.0 Package Operation

4.1 Manually Generate One CCDA Document

CCDA documents are generated automatically and the process does not require manual intervention. However, on occasion, the site manager may be asked to regenerate the CCDA document for one or more patients. Select the Generate CCD for a single patient (BCCD CCDGENONE) option in the CCDA Menu (BCCD Menu). Enter the patient by name, Social Security Number (SSN), date of birth, or chart number. If multiple patients match the input criteria, select the correct patient from the list displayed as shown in Figure 4-1:

```
Select CCDA Menu Option: one Generate CCD for a single patient
ENTER NAME, SSN, DOB OR CHART#: BCCD, PATIENT
  1 BCCD, PATIENT M 11-07-1911 XXX-XX-0313 ADB 129636
      DB 129637(I)
      URA 129638(I)
  2 BCCD, PATIENT M 08-23-1935 XXX-XX-3756 DB 173511
  3 BCCD, PATIENT M 04-13-1953 XXX-XX-3456 DB 123456
  4 BCCD, PATIENT A M 01-01-1977 XXX-XX-8262 ADB 137983
      DB 137984
      URA 137985
  5 BCCD, PATIENT F SR M 11-14-1961 XXX-XX-4124 ADB 108953
      DB 108954
      URA 108955

ENTER '^' TO STOP, OR
CHOOSE 1-5:
CHOOSE 1-5: 2
BCCD, PATIENT M 08-23-1935 XXX-XX-3756 DB 173511
CCD request has been scheduled for patient SMITH, JOHN
```

Figure 4-1: Manually regenerating the CCDA document for one patient

**Note:** The Generate CCD for a single patient option only works if the RPMS site is a part of the IHS HIE. Accessing this option at a site that is not a part of the IHS HIE will generate the error message shown in Figure 4-2:

```
Select CCDA Menu Option: one Generate CCD for a single patient
No receiving location is currently defined for CCD documents. Your site needs to be associated with a receiving location in order to be able to upload CCD documents. If Support has provided you with the URL of your receiving location, please consult the CCDA Installation Guide for instructions on how to enter this information into the system.

If Support hasn't provided this URL to your site, then your site is a "pull" site and you will be unable to generate CCDA on demand. This will not affect other CCDA functionality.
```

Figure 4-2: Error message when trying to manually regenerate a CCDA document at a site that is not a part of the IHS HIE
4.2 Manually Regenerate All CCDA Documents

If the RPMS site is a part of the IHS HIE, then, on rare occasions, the Help Desk may ask the Site Manager to regenerate CCDA documents for all patients in the database. When this happens, select the Generate CCD documents for all patients in RPMS (BCCD CCDALLGEN) option in the CCDA Menu (BCCD Menu). The system will check if there is enough disk space for this process and display the warning shown in Figure 4-3:

Select CCDA Menu Option: ALL  Generate CCD documents for all patients in RPMS
Checking free space...

Generation and transmission of CCD documents for all patients may take in excess of 2 days. It may also make extensive use of system resources. Please make sure that your system is not overloaded while this process is running as this may impact system performance.

Generation of all CCD documents will be done during the next CCDA nightly job which is currently scheduled to run at 01:00

Schedule All Patients? Yes// N (No)  Not scheduled

Figure 4-3: Manually Regenerating All CCDA Documents

Enter Y (Yes) or N (No) to confirm regeneration of CCDA documents for all patients.

Accessing this option at a site that is not a part of the IHS HIE will generate the error message shown in Figure 4-4:

Select CCDA Menu Option: ALL  Generate CCD documents for all patients in RPMS
No receiving location is currently defined for CCD documents. Your site needs to be associated with a receiving location in order to be able to upload CCD documents. If Support has provided you with the URL of your receiving location, please consult the CCDA Installation Guide for instructions on how to enter this information into the system.

If Support hasn't provided this URL to your site, then your site is a "pull" site and you will be unable to generate CCDA on demand. This will not affect other CCDA functionality.

Figure 4-4: Error message when trying to manually regenerate all CCDA documents at a site that is not a part of the IHS HIE
4.3 Manually Regenerate CCDA Documents for a Specified Date Range

If the RPMS site is a part of the IHS HIE, then, on rare occasions, the Help Desk may ask the Site Manager to regenerate CCDA documents for all visits updated in a specified date range. When this happens, select the Generate CCD documents for a specified date range (BCCD GEN CCD DATE RANGE) option in the CCDA Menu (BCCD Menu). The application will ask the user for the start date/time and end date/time for the range. The application will then check whether the date range will result in an excessive number of documents generated. If the number of documents would exceed the maximum number allowed, the system will display the message shown in Figure 4-5:

```
Select CCDA Menu <TEST ACCOUNT> Option: DAT Generate CCD documents for a specified date range
This option uploads CCDA documents for a given date/time range. It should be used only when instructed to by the support team.
START DATE/TIME: 1/01/2017@2100 (JAN 01, 2017@21:00)
END DATE/TIME: 11/01/2017@2100 (NOV 01, 2017@21:00)
Identifying visits... The selected date range will result in too many documents.
Please enter a smaller range.
```

Figure 4-5: Generate CCD documents for a specified date range option when selected range is too big

If the number of documents is within the allowed limit, the system will queue the document requests and display the message shown in Figure 4-6:

```
Select CCDA Menu <TEST ACCOUNT> Option: DAT Generate CCD documents for a specified date range
This option uploads CCDA documents for a given date/time range. It should be used only when instructed to by the support team.
START DATE/TIME: 1/01/2017@2100 (JAN 01, 2017@21:00)
END DATE/TIME: 11/01/2017@2100 (NOV 01, 2017@21:00)
Identifying visits...
100 visits queued.
```

Figure 4-6: Generate CCD documents for a specified date range option when selected range is sufficiently small
5.0 CCDA Audit Log

The CCDA application contains the global ^BCCD.Audit.AuditLogD, which is used to store the CCDA audit log. The CCDA Audit Log table contains audit information about specific requests for CCDA data that come from the EHR GUI, nightly uploads to the HIE, and for documents generated for Data Export. Prior to Patch 6 of Version 1, the audit log was stored in the CCDA database. Patch 6 of Version 1 moved the audit log to the RPMS database to make it easier for sites to journal and back up the audit log.

Please refer to the AU - Audit and Accountability section in the CCDA Security Manual for instructions to view the data stored in the CCDA Audit Log.
6.0 Limits on the Data Included in CCDA Documents

CCDA documents should contain “current and relevant historical” summary information about patients, which means only recent clinical data is extracted from RPMS and included in CCDA documents except as noted below.

The five types of CCDA documents are:

- Clinical Summary (CS)
- Transitions of Care (ToC)
- Data Export Summary (DP)
- Health Information Exchange Document (HIE)
- Case Reporting (CR) – Sent to Division of Epidemiology

Table 6-1 describes the time ranges and data limitations used to extract RPMS data for individual CCDA modules.

Table 6-1: CCDA Data Limitations

<table>
<thead>
<tr>
<th>CCDA Module</th>
<th>CCDA Document</th>
<th>Data Extract</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allergies and Intolerances</td>
<td>All</td>
<td>• Extract all Allergies (active and inactive) for the patient.</td>
</tr>
<tr>
<td>CCDA Module</td>
<td>CCDA Document</td>
<td>Data Extract</td>
</tr>
<tr>
<td>--------------------------</td>
<td>---------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Assessment and Plan</td>
<td>All</td>
<td>• For each Problem that is POV for the encounter, extract:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• The Care Plan</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• The Visit Instructions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• The Treatment/Regimen</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• The Request/Consultations</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• The Ordered/Pending Lab Tests,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• The Lab Tests Awaiting Results</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• In addition, extract all the below whether associated with a problem or not:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Ordered/Pending Lab and Radiology Tests</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• All Lab/Radiology Tests awaiting Results</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• All Referrals</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• All Requests/Consultations</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• All Future Appointments</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Finally, extract all Treatment/Regimen not associated with a problem</td>
</tr>
<tr>
<td>Care Team</td>
<td>All</td>
<td>• Extract the Visit Provider, Attending and Admitting providers (Inpatient), and Referring Providers associated with the visit.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract all Specialty Providers associated with the patient.</td>
</tr>
<tr>
<td>Case Reporting</td>
<td>CR only</td>
<td>• This section includes the trigger information that prompted the creation of the Case Reporting document.</td>
</tr>
<tr>
<td>Consultation Notes</td>
<td>All</td>
<td>• The contents of any signed TIU document associated with the visit and defined in the CCDA CLINICAL PARAMETERS option under the CLINICAL NOTE for CONSULTATION</td>
</tr>
<tr>
<td>Discharge Diagnoses</td>
<td>All (Inpatient only)</td>
<td>• For inpatient only, extract the primary of Purpose of Visit</td>
</tr>
<tr>
<td>Discharge Summary Notes</td>
<td>All</td>
<td>• The contents of any signed TIU document associated with the visit and defined in the CCDA CLINICAL PARAMETERS option under the CLINICAL NOTE for DISCHARGE SUMMARY/HOSPITAL COURSE</td>
</tr>
<tr>
<td>CCDA Module</td>
<td>CCDA Document</td>
<td>Data Extract</td>
</tr>
<tr>
<td>----------------------</td>
<td>---------------</td>
<td>-----------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Functional Status</td>
<td>CS</td>
<td>• Not extracted.</td>
</tr>
<tr>
<td>Functional Status</td>
<td>ToC, DP, HIE, CR</td>
<td>• Fall Risk: Extract the most recent Fall Risk entry for the patient from the Examinations file</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Elder Care Questions: Extract the most recent entry for the patient.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Problems: Extract the subset of SNOMED codes determined to be related to functional status (EHR IPL FUNCTIONAL STATUS).</td>
</tr>
<tr>
<td>General Header</td>
<td>All</td>
<td>• Visit Date and Location are visit-specific.</td>
</tr>
<tr>
<td>Goals</td>
<td>All</td>
<td>• Extract all Patient Goals that have a status of Active or Maintaining Goal.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract all active or met Care Plan goals that are associated with the visit.</td>
</tr>
<tr>
<td>Health Concerns</td>
<td>All</td>
<td>• Extract Chronic, Episodic, Environmental/Social History, Personal History/Inactive, and Sub-acute problems.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract any problems used as POV for that Encounter (mark with an asterisk).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract any POVs for that Encounter (mark with an asterisk).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract all Patient Goals that have a status of Active or Maintaining Goal</td>
</tr>
<tr>
<td>History and Physical Notes</td>
<td>All</td>
<td>• The contents of any signed TIU document associated with the visit and defined in the CCDA CLINICAL PARAMETERS option under the CLINICAL NOTE for H&amp;P</td>
</tr>
<tr>
<td>Hospital Course</td>
<td>All (Inpatient only)</td>
<td>• Extract the Hospital Course section (see section 2.2) for the Discharge Summary note associated with the visit (see the Discharge Summary Notes definition above to see how the Discharge Summary notes are determined).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract the Hospital Course section (see section 2.2) for the Discharge Summary Addendums associated with the visit.</td>
</tr>
<tr>
<td>CCDA Module</td>
<td>CCDA Document</td>
<td>Data Extract</td>
</tr>
<tr>
<td>------------------------------</td>
<td>---------------</td>
<td>-------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Hospital Discharge Instructions</td>
<td>CS</td>
<td>Not extracted.</td>
</tr>
<tr>
<td>Hospital Discharge Instructions</td>
<td>ToC, DP, CR, HIE (Inpatient only)</td>
<td>The contents of any signed TIU document associated with the visit and defined in the CCDA CLINICAL PARAMETERS option under the CLINICAL NOTE for DISCHARGE INSTRUCTIONS</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extract Discharge Information Addendums associated with the visit.</td>
</tr>
<tr>
<td>Immunizations</td>
<td>All</td>
<td>Extract all patient refusals and all immunizations, including Immunizations 'due'.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extract only the last two Influenza immunizations.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extract Immunization Refused or Declined by the patient.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extract Immunizations Due.</td>
</tr>
<tr>
<td>Medical Equipment</td>
<td>All</td>
<td>Extract all active or expired Patient Implanted Devices.</td>
</tr>
<tr>
<td>Medications</td>
<td>CS</td>
<td>Extract all Active, Held, Suspended, and Pending Outpatient Prescriptions.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extract all Given During Visit medications are extracted.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extract all Outside Medications.</td>
</tr>
<tr>
<td>Medications</td>
<td>ToC, DP, CR</td>
<td>Same as CS Medication but also include:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extract recently expired (last 30 days) for chronic medications.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Inpatient (Unit Dose, IVs, Pending) that were Started between the Admit and Discharge (or Visit Date for ambulatory).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extract all Historical Medications from past 5 years.</td>
</tr>
<tr>
<td>Medications</td>
<td>HIE</td>
<td>Same as ToC/DP/CR but also include all Historical Medications from past 5 years.</td>
</tr>
<tr>
<td>Mental/Cognitive Status</td>
<td>CS</td>
<td>Not extracted.</td>
</tr>
<tr>
<td>CCDA Module</td>
<td>CCDA Document</td>
<td>Data Extract</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>---------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Mental/Cognitive Status</td>
<td>ToC, DP, HIE, CR</td>
<td>• Mini Mental Status Exam: Extract the most recent entry for the patient from the Measurement file.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Problems: Extract the subset of SNOMED codes determined to be related to cognitive status (EHR IPL COGNITIVE STATUS).</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Patient</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Demographics for the requested patient.</td>
</tr>
<tr>
<td>Problems/Encounter Diagnoses</td>
<td>CS</td>
<td>• Extract Chronic, Episodic, and Sub-acute problems.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• The CCDA Site Parameter ADD SOCIAL ENVIRONMENT PROB allows the sites to determine whether they want to extract social/environmental problems (see Section 3.1).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• The CCDA Site Parameter ADD INACTIVE PERS HISTORY PROB allows the sites to determine whether they want to extract personal history with inactive status problems (see Section 3.1).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract any POVs for that Encounter (mark with an asterisk).</td>
</tr>
<tr>
<td>Problems/Encounter Diagnoses</td>
<td>ToC, DP, HIE, CR</td>
<td>• Extract chronic and sub-acute problems.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract social and environmental problems.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract personal history with inactive status problems.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract any POVs for that Encounter (mark with an asterisk).</td>
</tr>
<tr>
<td>Procedures and Interventions</td>
<td>CS</td>
<td>• Extract the procedures done on today’s visit (V Procedure and V CPT).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract Radiology exams associated with the visit.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Procedure Note: The contents of any signed TIU document associated with the visit and defined in the CCDA CLINICAL PARAMETERS option under the CLINICAL NOTE for PROCEDURE notes.</td>
</tr>
<tr>
<td>CCDA Module</td>
<td>CCDA Document</td>
<td>Data Extract</td>
</tr>
<tr>
<td>------------------------------</td>
<td>--------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| Procedures and Interventions | ToC, DP, HIE, CR | • Extract all significant procedures for CPT codes entered in the APCH HS MAJOR PROCEDURES CPTS taxonomy. For ICD-9/ICD-10 codes, use the same ‘significant procedure’ logic as is used by the Health Summary.  
• Extract all Radiology exams for the patient. |
| Progress Notes               | All          | • Any signed TIU document associated with the visit and not defined in the CCDA CLINICAL PARAMETERS option under the CLINICAL NOTE definitions and not specifically excluded in the SITE NOTE EXCLUSION parameter |
| Reason for Referral          | CS           | • Not extracted.                                                             |
| Reason for Referral          | ToC, DP, HIE, CR | • Extract referral information for the referrals associated with the visits. |
| Reason for Visit/Hospitalization | CS, ToC     | • Reason for Visit (Ambulatory), extract the Chief complaint.  
• Reason for Hospitalization (inpatient) get the Admit Diagnosis associated with the inpatient stay |
| Reason for Visit/Hospitalization | DP, CR      | • Not extracted.                                                             |
| Recent Encounters            | All          | • Extract the visit for which the document is being created  
• Extract all Hospitalization, Observation, Nursing Home, Same Day Surgery encounters going back one year from the date of the visit for which the document is being created.  
• Extract all Ambulatory and TeleMedicine encounters that are workload reportable going back one year going back one year from the date of the visit for which the document is being created.  
• For each encounter, extract the Purpose of Visit for the encounter. |
<table>
<thead>
<tr>
<th>CCDA Module</th>
<th>CCDA Document</th>
<th>Data Extract</th>
</tr>
</thead>
</table>
| Recent Test Results | CS            | - Extract lab results that are modified or resulted. For ambulatory, extract labs resulted with the past 15 day for the date of the visit. For inpatient stays, extract all labs resulted during the inpatient stay.  
- Extract Radiology results for the current visit that has an impression, and the report has been verified. For inpatient stay, only get the results of the first and last Radiology exam.  
- Extract Radiology Results with abnormal results that are within thirteen months of the earliest visit date.  
- Does not extract skin tests. |
| Recent Test Results | ToC, DP, CR    | - Extract resulted tests from Laboratory. For Ambulatory, results are not visit-specific. Extract the last two results for each atomic test and each panel that occurred within the last 30 days. For Inpatient, results are visit-specific. Extract the initial and last resulted test associated with the inpatient stay.  
- Extract Radiology results for the current visit that has an impression, and the report has been verified. For inpatient stay, only get the results of the first and last Radiology exam.  
- Extract Radiology Results with abnormal results that are within thirteen months of the earliest visit date.  
- Extract skin tests that occurred in the previous year, except for the TB skin test that has no time limit. |
<table>
<thead>
<tr>
<th>CCDA Module</th>
<th>CCDA Document</th>
<th>Data Extract</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recent Test Results</td>
<td>HIE</td>
<td>• Extract resulted tests from Laboratory. For Ambulatory, results are not visit-specific. Extract the last five results for each atomic test and each panel that occurred within the last year. For Inpatient, results are visit-specific. Extract the initial and last three resulted test associated with the inpatient stay.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract Radiology results for the current visit that has an impression, and the report has been verified. For inpatient stay, only get the results of the first and last Radiology exam.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract Radiology Results with abnormal results that are within thirteen months of the earliest visit date.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract skin tests that occurred in the previous year, except for the TB skin test that has no time limit.</td>
</tr>
<tr>
<td>Recent Vital Signs</td>
<td>CS</td>
<td>• Get only the most recent three vital signs per type. Display only Height, Weight, Blood Pressure, and Body Mass Index (BMI).</td>
</tr>
<tr>
<td>Recent Vital Signs</td>
<td>ToC, DP, HIE</td>
<td>• Extract measurements using the CCDA list of allowable measurements - Body Mass Index (BMI), Blood Pressure (BP), Head Circumference (HC), Height (HT), O2 Saturation (O2), Pulse (PU), Respiratory Rate (RS), Temperature (TMP), and Weight (WT).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• For O2 Saturation measurement, also extract the Inhaled Oxygen Concentration, Inhaled Flow Rate, and Method modifiers.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Extract Pain measurement, even though it is not specified by CCDA.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Ambulatory vital signs are not visit-specific. Extract the last three measurements.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Inpatient vital signs are visit-specific. Extract the initial and last three associated with the inpatient stay.</td>
</tr>
<tr>
<td>CCDA Module</td>
<td>CCDA Document</td>
<td>Data Extract</td>
</tr>
<tr>
<td>------------------------------</td>
<td>---------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Security and Privacy</td>
<td>CS and ToC only</td>
<td>• Standard text - “This information, except for Alcohol and Drug Abuse Patient Records as defined in 42 Code of Federal Regulations, Part 2, may be subject to redisclosure by the recipient and may no longer be protected by the Health Insurance Portability and Accountability Act Privacy Rule of 1996 and the Privacy Act of 1974, as amended.”</td>
</tr>
<tr>
<td>Prohibitions</td>
<td>All</td>
<td>• Get the most recent smoking status from V Health Factors.</td>
</tr>
</tbody>
</table>
7.0 Manage E-Mail Notifications

This step allows the user to configure the CCDA Ensemble production to automatically send e-mail notifications about new errors to selected users. A notification is typically sent as soon as a new error occurs, but the number of e-mails is limited to one per 15 minutes in order to prevent flooding the recipients’ mailboxes with multiple errors. This e-mail notification functionality is only available on systems that have an e-mail server configured. This managing e-mail notifications step is optional and may be skipped if the sites do not want to be notified about CCDA errors by e-mail. For requests coming from the EHR GUI, the site manager will be able to view errors in the CCDA Audit Log, which is described in the CCDA Security Manual, even if e-mail notifications are not set up.

Check if the site’s e-mail server requires a user to sign on. If it does require a user to sign on, the credentials will need to be set up first. To set up the user credentials, sign on to HealthShare’s Management Portal as an administrator and follow these instructions:

1. Switch to the CCDA namespace.

2. In the Namespace Chooser box, choose the appropriate CCDA namespace (the namespace will consist of “CCDA” concatenated with the name of the site’s RPMS namespace).

3. On the Management Portal screen, click Ensemble, then click Configure >>, then click Credentials.

Figure 7-1: Management Portal, Credential Options
Figure 7-2 shows the **Credentials Viewer** screen.

4. In the **ID** field enter an arbitrary ID that will identify the site’s e-mail server, e.g., `mail-server` (this ID will later be utilized in the **Credentials** field of the e-mail notification screen).

5. In the **Username** field, the HealthShare username will be defaulted; however, change the HealthShare username to a valid username on the e-mail server that will allow the user to sign on to the server and send e-mails.

6. In the **Password** field, enter the password for the username that was entered in the previous step.

7. Leave the **Business Partner** field blank.

8. Click **Save** (the newly entered credential will be displayed in the list in the center of the screen).

9. Once the credentials have been configured (or if the site’s e-mail server does not require authentication), sign onto HealthShare’s Management Portal as the administrator.

10. Switch to the CCDA namespace. The namespace will consist of “CCDA” concatenated with the name of the site’s RPMS namespace.

11. On the **Management Portal** page, click **Ensemble**, then click **Configure >>**, then click **Production**.
12. On the Production Configuration screen, click Open.

13. In the Finder Dialog, click BCCD, then click Prod, then click Production to select the CCDA production (the Production Configuration screen will update to display the items for the CCDA production).
14. Single-click on EmailAlert in the Operations column (the details of this process will be displayed in the Production Settings box on the right side of the page).

15. In the EmailAlert box on the right-hand side, be sure the Basic Settings and Additional Settings sections are expanded, and then enter values in Table 7-1:

Table 7-1: EmailAlert settings

<table>
<thead>
<tr>
<th>Field Name</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Enabled</td>
<td>Check the box to enable email alerts</td>
</tr>
<tr>
<td>SMTP Server</td>
<td>IP address or name of the email server at the site</td>
</tr>
<tr>
<td>SMTP Port</td>
<td>Port number used by the site’s email server. The default is 25</td>
</tr>
<tr>
<td>Credentials</td>
<td>Only required if the e-mail server requires authentication (see instructions above)</td>
</tr>
<tr>
<td>Recipient</td>
<td>A comma-delimited list of email addresses that Ensemble will be sending alerts to, e.g., <a href="mailto:John.Doe@ihs.gov">John.Doe@ihs.gov</a>, <a href="mailto:Jane.Doe@ihs.gov">Jane.Doe@ihs.gov</a></td>
</tr>
<tr>
<td>From</td>
<td>The e-mail address that the alerts will appear as coming from, e.g., <a href="mailto:CCDA@sitename.ihs.gov">CCDA@sitename.ihs.gov</a></td>
</tr>
</tbody>
</table>

If the site has a functional e-mail server but the user does not have some of the information in the EmailAlert settings table above, contact the Help Desk. **Do not modify any other values on this screen since it can invalidate the Ensemble production.**
16. Once all the required data is entered, click **Apply** at the top of the **EmailAlert** box.

Figure 7-6 contains a sample screenshot which shows what the screen will look like during the configuration process.

![Figure 7-6: Ensemble e-mail server configuration](image-url)
8.0 References and Sources


Office of the National Coordinator for Health Information Technology (ONC) 2015 Certification of Health Information Technology (CHIT): https://www.healthit.gov/topic/certification-ehrs/certification-health-it
Appendix A  Rules of Behavior

The Resource and Patient Management (RPMS) system is a United States Department of Health and Human Services (HHS), Indian Health Service (IHS) information system that is FOR OFFICIAL USE ONLY. The RPMS system is subject to monitoring; therefore, no expectation of privacy shall be assumed. Individuals found performing unauthorized activities are subject to disciplinary action including criminal prosecution.

All users (Contractors and IHS Employees) of RPMS will be provided a copy of the Rules of Behavior (RoB) and must acknowledge that they have received and read them prior to being granted access to a RPMS system, in accordance IHS policy.

- For a listing of general ROB for all users, see the most recent edition of IHS General User Security Handbook (SOP 06-11a).
- For a listing of system administrators/managers rules, see the most recent edition of the IHS Technical and Managerial Handbook (SOP 06-11b).

Both documents are available at this IHS website: https://home.ihs.gov/security/index.cfm.

**Note:** Users must be logged on to the IHS D1 Intranet to access these documents.

The ROB listed in the following sections are specific to RPMS.

A.1  All RPMS Users

In addition to these rules, each application may include additional RoBs that may be defined within the documentation of that application (e.g., Dental, Pharmacy).

A.1.1  Access

RPMS users shall:

- Only use data for which you have been granted authorization.
- Only give information to personnel who have access authority and have a need to know.
- Always verify a caller’s identification and job purpose with your supervisor or the entity provided as employer before providing any type of information system access, sensitive information, or nonpublic agency information.
- Be aware that personal use of information resources is authorized on a limited basis within the provisions Indian Health Manual Part 8, “Information Resources Management,” Chapter 6, “Limited Personal Use of Information Technology Resources.”
RPMS users shall not:

- Retrieve information for someone who does not have authority to access the information.
- Access, research, or change any user account, file, directory, table, or record not required to perform their official duties.
- Store sensitive files on a PC hard drive, or portable devices or media, if access to the PC or files cannot be physically or technically limited.
- Exceed their authorized access limits in RPMS by changing information or searching databases beyond the responsibilities of their jobs or by divulging information to anyone not authorized to know that information.

A.1.2 Information Accessibility

RPMS shall restrict access to information based on the type and identity of the user. However, regardless of the type of user, access shall be restricted to the minimum level necessary to perform the job.

RPMS users shall:

- Access only those documents they created and those other documents to which they have a valid need-to-know and to which they have specifically granted access through an RPMS application based on their menus (job roles), keys, and FileMan access codes. Some users may be afforded additional privileges based on the functions they perform, such as system administrator or application administrator.
- Acquire a written preauthorization in accordance with IHS policies and procedures prior to interconnection to or transferring data from RPMS.

A.1.3 Accountability

RPMS users shall:

- Behave in an ethical, technically proficient, informed, and trustworthy manner.
- Log out of the system whenever they leave the vicinity of their personal computers (PCs).
- Be alert to threats and vulnerabilities in the security of the system.
- Report all security incidents to their local Information System Security Officer (ISSO)
- Differentiate tasks and functions to ensure that no one person has sole access to or control over important resources.
- Protect all sensitive data entrusted to them as part of their government employment.
• Abide by all Department and Agency policies and procedures and guidelines related to ethics, conduct, behavior, and information technology (IT) information processes.

A.1.4 Confidentiality

RPMS users shall:

• Be aware of the sensitivity of electronic and hard copy information and protect it accordingly.

• Store hard copy reports/storage media containing confidential information in a locked room or cabinet.

• Erase sensitive data on storage media prior to reusing or disposing of the media.

• Protect all RPMS terminals from public viewing at all times.

• Abide by all Health Insurance Portability and Accountability Act (HIPAA) regulations to ensure patient confidentiality.

RPMS users shall not:

• Allow confidential information to remain on the PC screen when someone who is not authorized to that data is in the vicinity.

• Store sensitive files on a portable device or media without encrypting.

A.1.5 Integrity

RPMS users shall:

• Protect their systems against viruses and similar malicious programs.

• Observe all software license agreements.

• Follow industry standard procedures for maintaining and managing RPMS hardware, operating system software, application software, and/or database software and database tables.

• Comply with all copyright regulations and license agreements associated with RPMS software.

RPMS users shall not:

• Violate federal copyright laws.

• Install or use unauthorized software within the system libraries or folders.

• Use freeware, shareware, or public domain software on/with the system without their manager’s written permission and without scanning it for viruses first.
A.1.6 System Logon
RPMS users shall:

• Have a unique User Identification/Account name and password.
• Be granted access based on authenticating the account name and password entered.
• Be locked out of an account after five successive failed login attempts within a specified time period (e.g., one hour).

A.1.7 Passwords
RPMS users shall:

• Change passwords a minimum of every 90 days.
• Create passwords with a minimum of eight characters.
• If the system allows, use a combination of alpha-numeric characters for passwords, with at least one uppercase letter, one lower case letter, and one number. It is recommended, if possible, that a special character also be used in the password.
• Change vendor-supplied passwords immediately.
• Protect passwords by committing them to memory or store them in a safe place (do not store passwords in login scripts or batch files).
• Change passwords immediately if password has been seen, guessed, or otherwise compromised, and report the compromise or suspected compromise to their ISSO.
• Keep user identifications (IDs) and passwords confidential.

RPMS users shall not:

• Use common words found in any dictionary as a password.
• Use obvious readable passwords or passwords that incorporate personal data elements (e.g., user’s name, date of birth, address, telephone number, or social security number; names of children or spouses; favorite band, sports team, or automobile; or other personal attributes).
• Share passwords/IDs with anyone or accept the use of another’s password/ID, even if offered.
• Reuse passwords. A new password must contain no more than five characters per eight characters from the previous password.
• Post passwords.
• Keep a password list in an obvious place, such as under keyboards, in desk drawers, or in any other location where it might be disclosed.
• Give a password out over the phone.

A.1.8 Backups
RPMS users shall:
• Plan for contingencies such as physical disasters, loss of processing, and disclosure of information by preparing alternate work strategies and system recovery mechanisms.
• Make backups of systems and files on a regular, defined basis.
• If possible, store backups away from the system in a secure environment.

A.1.9 Reporting
RPMS users shall:
• Contact and inform their ISSO that they have identified an IT security incident and begin the reporting process by providing an IT Incident Reporting Form regarding this incident.
• Report security incidents as detailed in the IHS Incident Handling Guide (SOP 05-03).
RPMS users shall not:
• Assume that someone else has already reported an incident. The risk of an incident going unreported far outweighs the possibility that an incident gets reported more than once.

A.1.10 Session Timeouts
RPMS system implements system-based timeouts that back users out of a prompt after no more than 5 minutes of inactivity.
RPMS users shall:
• Utilize a screen saver with password protection set to suspend operations at no greater than 10 minutes of inactivity. This will prevent inappropriate access and viewing of any material displayed on the screen after some period of inactivity.

A.1.11 Hardware
RPMS users shall:
• Avoid placing system equipment near obvious environmental hazards (e.g., water pipes).
• Keep an inventory of all system equipment.
• Keep records of maintenance/repairs performed on system equipment.
RPMS users shall not
• Eat or drink near system equipment.

A.1.12 Awareness
RPMS users shall:
• Participate in organization-wide security training as required.
• Read and adhere to security information pertaining to system hardware and software.
• Take the annual information security awareness.
• Read all applicable RPMS manuals for the applications used in their jobs.

A.1.13 Remote Access
Each subscriber organization establishes its own policies for determining which employees may work at home or in other remote workplace locations. Any remote work arrangement should include policies that:
• Are in writing.
• Provide authentication of the remote user through the use of ID and password or other acceptable technical means.
• Outline the work requirements and the security safeguards and procedures the employee is expected to follow.
• Ensure adequate storage of files, removal, and nonrecovery of temporary files created in processing sensitive data, virus protection, and intrusion detection, and provide physical security for government equipment and sensitive data.
• Establish mechanisms to back up data created and/or stored at alternate work locations.
Remote RPMS users shall:
• Remotely access RPMS through a virtual private network (VPN) whenever possible. Use of direct dial in access must be justified and approved in writing and its use secured in accordance with industry best practices or government procedures.
Remote RPMS users shall not:
• Disable any encryption established for network, internet, and Web browser communications.
A.2 RPMS Developers

RPMS developers shall:

- Always be mindful of protecting the confidentiality, availability, and integrity of RPMS when writing or revising code.
- Always follow the IHS RPMS Programming Standards and Conventions (SAC) when developing for RPMS.
- Only access information or code within the namespaces for which they have been assigned as part of their duties.
- Remember that all RPMS code is the property of the U.S. Government, not the developer.
- Not access live production systems without obtaining appropriate written access and shall only retain that access for the shortest period possible to accomplish the task that requires the access.
- Observe separation of duties policies and procedures to the fullest extent possible.
- Document or comment all changes to any RPMS software at the time the change or update is made. Documentation shall include the programmer’s initials, date of change, and reason for the change.
- Use checksums or other integrity mechanism when releasing their certified applications to assure the integrity of the routines within their RPMS applications.
- Follow industry best standards for systems they are assigned to develop or maintain and abide by all Department and Agency policies and procedures.
- Document and implement security processes whenever available.

RPMS developers shall not:

- Write any code that adversely impacts RPMS, such as backdoor access, “Easter eggs,” time bombs, or any other malicious code or make inappropriate comments within the code, manuals, or help frames.
- Grant any user or system administrator access to RPMS unless proper documentation is provided.
- Release any sensitive agency or patient information.

A.3 Privileged Users

Personnel who have significant access to processes and data in RPMS, such as, system security administrators, systems administrators, and database administrators, have added responsibilities to ensure the secure operation of RPMS.
Privileged RPMS users shall:

- Verify that any user requesting access to any RPMS system has completed the appropriate access request forms.
- Ensure that government personnel and contractor personnel understand and comply with license requirements. End users, supervisors, and functional managers are ultimately responsible for this compliance.
- Advise the system owner on matters concerning information technology security.
- Assist the system owner in developing security plans, risk assessments, and supporting documentation for the certification and accreditation process.
- Ensure that any changes to RPMS that affect contingency and disaster recovery plans are conveyed to the person responsible for maintaining continuity of operations plans.
- Ensure that adequate physical and administrative safeguards are operational within their areas of responsibility and that access to information and data is restricted to authorized personnel on a need-to-know basis.
- Verify that users have received appropriate security training before allowing access to RPMS.
- Implement applicable security access procedures and mechanisms, incorporate appropriate levels of system auditing, and review audit logs.
- Document and investigate known or suspected security incidents or violations and report them to the ISSO, Chief Information Security Officer (CISO), and systems owner.
- Protect the supervisor, superuser, or system administrator passwords.
- Avoid instances where the same individual has responsibility for several functions (i.e., transaction entry and transaction approval).
- Watch for unscheduled, unusual, and unauthorized programs.
- Help train system users on the appropriate use and security of the system.
- Establish protective controls to ensure the accountability, integrity, confidentiality, and availability of the system.
- Replace passwords when a compromise is suspected. Delete user accounts as quickly as possible from the time that the user is no longer authorized system. Passwords forgotten by their owner should be replaced, not reissued.
- Terminate user accounts when a user transfers or has been terminated. If the user has authority to grant authorizations to others, review these other authorizations. Retrieve any devices used to gain access to the system or equipment. Cancel logon IDs and passwords and delete or reassign related active and backup files.
• Use a suspend program to prevent an unauthorized user from logging on with the current user's ID if the system is left on and unattended.

• Verify the identity of the user when resetting passwords. This can be done either in person or having the user answer a question that can be compared to one in the administrator’s database.

• Shall follow industry best standards for systems they are assigned to and abide by all Department and Agency policies and procedures.

Privileged RPMS users shall not:

• Access any files, records, systems, etc., that are not explicitly needed to perform their duties

• Grant any user or system administrator access to RPMS unless proper documentation is provided.

• Release any sensitive agency or patient information.
Glossary

**BCCD**
Namespace for CCDA files, routines and classes.

**BMW**
Namespace for classes generated from FileMan files.

**Hypertext Transfer Protocol**
A widely used communication protocol on the World Wide Web.

**Simple Object Access Protocol**
The communication protocol and message format used in communicating with web services.

**Software Quality Assurance**
The office within the IHS Office of Information Technology responsible for ensuring that the system conforms to RPMS Programming Standards and Conventions.

**V Codes**
V codes (codes V01–V91) are used to describe encounters with circumstances other than disease or injury. V codes are used either as a first listed (primary) or contributing (secondary) code depending on the situation.

**Web service**
An application programming interface (API) that allows communication with an application via SOAP messages over HTTP.

**Web Service Description Language**
A file that describes a web service’s API.
## Acronym List

<table>
<thead>
<tr>
<th>Acronym</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>API</td>
<td>Application Programmer Interface</td>
</tr>
<tr>
<td>BMI</td>
<td>Body Mass Index</td>
</tr>
<tr>
<td>CCD</td>
<td>Continuity of Care Document</td>
</tr>
<tr>
<td>CCDA</td>
<td>Consolidated Clinical Document Architecture</td>
</tr>
<tr>
<td>CDA R2</td>
<td>Clinical Document Architecture Release 2</td>
</tr>
<tr>
<td>CHIT</td>
<td>Certified Health Information Technology</td>
</tr>
<tr>
<td>CPT</td>
<td>Current Procedural Terminology</td>
</tr>
<tr>
<td>CS</td>
<td>Clinical Summary</td>
</tr>
<tr>
<td>DP</td>
<td>Data Export, formerly Data Portability</td>
</tr>
<tr>
<td>DTL</td>
<td>Data Transformation Language</td>
</tr>
<tr>
<td>EHR</td>
<td>Electronic Health Record</td>
</tr>
<tr>
<td>GUI</td>
<td>Graphical User Interface</td>
</tr>
<tr>
<td>HIE</td>
<td>Health Information Exchange</td>
</tr>
<tr>
<td>HIPAA</td>
<td>Health Insurance Portability and Accountability Act</td>
</tr>
<tr>
<td>HL7</td>
<td>Health Level Seven</td>
</tr>
<tr>
<td>HTTP</td>
<td>Hypertext Transfer Protocol</td>
</tr>
<tr>
<td>I/T/U</td>
<td>IHS direct, tribal, and urban facilities. Generally, refers to all components of the Indian healthcare system</td>
</tr>
<tr>
<td>IHS</td>
<td>Indian Health Service</td>
</tr>
<tr>
<td>KIDS</td>
<td>Kernel Installation and Distribution System</td>
</tr>
<tr>
<td>OIT</td>
<td>Office of Information Technology</td>
</tr>
<tr>
<td>PHR</td>
<td>Personal Health Record</td>
</tr>
<tr>
<td>POV</td>
<td>Purpose of Visit</td>
</tr>
<tr>
<td>RPMS</td>
<td>Resource and Patient Management System</td>
</tr>
<tr>
<td>SAC</td>
<td>Standards and Conventions</td>
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<tr>
<td>SME</td>
<td>Subject Matter Expert</td>
</tr>
<tr>
<td>SNOMED CT</td>
<td>Systematized Nomenclature of Medicine--Clinical Terms</td>
</tr>
<tr>
<td>TB</td>
<td>Tuberculosis</td>
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<tr>
<td>ToC</td>
<td>Transitions of Care</td>
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<td>WS</td>
<td>Web Service</td>
</tr>
<tr>
<td>WSDL</td>
<td>Web Services Description Language</td>
</tr>
<tr>
<td>XML</td>
<td>Extensible Markup Language</td>
</tr>
</tbody>
</table>
Contact Information

If you have any questions or comments regarding this distribution, please contact the IHS IT Service Desk.

**Phone:** (888) 830-7280 (toll free)
**Web:** [https://www.ihs.gov/itsupport/](https://www.ihs.gov/itsupport/)
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