



Electronic Prescribing Information for Transitional eNRM C Software Vendors

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Transitional eNRM C (Electronic National Residential Medication Chart) software vendors will go through the following process to have their product approved as a Transitional eNRM C Product:

1. Vendor engages their preferred Prescription Delivery Service (PDS).
2. Vendor ensures product meets the technical specifications of the chosen PDS.
3. Vendor ensures product meets all Conformance Profile (CP) v3.0.1 conformance requirements for a medication chart prescribing system.
4. Vendor gains access to PDS's eNRM C transition approved CPv3.0.1 test environment.
5. Vendor completes all relevant conformance test cases in the Conformance Test Specifications (CTS) v3.0.3.
6. Vendor submits test evidence to the Agency for assessment. See instructions below.
7. The Agency reviews test evidence and results (around 2 – 3 weeks).
8. The Agency will advise the Department of Health and Aged Care of the outcome.
9. The Department of Health and Aged Care will send a Deed of Agreement with 'self-declaration' of conformance with Transitional eNRM C conformance requirements to the vendor for signing.
10. Once signed, the Agency will list the approved Transitional eNRM C Product on the Transitional eNRM C Conformance Register.
11. Vendors can start operating under the conditions of the Transitional Arrangement, as per the Legislative Instrument. For clarifications about the legislative framework for electronic prescribing/eNRM C, please contact the Department of Health and Aged Care at eNRM C@health.gov.au.
12. If formally operating under the eNRM C Trial conditions, the Department of Health and Aged Care will remove the vendor from the Schedule on the Special Arrangement.

Instructions how to prepare test evidence:

Clear and unambiguous test evidence increases the chance of fast processing.

1. Test evidence and a clear explanation of evidence for all the requirements in scope should be provided.
2. When XML or screenshots are provided, the relevant fields must be marked clearly.
3. If data fields in XML or screenshots are named differently to the names in the conformance requirements, please provide clear information.
4. For the test cases that represent flow or change of state (i.e. before and after logging in), test evidence for all the relevant steps and expected results should be provided and described.

Instructions how to send test evidence to the Agency for assessment:

1. Vendors send test evidence and completed CTS v3.0.3 to help@digitalhealth.gov.au.
2. If the file is too large to be sent via email, vendors may request the Agency to set up a large file transfer space via GovTeams. Email help@digitalhealth.gov.au.

Important Note about Transitional eNRM Assessment:

While the Agency endeavours to review your Transitional eNRM test evidence thoroughly, it should be noted that it is your organisation responsibility to ensure that your software product meets all relevant mandatory conformance requirements. Software entered on the Transitional eNRM Register of Conformity does not guarantee your software product will be conformant to Electronic Prescribing Conformance Profile v3.0.1. You will still be required to undergo further assessment at the time of registering for Electronic Prescribing Conformance Profile v3.0.1.

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Australian Digital Health Agency ABN 84 425 496 912, Level 17, 1 Eagle Street, Brisbane City, QLD 4000 digitalhealth.gov.au
Telephone 1300 901 001 or email help@digitalhealth.gov.au

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