
Clinical Document Conformance Report

(for producers of clinical documents)

Prepared by

for

Report identifier:

Test date: Friday, 19 April 2024

Acknowledgements

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Disclaimer

Before formally declaring conformity to the System Operator of the My Health Record system, developers need to perform additional tests to ensure full coverage of all requirements. Please refer to the Clinical Package Validator Product Data Sheet for more detailed information about the Validator's coverage of requirements and conformance test cases.

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1 Preface

1.1 Background

A developer who wants their health software systems to send clinical information to, or receive clinical information from, another health software system or the My Health Record System must assess whether their software conforms to the relevant requirements.

The Clinical Package Validator (Validator) has been developed as a tool for testing whether a health software system is producing conformant clinical documents. The Validator applies a number of tests to a sample clinical document and reports the outcomes of these tests in this report. These tests assess whether the health software system is:

1. Producing sample clinical documents that conform to specifications for the document's syntax and structure;
2. Producing sample clinical documents with data element values that are the same as the test data that was entered into the health software system;
3. Producing clinical packages that conform to the requirements for sending a clinical package of information to the My Health Record or to another health software system;
4. Producing sample clinical documents that conform to the requirements for document authoring; and
5. Using valid code sets (e.g. clinical terminology).

The Validator tests one sample clinical document at a time and each sample clinical document is populated with values from a single set of test data. A tester who is assessing the ability of a health software system to generate a particular type of clinical document may need to run the Validator several times because several sets of test data may have been defined for a type of clinical document. This process must be repeated for each type of clinical document that the health software system produces.

This document is generated each time the tester runs the Validator. While most of the information in this report is generated automatically, the tester should also supply any missing information, and include any comments if needed. This document contains the results from each test that was applied by the Validator, and a summary of the overall result of each test that was applied. To help the tester to analyse the results, an image of the rendered sample clinical document and the sample clinical document XML are included in the appendices.





1.2 Document purpose

A tester may use the information in this document to determine whether a health software system is producing clinical documents that conform to the relevant requirements.

2 Assessment Details

<i>Developer name</i>	
<i>Name of implementation under test</i>	
<i>Version of implementation under test</i>	
<i>Test environment</i>	
<i>Location of assessment (address)</i>	
<i>Tester name</i>	
<i>Tester organisation</i>	
<i>Date & time of assessment</i>	Friday, 19 April 2024 - 11:08:02 AM
<i>Context of test (MHR or P2P)</i>	MHR
<i>Input File</i>	C:\Users\MuhammadAbubakar\StrongRoom\mhr-b2b-client-node\src\doc.xml
<i>Other information</i>	
<i>Clinical Package Validator version</i>	3.3
<i>Template Package ID</i>	Specialist Letter 1A (1.2.36.1.2001.1006.1.16615.28 ver 35416)
<i>Document Template ID</i>	1.2.36.1.2001.1001.101.100.1002.132
<i>Clinical document type</i>	Specialist Letter
<i>Target conformance level</i>	1A

3 Symbols

Symbol	Explanation
	Fail
	Warning
	Pass
	Not Run

4 Terminologies Used

Australian Medicines Terminology (AMT), 2.56

Australian Medicines Terminology (AMT), <http://snomed.info/sct/900062011000036108/version/20151130>

Australian PBS Code, 20230501

SNOMED CT-AU, <http://snomed.info/sct/32506021000036107/version/20230531>

Health Care Facility Type Code

ANZSCO Type Code

SNOMED CT

Australian Medicines Terminology (AMT)

Australian Vaccine Code

HL7 Identifier Type Code

Clinical specialties

Australian PBS Manufacturer Code

Australian PBS Code

5 Clinical Document Test Results Summary

	Overall result	Error count	Warning count	Comments
<i>Template Validation</i>	✓	0	0	
<i>Package Validation</i>	n/a	n/a		

	Overall result	Total count	Error count	Warning count	Hint count	Finding count	Recommendation count
<i>Additional Rules Validation</i>	—		0				

6 **Template Validation Report**

No Template Errors Detected

7 **Additional Rules Report**

Not Run

8 **Package Report**

Input File not a CDA Package

9 Signature File Information

Name	Description
Signing Time	
Approver Person Id	
Approver Person Name	
NASH Certificate ?	
Subject	
Valid From	
Valid To	
Certificate Policies	
Other signature files in package	

Appendix A: Rendered Clinical Document

See external file: C:\Users\MuhammadAbubakar\StrongRoom\mhr-b2b-client-node\src\doc.xml_Report-1A-20240419110758.htm