# Common – Clinical Document Release Note v1.1.3

18 August 2014 Approved for external information

## EP-1754:2014 Common - Clinical Document v1.1.3

## **Release rationale**

This incremental release of the Common – Clinical Documents end product introduces the *Template Package Directory*.

The purpose of this new product component is to provide implementers of clinical document specifications and other stakeholders with a comprehensive overview of available template packages and their current status.

The template package directory also contains hyperlinks referring to the download locations of all template packages. For each template package, the hyperlink refers to the location of the template package library containing the template package. Template package libraries are published for each clinical document type and are contained in the end product for the document type.

## **Package inclusions**

#### New

Identifier	Name
NEHTA-1738:2014	Common – Clinical Document – Template Package Directory v1.0

### Updated (supersedes previous version)

Identifier	Name	
NEHTA-1758:2014	Common – Clinical Document - Release Note v1.1.3	
No change		
Identifier	Name	
NEHTA-1199:2012	Clinical Documentation - CDA Rendering Specification v1.0	
NEHTA-1446:2013	Clinical Documents - Common Conformance Profile v1.4	
NEHTA-1165:2012	Common - CDA - FAQ MIMS Coding Requirements v1.0	
NEHTA-1096:2011	Common - CDA - FAQ OIDs For HL7 v1.0	
NEHTA-1097:2011	Common - CDA - Implementation Guidance - Representing Coding in CDA Documents	

Identifier	Name	
NEHTA-1331:2012	<i>Common - Clinical Document - Conformance Test Specification for CDA Packaging v1.4</i>	
NEHTA-1330:2012	<i>Common - Clinical Document - Conformance Test Specification for CDA Rendering v1.3</i>	
NEHTA-1329:2012	<i>Common - Clinical Document - Conformance Test Specification for Clinical Documents v1.2</i>	
NEHTA-1270:2013	Common - Clinical Document - FAQ - Clarification on Messaging and CDA Packaging v1.4	
NEHTA-1255:2012	Common - Clinical Document - FAQ Appropriate use of date and date-time values in Clinical Documents v1.0	
NEHTA-1276:2013	Common - Clinical Document - FAQ Hash value verification v1.0	
NEHTA-1275:2013	Common - Clinical Document - FAQ Pathology Date Time v1.1	
NEHTA-1277:2013	Common - Clinical Document - FAQ Patient Medications v1.1	
NEHTA-1271:2013	Common - Clinical Document - FAQ Rendering Specification v2.0	
NEHTA-1278:2013	Common - Clinical Document - FAQ Representing patient IDs in CDA documents v1.1	
NEHTA-1256:2012	Common - Clinical Document - FAQ Trusted source on a CDA Package v1.0	
NEHTA-1564:2014	Common – Clinical Document – PCEHR Usability Recommendations v1.1	
NEHTA-1328:2013	<i>Common - Clinical Document - Supplementary Notes for Implementers Relating to Clinical Document Presentation v1.0</i>	

## Scope

The scope of this end product has not changed as part of this release.

## Stakeholders

Stakeholder groups involved in the development of this release include:

- Commonwealth Department of Health
- Australian College of Rural and Remote Medicine (ACRRM)
- Australian Medical Association (AMA)
- Australian Medicare Local Alliance (AMLA)
- Aboriginal Medical Services Alliance Northern Territory (AMSANT)
- Australian Primary Health Care Nurses Association (APNA)
- Improvement Foundation (Australia) (IF)
- Royal Australian College of General Practitioners (RACGP)

### Audience

- Implementers of clinical systems producing or consuming clinical documents
- Senior managers and policy makers, clinical experts, health information managers, IT operations and support teams, and system integrators
- Technical and non-technical readers

## **Known issues**

None known

## Support

For further support or to provide feedback, please email <u>help@nehta.gov.au</u>.

## **Future releases**

Increased uptake and implementation of the specifications provided as part of this end product are expected to result in the need to further update and improve these specifications. Any such updates will be managed through the joint change control process operated by the Commonwealth Department of Health.

In addition to changes managed through the joint change control process, NEHTA may provide supplementary implementation guidance for the specifications of this end product. Such information will be added to the end product as additional or updated product components and published as an incremental release of the end product (version identifier 1.1.x).

## Previous releases

## EP-1589:2014 Common - Clinical Document v1.1.2

Release note: NEHTA-1592:2014, 5 May 2014

## **Release rationale**

This incremental release of the Common - Clinical Document end product introduces *Clinical Documents – PCEHR Usability Recommendations*. This new product component replaces the *Supplementary Guidance for Implementers* product component introduced with the previous version of this end product.

The *PCEHR Usability Recommendations* document contains implementation guidance previously published in *Supplementary Guidance for Implementers*. The new format, "usability recommendations", makes it easier for implementers to assess whether their software conforms to the guidance.

PCEHR *usability recommendations* are not part of PCEHR *conformance requirements*. Only the latter are used as the basis for conformance assessments performed as a prerequisite to PCEHR system integration. PCEHR usability recommendations can be used by implementers to perform usability assessments on a voluntary basis, for example, with the aim of providing their users with a consistently high level of usability.

The *PCEHR Usability Recommendations* document also contains additional guidance for implementers, developed as part of NEHTA's Clinical Usability Program (CUP) Release 2.

Please refer to the Capabilities section on page 7 for more details.

This release also removes the developer resource product components and related product data sheets. These have been republished in two new end products:

- <u>Clinical Documents Integration Toolkit v1.0</u>
- Secure Messaging Integration Toolkit v1.0

In addition, the document *Reference Platform - Vendor End 2 End Portal v1.4* has been archived as it is no longer relevant. *eSignature - Java Library v1.1.0* has also been archived since it is included in the libraries in the new integration toolkits

## Package inclusions

### New

Identifier	Name
NEHTA-1564:2014	Common – Clinical Documents – PCEHR Usability Recommendations v1.1

### Updated (supersedes previous version)

Identifier	Name
NEHTA-1592:2014	Common – Clinical Document - Release Note v1.1.2

### Archived

Identifier	Name
NEHTA-1476:2013	Clinical Documents - Supplementary Guidance for Implementers v1.0

Identifier	Name	
NEHTA-1217:2012	Reference Platform - Vendor End 2 End Portal v1.4	
NEHTA-1288:2013	eSignature - Java Library v1.1.0	
No change		
Identifier	Name	
NEHTA-1199:2012	Clinical Documentation - CDA Rendering Specification v1.0	
NEHTA-1446:2013	Clinical Documents - Common Conformance Profile v1.4	
NEHTA-1165:2012	Common - CDA - FAQ MIMS Coding Requirements v1.0	
NEHTA-1096:2011	Common - CDA - FAQ OIDs For HL7 v1.0	
NEHTA-1097:2011	<i>Common - CDA - Implementation Guidance - Representing Coding in CDA Documents v1.0</i>	
NEHTA-1229:2011	Common - Clinical Document - CDA Package v1.0	
NEHTA-1226:2011	Common - Clinical Document - Clinical Package v1.0	
NEHTA-1331:2012	<i>Common - Clinical Document - Conformance Test Specification for CDA Packaging v1.4</i>	
NEHTA-1330:2012	<i>Common - Clinical Document - Conformance Test Specification for CDA Rendering v1.3</i>	
NEHTA-1329:2012	<i>Common - Clinical Document - Conformance Test Specification for Clinical Documents</i> v1.2	
NEHTA-1270:2013	Common - Clinical Document - FAQ - Clarification on Messaging and CDA Packaging v1.4	
NEHTA-1255:2012	Common - Clinical Document - FAQ Appropriate use of date and date-time values in Clinical Documents v1.0	
NEHTA-1276:2013	Common - Clinical Document - FAQ Hash value verification v1.0	
NEHTA-1275:2013	Common - Clinical Document - FAQ Pathology Date Time v1.1	
NEHTA-1277:2013	Common - Clinical Document - FAQ Patient Medications v1.1	
NEHTA-1271:2013	Common - Clinical Document - FAQ Rendering Specification v2.0	
NEHTA-1278:2013	Common - Clinical Document - FAQ Representing patient IDs in CDA documents v1.1	
NEHTA-1256:2012	Common - Clinical Document - FAQ Trusted source on a CDA Package v1.0	
NEHTA-1328:2013	<i>Common - Clinical Document - Supplementary Notes for Implementers Relating to Clinical Document Presentation v1.0</i>	

## Moved to new end products

### Republished in Clinical Documents Integration Toolkit v1.0

The following product components have been removed from this end product and republished under the identifier shown.

Superseded and republished

Identifier	Name and version
NEHTA-1601:2014	CDA Document Library - Product Data Sheet v1.1
NEHTA-1605:2014	CDA Packaging Library - Product Data Sheet v1.1
NEHTA-1606:2014	Generic CDA Style Sheet - Product Data Sheet v1.1
NEHTA-1598:2014	Australian CDA Schema Extension 3.0 - Compilable CDA Schema v20140219
NEHTA-1599:2014	Australian CDA Schema Extension 3.0 - CDA Schema v20140219
NEHTA-1600:2014	CDA Document LibraryNET Sample Code v4.0.1
NEHTA-1603:2014	CDA Packaging LibraryNET Sample Code v1.4.0

Republished (same version, new identifier)

New identifier	Name and version	Old identifier
NEHTA-1602:2013	CDA Packaging Library - Java Sample Code v1.2.2	NEHTA-1285:2013
NEHTA-1608:2013	Generic CDA Style Sheet - Stylesheet With CSS v1.2.8	NEHTA-1496:2013
NEHTA-1607:2013	Generic CDA Style Sheet - Stylesheet No CSS v1.2.8	NEHTA-1497:2013

### Republished in Secure Messaging Integration Toolkit v1.0

The following product components have been removed from this end product and republished under the identifier shown.

Superseded and republished

Identifier	Name and version
NEHTA-1627:2014	HL7 MDM Library - Product Data Sheet v1.1

Republished (same version, new identifier)

New identifier	identifier Name and version Old identif	
NEHTA-1626:2013	HL7 MDM Library - Java Sample Code v1.1.4	NEHTA-1286:2013
NEHTA-1625:2013	HL7 MDM LibraryNET Sample Code v1.0.6	NEHTA-1287:2013

### Scope

The scope of the Common Clinical Document end product has not changed as part of this release.

## Stakeholders

Stakeholder groups involved in the development of this release include:

- Commonwealth Department of Health
- Australian College of Rural and Remote Medicine (ACRRM)

- Australian Medical Association (AMA)
- Australian Medicare Local Alliance (AMLA)
- Aboriginal Medical Services Alliance Northern Territory (AMSANT)
- Australian Primary Health Care Nurses Association (APNA)
- Improvement Foundation (Australia) (IF)
- Royal Australian College of General Practitioners (RACGP)

### Audience

- Implementers of clinical systems producing or consuming clinical documents
- Senior managers and policy makers, clinical experts, health information managers, IT operations and support teams, and system integrators
- Technical and non-technical readers

### Known issues

None known

### Capabilities

### **CUP Release 2**

CUP Release 2 has been focused on resolving key usability issues with clinical information systems used by general practitioners. The usability recommendations provided as part of this release represents the outcomes of several workshops with clinical consultation groups.

Information related to all types of clinical documents is documented in the new product component *Clinical Documents – PCEHR Usability Recommendations v1.1*, which replaces the document created for CUP Release 1 (*Clinical Documents – Supplementary Guidance for Implementers v1.0*).

Note that additional usability recommendations resulting from CUP Release 2 have been released as updates to other end products, namely *Event Summary* v1.3.1 and *Shared Health Summary* v1.4.3.

## EP-1477:2013 Common - Clinical Document v1.1.1

Release note: NEHTA-1480:2013, 24 October 2013

## **Release rationale**

This incremental release of the Common Clinical Document end product introduces supplementary guidance for the implementation of clinical documents, representing a key outcome of NEHTA's Clinical Usability Program (CUP) Release 1.

Aligning with the sets of template packages supported by PCEHR Releases 3.5 and 4, updated versions of the CDA Document Library sample code have been provided.

This release also introduces a number of product data sheets, each containing the description of a type of non-document product associated with this end product.

Please refer to the Capabilities section for more details.

## **Package inclusions**

#### New

Identifier	Name	Version
NEHTA-1476:2013	Clinical Documents - Supplementary Guidance for Implementers	1.0
NEHTA-1410:2013	CDA Document Library - Product Data Sheet	1.0
NEHTA-1403:2013	CDA Packaging Library - Product Data Sheet	1.0
NEHTA-1408:2013	Generic CDA Style Sheet - Product Data Sheet	1.0
NEHTA-1407:2013	MDM Client Library – Product Data Sheet	1.0

### Updated (supersedes previous version)

Identifier	Name	Version
NEHTA-1480:2013	Common – Clinical Document - Release Note	1.1.1
NEHTA-1486:2013	CDA Document Library Sample Code .net Client	3.0.9
NEHTA-1487:2013	CDA Document Library Sample Code .net Client	3.1.0
NEHTA-1496:2013	Generic CDA Stylesheet - CSS	1.2.8
NEHTA-1497:2013	Generic CDA Stylesheet	1.2.8

### No change

Identifier	Name	Version
NEHTA-1446:2013	Clinical Documents - Common Conformance Profile	1.4
NEHTA-1097:2011	Implementation Guidance - Representing Coding in CDA Documents	1.0
NEHTA-1096:2011	FAQ OIDs For HL7	1.0
NEHTA-1165:2012	FAQ MIMS Coding Requirements	1.0
NEHTA-1191:2012	Australian CDA Schemas	3.0
NEHTA-1192:2012	Australian CDA Compilable Schema	3.0

Identifier	Name	Version
NEHTA-1199:2012	CDA Rendering Specification	1.0
NEHTA-1217:2012	Reference Platform - Vendor End 2 End Portal	1.4
NEHTA-1229:2011	CDA Package	1.0
NEHTA-1255:2012	FAQ Appropriate use of date and date-time values in Clinical Documents	1.0
NEHTA-1256:2012	FAQ Trusted source on a CDA Package	1.0
NEHTA-1226:2011	Clinical Package	1.0
NEHTA-1271:2013	FAQ Rendering Specification	2.0
NEHTA-1275:2013	FAQ Pathology Date Time	1.1
NEHTA-1278:2013	FAQ Representing Patient IDs in CDA documents	1.1
NEHTA-1270:2013	FAQ - Clarification on Messaging and CDA Packaging	1.4
NEHTA-1276:2013	FAQ Hash Value Verification	1.0
NEHTA-1277:2013	FAQ Patient Medications	1.1
NEHTA-1288:2013	eSignature - Java Library	1.1.0
NEHTA-1285:2013	CDA Packaging Library .java Client	1.2.2
NEHTA-1287:2013	HL7 MDM Library .net	1.0.6
NEHTA-1286:2013	HL7 MDM Library java	1.1.4
NEHTA-1328:2013	Supplementary Notes for Implementers Relating to Clinical Document Presentation	1.0
NEHTA-1329:2012	Conformance Test Specification for Clinical Documents	1.2
NEHTA-1330:2012	Conformance Test Specification for CDA Rendering	1.3
NEHTA-1331:2012	Conformance Test Specification for CDA Packaging	1.4
NEHTA-1378:2013	CDA Packaging Library .net Client v1.3.7	1.3.7

### Removed

None

### Scope

The scope of this end product has not changed as part of this release.

## Stakeholders

Stakeholder groups involved in the development of this release include:

- Commonwealth Department of Health
- Commonwealth Department of Human Services
- ACRRM
- AMA
- AMSANT
- APNA
- RACGP

## Audience

- Implementers of clinical systems producing or consuming clinical documents
- Senior managers and policy makers, clinical experts, health information managers, IT operations and support teams, and system integrators
- Technical and non-technical readers

## **Known issues**

None known

## Capabilities

### CUP Release 1

CUP Release 1 has been focused on resolving key usability issues with clinical information systems used by general practitioners. The guidance provided as part of this release represents the outcomes of several workshops with clinical consultation groups.

It is documented in the new product component *Clinical Documents – Supplementary Guidance for Implementers v1.0*.

Note that additional guidance resulting from CUP Release 1 is released as updates to other specification products, namely Shared Health Summary v1.4.1 and PCEHR B2B Gateway Service.

Complementing this guidance are updated versions of the Generic CDA Stylesheet including its associated CSS file.

### CDA Document Library for PCEHR Releases 3.5 and 4

The sample code for the CDA Document Library has been updated and is now provided in two different versions as part of this release of the *Common – Clinical Document* end product. Version 3.0.9 is aligned with the template packages supported by PCEHR Release 3.5; version 3.1.0 is aligned with the template packages supported by PCEHR Release 4.

The tables below provide overviews of the template packages supported by versions 3.0.9 and 3.1.0 of the CDA Document Library sample code, respectively.

CDA Doc Lib	Document Type	Conf. Level	HPI-Is	Template Package ID	Template Pkg Version
3.0.9	Shared Health Summary	ЗA	enforced	1.2.36.1.2001.1006.1.16565.3	42
	Event Summary	3A	enforced	1.2.36.1.2001.1006.1.16473.7	999
	Discharge Summary	1A	relaxed	1.2.36.1.2001.1006.1.20000.13	31147
		1B	relaxed	1.2.36.1.2001.1006.1.20000.14	31147
		2	relaxed	1.2.36.1.2001.1006.1.20000.15	31147
		3A	relaxed	1.2.36.1.2001.1006.1.20000.16	31147
		3B	relaxed	1.2.36.1.2001.1006.1.20000.17	31147
	eReferral	1A	enforced	1.2.36.1.2001.1006.1.21000.12	145
		1B	enforced	1.2.36.1.2001.1006.1.21000.9	142
		2	enforced	1.2.36.1.2001.1006.1.21000.10	143
		3A	enforced	1.2.36.1.2001.1006.1.21000.11	144

Table 1: Template packages supported by CDA Document Library v3.0.9

CDA Doc Lib	Document Type	Conf. Level	HPI-Is	Template Package ID	Template Pkg Version
	Specialist Letter	1A	enforced	1.2.36.1.2001.1006.1.16615.12	149
		1B	enforced	1.2.36.1.2001.1006.1.16615.9	146
		2	enforced	1.2.36.1.2001.1006.1.16615.10	147
		3A	enforced	1.2.36.1.2001.1006.1.16615.11	148
	PCEHR Dispense Record	3A	enforced	1.2.36.1.2001.1006.1.171.1	30988
	PCEHR Prescription Record	3A	enforced	1.2.36.1.2001.1006.1.170.1	30979

CDA **HPI-Is Document Type** Conf. Template Package ID Template Doc Level Pka Lib Version 1.2.36.1.2001.1006.1.16565.4 3.1.0 Shared Health Summary 3A relaxed 32620 3B relaxed 1.2.36.1.2001.1006.1.16565.5 32620 3A enforced 1.2.36.1.2001.1006.1.16565.6 32620 3B enforced 1.2.36.1.2001.1006.1.16565.7 32620 3A relaxed 1.2.36.1.2001.1006.1.16473.9 32620 **Event Summary** 3B relaxed 1.2.36.1.2001.1006.1.16473.8 32620 3A enforced 32620 1.2.36.1.2001.1006.1.16473.10 3B enforced 1.2.36.1.2001.1006.1.16473.11 32620 **Discharge Summary** 1A relaxed 1.2.36.1.2001.1006.1.20000.18 32620 1.2.36.1.2001.1006.1.20000.19 1B relaxed 32620 2 1.2.36.1.2001.1006.1.20000.20 32620 relaxed 3A relaxed 1.2.36.1.2001.1006.1.20000.21 32620 3B relaxed 1.2.36.1.2001.1006.1.20000.22 32620 1A enforced 1.2.36.1.2001.1006.1.20000.23 32620 enforced 1.2.36.1.2001.1006.1.20000.24 32620 1B enforced 2 1.2.36.1.2001.1006.1.20000.25 32620 1.2.36.1.2001.1006.1.20000.26 3A enforced 32620 1.2.36.1.2001.1006.1.20000.2732620 3B enforced eReferral 1A relaxed 1.2.36.1.2001.1006.1.21000.13 32624 1B relaxed 1.2.36.1.2001.1006.1.21000.14 32624 2 relaxed 1.2.36.1.2001.1006.1.21000.15 32624 3A relaxed 1.2.36.1.2001.1006.1.21000.16 32624 3B relaxed 1.2.36.1.2001.1006.1.21000.17 32624 enforced 32624 1A 1.2.36.1.2001.1006.1.21000.18 enforced 1B 1.2.36.1.2001.1006.1.21000.19 32624 1.2.36.1.2001.1006.1.21000.20 2 enforced 32624 3A enforced 1.2.36.1.2001.1006.1.21000.21 32624

### Table 2: Template packages supported by CDA Document Library v3.1.0

enforced

1.2.36.1.2001.1006.1.21000.22

3B

32624

CDA Doc Lib	Document Type	Conf. Level	HPI-Is	Template Package ID	Template Pkg Version
	Specialist Letter	1A	relaxed	1.2.36.1.2001.1006.1.16615.13	32624
		1B	relaxed	1.2.36.1.2001.1006.1.16615.14	32624
		2	relaxed	1.2.36.1.2001.1006.1.16615.15	32624
		ЗA	relaxed	1.2.36.1.2001.1006.1.16615.16	32624
		3B	relaxed	1.2.36.1.2001.1006.1.16615.17	32624
		1A	enforced	1.2.36.1.2001.1006.1.16615.18	32624
		1B	enforced	1.2.36.1.2001.1006.1.16615.19	32624
		2	enforced	1.2.36.1.2001.1006.1.16615.20	32624
		ЗA	enforced	1.2.36.1.2001.1006.1.16615.21	32624
		3B	enforced	1.2.36.1.2001.1006.1.16615.22	32624
	PCEHR Dispense Record	3A	relaxed	1.2.36.1.2001.1006.1.171.2	32566
		3A	enforced	1.2.36.1.2001.1006.1.171.3	32566
	PCEHR Prescription Record	3A	relaxed	1.2.36.1.2001.1006.1.170.2	32566
		3A	enforced	1.2.36.1.2001.1006.1.170.3	32566

### **Product Data Sheets**

This release also introduces a number of Product Data Sheets (PDS). The purpose of a PDS is to provide a description of a type of deliverable that is not in document format.

## EP-1457:2013 Common – Clinical Document v1.1

Release note: NEHTA-1473:2013, 09 October 2013

## Release rationale

This release of the Common Clinical Document end product introduces updates to the conformance profile for Common Clinical Documents, as mandated by the following approved change requests.

More detailed information about the referenced change requests is provided in the Capabilities section of this document and can be accessed by following the provided hyperlinks.

Change Request ID	Change request title	Impact on this release
CCB-0116	Relaxation of the mandatory use of HPI-Is in uploaded documents	New conformance requirements added for local identifiers.
CCB-0222	Support for CSP Certificates in CDA Documents	Conformance requirements regarding digital signatures previously contained in document-type specific Conformance Profiles have been consolidated and revised in this version of the Common Conformance Profile.
		New conformance requirements added for Legal Authenticator, Approver and Custodian.

In addition to these changes, the structure of the document has been modified to improve clarity and readability. This structural change does not affect the contents of any of the conformance requirements.

## Package inclusions

### New

None

### Updated (supersedes previous version)

Identifier	Name	Version
NEHTA-1473:2013	Common – Clinical Document - Release Note	1.1
NEHTA-1446:2013	Clinical Documents - Common Conformance Profile	1.4

### No change

Identifier	Name	Version
NEHTA-1097:2011	Implementation Guidance - Representing Coding in CDA Documents	1.0
NEHTA-1096:2011	FAQ OIDs For HL7	1.0
NEHTA-1165:2012	FAQ MIMS Coding Requirements	1.0
NEHTA-1191:2012	Australian CDA Schemas	3.0
NEHTA-1192:2012	Australian CDA Compilable Schema	3.0
NEHTA-1199:2012	CDA Rendering Specification	1.0

Identifier	Name	Version
NEHTA-1200:2012	Generic CDA Stylesheet	1.1.12
NEHTA-1217:2012	Reference Platform - Vendor End 2 End Portal	1.4
NEHTA-1229:2011	CDA Package	1.0
NEHTA-1255:2012	FAQ Appropriate use of date and date-time values in Clinical Documents	1.0
NEHTA-1256:2012	FAQ Trusted source on a CDA Package	1.0
NEHTA-1226:2011	Clinical Package	1.0
NEHTA-1271:2013	FAQ Rendering Specification	2.0
NEHTA-1275:2013	FAQ Pathology Date Time	1.1
NEHTA-1278:2013	FAQ Representing Patient IDs in CDA documents	1.1
NEHTA-1270:2013	FAQ - Clarification on Messaging and CDA Packaging	1.4
NEHTA-1276:2013	FAQ Hash Value Verification	1.0
NEHTA-1277:2013	FAQ Patient Medications	1.1
NEHTA-1288:2013	eSignature - Java Library	1.1.0
NEHTA-1283:2013	Generic CDA Stylesheet	1.2.7
NEHTA-1282:2013	Generic CDA Stylesheet - CSS	1.2.7
NEHTA-1285:2013	CDA Packaging Library .java Client	1.2.2
NEHTA-1287:2013	HL7 MDM Library .net	1.0.6
NEHTA-1286:2013	HL7 MDM Library java	1.1.4
NEHTA-1328:2013	Supplementary Notes for Implementers Relating to Clinical Document Presentation	1.0
NEHTA-1329:2012	Conformance Test Specification for Clinical Documents	1.2
NEHTA-1330:2012	Conformance Test Specification for CDA Rendering	1.3
NEHTA-1332:2013	CDA Document Library Sample Code .net Client	3.0.7
NEHTA-1331:2012	Conformance Test Specification for CDA Packaging	1.4
NEHTA-1378:2013	CDA Packaging Library .net Client v1.3.7	1.3.7

### Removed

None

### Scope

The scope of the Common Clinical Document end product has not been changed as part of this release.

## Stakeholders

The following stakeholders have been involved in the development of this release:

- DOHA
- Accenture
- CCA Governance Group

### Audience

- Implementers of clinical systems producing or consuming Consumer Entered Information clinical documents
- Senior managers and policy makers, clinical experts, health information managers, IT operations and support teams, and system integrators
- Technical and non-technical readers

## Capabilities

The following sections provide additional details for each of the change requests addressed in this release.

### CCB-0116

The change request introduces the temporary and limited relaxation of the mandatory requirement to include HPI-Is for a number of clinical document types. A similar relaxation had already been applied to eDischarge Summary documents with release 1.4 of the eDischarge Summary end product.

New conformance requirements have been added to the specific Conformance Profiles of the affected document types that require local identifiers to be included in a clinical document wherever an HPI-I has been omitted. Complementary conformance requirements clarifying the semantics of local identifiers have been introduced in the new version 1.4 of the *Clinical Documents – Common Conformance Profile* document in this release.

### CCB-0222

The change request introduces support for digital signatures created with CSP digital certificates for all types of clinical documents.

The new version 1.4 of the *Clinical Documents – Common Conformance Profile* document in this release contains a consolidated and revised version of conformance requirements previously contained in the specific Conformance Profiles of the affected document types. This revised version introduces support for digital signatures created with CSP digital certificates.

New conformance requirements have been added to clarify the semantics of Legal Authenticator, Approver and Custodian.

## EP-1094:2011 Common - Clinical Document v1.0.2

Release note: NEHTA-1380:2013, 4 July 2013

## **Release rationale**

This incremental release includes:

- updated sample code to address a small change in the CDA packaging library
- updates to three FAQ title prefixes (document content is unchanged).

### Package inclusions

### Updated (supersedes previous version)

Identifier	Name	Version
NEHTA-1378:2013	CDA Packaging Library .net Client	1.3.7

### No change to the following inclusions

Identifier	Name	Version
NEHTA-1097:2011	Implementation Guidance - Representing Coding in CDA Documents	1.0
NEHTA-1096:2011	FAQ OIDs For HL7	1.0
NEHTA-1165:2012	FAQ MIMS Coding Requirements	1.0
NEHTA-1191:2012	Australian CDA Schemas	3.0
NEHTA-1192:2012	Australian CDA Compilable Schema	3.0
NEHTA-1199:2012	CDA Rendering Specification	1.0 07-
NEHTA-1200:2012	Generic CDA Stylesheet	1.1.12
NEHTA-1217:2012	Reference Platform - Vendor End 2 End Portal	1.4 23-
NEHTA-1229:2011	CDA Package	1.0
NEHTA-1255:2012	FAQ Appropriate use of date and date-time values in Clinical Documents	1.0
NEHTA-1256:2012	FAQ Trusted source on a CDA Package	1.0
NEHTA-1226:2011	Clinical Package	1.0
NEHTA-1263:2012	Common Conformance Profile	1.3
NEHTA-1271:2013	FAQ Rendering Specification	2.0
NEHTA-1275:2013	FAQ Pathology Date Time	1.1
NEHTA-1278:2013	FAQ Representing Patient IDs in CDA documents	1.1
NEHTA-1270:2013	FAQ - Clarification on Messaging and CDA Packaging	1.4
NEHTA-1276:2013	FAQ Hash Value Verification	1.0
NEHTA-1277:2013	FAQ Patient Medications	1.1
NEHTA-1288:2013	Java Library	1.1.0

Identifier	Name	Version
NEHTA-1283:2013	Generic CDA Stylesheet	1.2.7
NEHTA-1282:2013	Generic CDA Stylesheet - CSS	1.2.7
NEHTA-1285:2013	CDA Packaging Library .java Client	1.2.2
NEHTA-1287:2013	HL7 MDM Library .net	1.0.6
NEHTA-1286:2013	HL7 MDM Library java	1.1.4
NEHTA-1328:2013	Supplementary Notes for Implementers Relating to Clinical Document Presentation	1.0
NEHTA-1331:2012	Conformance Test Specification for CDA Packaging	1.4
NEHTA-1330:2012	Conformance Test Specification for CDA Rendering	1.3
NEHTA-1329:2012	Conformance Test Specification for Clinical Documents	1.2
NEHTA-1332:2013	CDA Document Library Sample Code .net Client [Update]	3.0.7

#### Removed

None

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