

Medicines and Poisons Act 2019

Extended Practice Authority 'Registered Nurses'



**Queensland
Government**

Version control

Version	Replaces version	Date approved	Commencement date
1	NA	12 August 2021	27 September 2021

Disclaimer: Please note that any material printed is regarded as an uncontrolled copy. It is the responsibility of the person printing the document to refer frequently to <https://www.health.qld.gov.au/system-governance/licences/medicines-poisons/legislation-standards/acts-regulation> for updates

Medicines and Poisons Act 2019
Extended Practice Authority 'Registered Nurses'

This extended practice authority (EPA) has been made by the Director General, Queensland Health under section 232 of the *Medicines and Poisons Act 2019*. It states the scope of the regulated activities with the regulated substances which a registered nurse is authorised to carry out for the purposes described in column 3 of the table under Schedule 7, Part 3 Division 2 of the Medicines and Poisons (Medicines) Regulation 2021 (the Medicines Regulation).

A term used in this EPA that is defined in the *Medicines and Poisons Act 2019* or the Medicines and Poisons (Medicines) Regulation 2021, has the meaning stated in the *Medicines and Poisons Act 2019* or Medicines Regulation.

1. Conditions – general

The following general conditions apply to a registered nurse in addition to any specific conditions under parts A, B, C and D of the EPA.

- 1.1. The registered nurse must ensure they have access to their applicable health management protocol, *Australian Immunisation Handbook*¹ and current guidelines, manuals or protocols adopted or established by their employer when acting under this EPA.
- 1.2. The registered nurse must act in accordance with a current health management protocol that applies to the dealings of the registered nurse and that complies with the requirements specified in Appendix 1.
- 1.3. The registered nurse must not give a treatment dose of any monitored medicine listed in Schedule 2 Part 4 of the Medicines and Poisons (Medicines) Regulation 2021 unless stated in Appendix 3 Column 3.
- 1.4. Before administering or giving a treatment dose of a medicine the registered nurse must be familiar with the contra-indication(s) and known side effects of the medicine and advise the patient accordingly.
- 1.5. For the administration of vaccines, the registered nurse must act in accordance with:
 - 1.5.1. the requirements for vaccine administration in the current online edition of the *Australian Immunisation Handbook* including for patient selection, patient consent, vaccine administration, documenting vaccination and follow up care; and
 - 1.5.2. the current online edition of the *Immunisation Schedule Queensland*².
- 1.6. Before vaccines are administered, the registered nurse must ensure the equipment and procedures detailed in the current online edition of the *Australian Immunisation Handbook* are in place.

¹ For current on-line Australian Immunisation Handbook see <https://immunisationhandbook.health.gov.au/>

² See <https://www.health.qld.gov.au/clinical-practice/guidelines-procedures/diseases-infection/immunisation/schedule>

- 1.7. Prior to performing a Tuberculin Skin Test (TST) or administering a bacillin Calmette-Guerin (BCG) vaccination for Tuberculosis (TB), registered nurses must demonstrate that they have completed the specified training and been assessed by a clinician as required by the Health Services Directive – Tuberculosis Control ³ and the Health Service Directive Protocol for the Control of Tuberculosis ⁴.
- 1.8. When vaccines are in the possession of the registered nurse, the registered nurse must ensure that the storage and transport of vaccines is in accordance with the *National vaccine storage guidelines: Strive for 5* ⁵.
- 1.9. The registered nurse who administers a vaccine must ensure:
 - 1.9.1. the vaccination is recorded on the Australian Immunisation Register (AIR) as soon as practicable and ideally at the time of vaccination
 - 1.9.2. any adverse events occurring following immunisation must be notified using the Adverse Event Following Immunisation (AEFI) form published on the Queensland Health website ⁶.
- 1.10. If Consumer Medicine Information (CMI) ⁷ is available for a particular medicine, the registered nurse must, where reasonably practicable, offer the information to each person to whom the registered nurse administers or gives a treatment dose of the medicine.

Note that a registered nurse is authorised under Schedule 7, Part 3 Division 2 of the Medicines and Poisons (Medicines) Regulation 2021 to administer an S2 or S3 medicine without an extended practice authority. This EPA authorises the giving of a treatment dose of S2 or S3 medicines in the circumstances specified the EPA.

2. Part A – specified services to address health and community need

- 2.1. A registered nurse may only administer or give a treatment dose of medicines under Part A of this EPA if the registered nurse is working for a Hospital and Health Service that uses a credentialing process meeting the requirements of the current Health Service Directive: Credentialing and defining the scope of clinical practice ⁸ or the current Australian Commission on Safety and Quality in Health Care *Standard for Credentialing and Defining the Scope of Clinical Practice* ⁹ to define a **credentialed scope of clinical practice**.
- 2.2. The registered nurse must be credentialed to perform a **specified service** ¹⁰ and must have a document stating their credentialed scope of clinical practice for the specified service.

³ https://www.health.qld.gov.au/_data/assets/pdf_file/0023/150935/qh-hsd-040.pdf

⁴ https://www.health.qld.gov.au/_data/assets/pdf_file/0024/155175/qh-hsdptl-040-1.pdf

⁵ For National vaccine storage guidelines: Strive for 5 see

<https://www.health.gov.au/resources/publications/national-vaccine-storage-guidelines-strive-for-5>

⁶ For Adverse Event Following Immunisation (AEFI) form see

https://www.health.qld.gov.au/_data/assets/pdf_file/0033/442968/aefi-reporting-form.pdf

⁷ Consumer Medicines Information is a leaflet that contains information on the safe and effective use of a prescription medicine, as well as some non-prescription medicines – see

<https://www.tga.gov.au/consumer-medicines-information-cmi>

⁸ [Credentialing and defining the scope of clinical practice \(health.qld.gov.au\)](https://www.health.qld.gov.au/credentialing-and-defining-the-scope-of-clinical-practice)

⁹ <https://www.safetyandquality.gov.au/publications-and-resources/resource-library/standard-credentialing-and-defining-scope-clinical-practice>

¹⁰ Specified service means service provided by a Hospital and Health Service to meet a health and community need that is facilitated by the extended practice authority.

- 2.3. The document stating the registered nurse's credentialed scope of clinical practice for the specified service must specify the following:
 - 2.3.1. the health management protocol that applies to the credentialed scope of clinical practice for the registered nurse; and
 - 2.3.2. any specific restrictions or requirements applied under the credentialing process or in accordance with the policies or procedures established for the Hospital and Health Service including:
 - 2.3.2.1. any activities or dealings with medicines that require either direct or indirect supervision by a senior health practitioner, or that may only be undertaken after consultation with a senior health practitioner.
 - 2.3.2.2. any restrictions, conditions, or other limitations on the registered nurse's authority to administer or give a treatment dose of a medicine e.g. that the registered nurse is not authorised to give deep intra-muscular injections.
 - 2.3.2.3. any qualifications and/or training that the registered nurse must hold; and
 - 2.3.3. the date the credentialed scope of clinical practice commences and the dates for review of the credentialed scope of clinical practice.
- 2.4. The document stating the registered nurse's credentialed scope of clinical practice for the specified service must be accessible by the registered nurse when the registered nurse is performing the credentialed service.
- 2.5. If the registered nurse acts within their credentialed scope of clinical practice and in accordance with the health management protocol specified in the document stating the registered nurse's credentialed scope of clinical practice, the registered nurse is authorised to:
 - 2.5.1. administer or give a treatment dose of a medicine listed in Appendix 2 column 1 by the route of administration stated in column 2.
 - 2.5.2. give a treatment dose of an S2 or S3 medicine to a patient requiring treatment or care for the specified service.
 - 2.5.3. administer vaccines listed in column 1 Appendix 4 for the Queensland Immunisation Program.
- 2.6. If the credentialed scope of clinical practice for a registered nurse includes immunisation with Japanese encephalitis vaccine or BCG vaccine, or tetanus immunoglobulin, the registered nurse is authorised to:
 - 2.6.1. administer Japanese encephalitis vaccine only if working under a Japanese Encephalitis immunisation program approved by the chief executive of the Torres and Cape Hospital and Health Service (TCHHS) or the Public Health Medical Officer of the TCHHS and in accordance with local procedures for the Japanese Encephalitis Vaccine Program for Outer Islands of the Torres Strait.
 - 2.6.2. administer BCG vaccine or tuberculin skin test only under a Tuberculosis immunisation program in accordance with the Health Services Directive – Tuberculosis Control and the Health Service Directive Protocol for the Control of Tuberculosis.
 - 2.6.3. administer tetanus immunoglobulin in accordance with the requirements for vaccine administration in the current online edition of the *Australian Immunisation Handbook*.

3. Part B – services in rural and isolated practice areas

- 3.1. A registered nurse may only administer or give a treatment dose of medicines under Part B of this EPA if the registered nurse:
- 3.1.1. is practicing in an isolated practice area or at a rural hospital ¹¹; and
 - 3.1.2. has completed a program of study relevant to the use of medicines in providing emergency and acute care in rural and isolated practice; and
 - 3.1.3. the program of study was previously recognised by the Nursing and Midwifery Board of Australia (NMBA) to enable the registration of the registered nurse to be endorsed as ‘qualified to obtain, supply and administer schedule 2,3,4 & 8 medicines for nursing practice in a rural and isolated practice area’ ¹² under the *Health Practitioner Regulation National Law (Queensland)*.
- 3.2. The registered nurse to whom Part B applies is authorised to:
- 3.2.1. administer a medicine listed in Appendix 3, Column 1 by the route of administration specified in Column 2 and in accordance with any applicable restrictions or conditions specified in Column 3 and in accordance with the health management protocol for the rural and remote ambulatory care settings approved for the Hospital and Health Service or other organisation by whom the registered nurse is employed (see Appendix 1).
 - 3.2.2. administer a medicine that is a vaccine that contains any of the antigens listed in Appendix 4, Column 1.
 - 3.2.3. administer Japanese encephalitis vaccine only if working under a Japanese Encephalitis immunisation program approved by the chief executive of the Torres and Cape Hospital and Health Service (TCHHS) or the Public Health Medical Officer of the TCHHS and in accordance with local procedures for the Japanese Encephalitis Vaccine Program for Outer Islands of the Torres Strait.
 - 3.2.4. administer BCG vaccine or tuberculin skin test only under a Tuberculosis immunisation program in accordance with the Health Services Directive – Tuberculosis Control and the Health Service Directive Protocol for the Control of Tuberculosis.
 - 3.2.5. administer tetanus immunoglobulin in accordance with the requirements for vaccine administration in the current online edition of the *Australian Immunisation Handbook*.
 - 3.2.6. give a treatment dose of a medicine in Appendix 3 where Column 3 states that a medicine may be given as a treatment dose and in accordance with any applicable restrictions or conditions specified in Column 3 and in accordance with the health management protocol for the for the rural and remote ambulatory care settings approved for the Hospital and Health Service or other organisation by whom the registered nurse is employed (see Appendix 1).

¹¹ See definitions of isolated practice area and rural hospital in the Medicines Regulation Schedule 22

¹² See NMBA former ‘Registration standard for endorsement for scheduled medicines registered nurses (rural and isolated practice)’
<https://www.ahpra.gov.au/documents/default.aspx?record=WD10%2f138&dbid=AP&checksum=0wq5Z4dBrn1c6oU16s2H2A%3d%3d>

3.2.7. give a treatment dose of an S2 or S3 medicine to a patient requiring treatment or care at a rural hospital or in an isolated practice area ¹³.

4. Part C – sexual or reproductive health services

- 4.1. A registered nurse may only administer or give a treatment dose of medicines under Part C of this EPA if the registered nurse has successfully completed:
 - 4.1.1. an approved program of study for endorsement as a Sexual Health Program Nurse with the former Queensland Nursing Council.
 - 4.1.2. a qualification in sexual health previously approved by the chief executive under the (repealed) Health (Drugs and Poisons) Regulation 1996.
- 4.2. To the extent necessary to provide sexual or reproductive health services, a registered nurse is authorised to administer or give a treatment dose of a medicine if any of the following apply:
 - 4.2.1. the services are provided under a sexual or reproductive health program carried out by a Hospital and Health Service.
 - 4.2.2. the services are provided under a sexual or reproductive health program that was certified under the (repealed) Health (Drugs and Poisons) Regulation 1996 until the date stated in the program to be the end of the certification.
 - 4.2.3. the service is provided under a general approval given for a sexual or reproductive health program under the *Medicines and Poisons Act 2019*.
- 4.3. The registered nurse to whom Part C of this EPA applies is authorised to:
 - 4.3.1. administer a medicine listed in Appendix 3, Column 1 by the route of administration specified in Column 2 and in accordance with any applicable restrictions or conditions specified in Column 4 and in accordance with an approved health management protocol for the service (see Appendix 1).
 - 4.3.2. administer a medicine that is a vaccine that contains any of the antigens listed in Appendix 4, Column 1 in accordance with the conditions in Column 2
 - 4.3.3. give a treatment dose of a medicine in Appendix 3 where Column 4 states that a medicine may be given and in accordance with any applicable restrictions or conditions specified in Column 4, and in accordance with the health management protocol for the service (see Appendix 1).
- 4.4. The registered nurse may only administer or give a treatment dose of an hormonal contraceptive or a contraceptive vaginal ring mentioned in Appendix 3 if the client has had an initial assessment and been prescribed a hormonal contraceptive or a contraceptive vaginal ring by an authorised prescriber at the sexual or reproductive health service where the client is currently attending.

¹³ See definitions of isolated practice area and rural hospital in the Medicines Regulation Schedule 22.

5. Part D – immunisation program services

- 5.1. A registered nurse may only administer the medicines under Part D of this EPA if the registered nurse has successfully completed any of the following:
 - 5.1.1. an approved program of study for endorsement as an Immunisation Program Nurse with the former Queensland Nursing Council.
 - 5.1.2. a qualification in immunisation previously approved by the chief executive under the (repealed) Health (Drugs and Poisons) Regulation 1996.
 - 5.1.3. an accredited immunisation training course that contains learning objectives equivalent to the domains in the *National Immunisation Education Framework for Health Professionals*¹⁴.
- 5.2. To the extent necessary to provide immunisation services, a registered nurse is authorised to administer a medicine if any of the following apply:
 - 5.2.1. the services are provided under an immunisation program carried out by a Hospital and Health Service.
 - 5.2.2. the services are provided under an immunisation program carried out by the Queensland Health.
 - 5.2.3. the services are provided under an immunisation program carried out by a local government.
 - 5.2.4. the services are provided under an immunisation program that was certified under the (repealed) Health (Drugs and Poisons) Regulation 1996 and until the date stated in the program to be the end of the certification.
 - 5.2.5. the services are provided under an immunisation program authorised under a general approval given to provide an immunisation program under the *Medicines and Poisons Act 2019*.
- 5.3. The registered nurse to whom Part D applies is authorised to:
 - 5.3.1. administer a medicine that is a vaccine that contains any of the antigens listed in Appendix 4, Column 1.
 - 5.3.2. administer Japanese encephalitis vaccine only if working under a Japanese Encephalitis immunisation program approved by the chief executive of the Torres and Cape Hospital and Health Service (TCHHS) or the Public Health Medical Officer of the TCHHS and in accordance with local procedures for the Japanese Encephalitis Vaccine Program for Outer Islands of the Torres Strait.
 - 5.3.3. administer BCG vaccine or tuberculin skin test only under a Tuberculosis immunisation program in accordance with the Health Services Directive – Tuberculosis Control and the Health Service Directive Protocol for the Control of Tuberculosis.
 - 5.3.4. administer tetanus immunoglobulin in accordance with the requirements for vaccine administration in the current online edition of the *Australian Immunisation Handbook*.

¹⁴ <https://www.health.gov.au/resources/publications/national-immunisation-education-framework-for-health-professionals>

Appendix 1 – Requirements for health management protocols

1. A health management protocol is a document approved and dated by the chief executive ¹⁵ of a Hospital and Health Service or the Chief Executive Officer ¹⁶ of a non-Queensland Health employing organisation that details the clinical use of medicines for services provided by a registered nurse under this EPA for patients of the Hospital and Health Service or other employing organisations.
2. A health management protocol must have been reviewed and endorsed by an inter-disciplinary health team comprising, at a minimum, a medical practitioner, a registered nurse and a pharmacist, and may include other identified professional personnel (an ***inter-disciplinary team***).
3. A health management protocol details the clinical use of medicines that may be administered or given as a treatment dose by a registered nurse under Appendix 2 or 3 of this Extended Practice Authority, and must include the following:
 - a) The procedures for clinical assessment, management, and follow up of patients, including the recommended medicine for the relevant clinical problem.
 - b) For each medicine in the health management protocol:
 - i. a clinical indication or time when medical referral/consultation must occur for that condition;
 - ii. the name, form and strength of the medicine and the condition/situation for which it is intended and any contraindications to the use of the medicine;
 - iii. the recommended dose of the medicine, the frequency of administration (including rate where applicable) and the route of administration of the medicine;
 - iv. for a medicine to be administered, the maximum dose of a medicine that may be administered or maximum duration of administration allowed without a prescription from an authorised prescriber;
 - v. for a medicine to be given as a treatment dose, the maximum quantity of or duration of treatment with a medicine that may be given without a prescription;
 - vi. the type of equipment and management procedures required for management of an emergency associated with the use of the medicine.
 - c) When to refer to a higher level of care for intervention or follow-up.
4. A health management protocol is either:
 - d) developed for the Hospital and Health Service or for a non-Queensland Health employing organisation by an inter-disciplinary team for the Hospital and Health Service or other employing organisation (***a local inter-disciplinary team***); or
 - e) a clinical guideline containing all the required content specified in paragraph 3 or a health management protocol developed by another entity's inter-disciplinary team and endorsed as the health management protocol for the service by the local inter-disciplinary team.
5. A health management protocol is ***current*** when it has been approved the chief executive of a Hospital and Health Service or the Chief Executive Officer of a non-Queensland Health employing organisation within two (2) years of the day a registered nurse acts in accordance with the health management protocol unless

¹⁵ refer to *Hospital and Health Services Boards Act 2011*

¹⁶ Chief Executive Officer means the highest-ranking executive or administrator in charge of the management of an organisation.

the current on-line edition of the *Primary Clinical Care Manual (PCCM)* ¹⁷ has been endorsed and approved as the a health management protocol for the service.

6. If PCCM is adopted as the health management protocol, the health management protocol is **current** when it has been approved the chief executive of a Hospital and Health Service or the Chief Executive Officer of a non-Queensland Health employing organisation within three (3) years of the day a registered nurse acts in accordance with the health management protocol.

Note: The current online edition of the PCCM published by Queensland Health contains the principal health management protocols for use in rural and remote ambulatory care settings.

¹⁷ For the Primary Clinical Care Manual - 10th see <https://www.health.qld.gov.au/rrcsu/clinical-manuals/primary-clinical-care-manual-pccm>.

Appendix 2 – Part A Medicines

Note: Monitored medicines are marked with #

Column 1 Medicine	Column 2 Approved route of administration
Aciclovir	Oral
Albendazole	Oral
Amoxicillin	Oral
Amoxicillin/clavulanic acid	Oral
Azithromycin	Oral
Benzathine penicillin (Bicillin L-A)	Intramuscular
Benzatropine	Intramuscular Oral
Box jellyfish anti-venom	Intravenous Intramuscular
Cefaclor	Oral
Cefalexin	Oral
Ceftriaxone	Intramuscular
Cefuroxime	Oral
Ciprofloxacin	Oral
Ciprofloxacin	Ear drops
Clindamycin	Oral
Clindamycin	Intravaginal
Clotrimazole	Intravaginal
Dexamethasone 0.5 mg / framycetin sulfate 5 mg / gramicidin 0.05 mg / mL	Ear drops
Diazepam #	Intravenous Oral Rectal
Dicloxacillin	Oral
Doxycycline	Oral
Erythromycin	Oral
Estradiol	Intravaginal
Estriol	Intravaginal
Ethinylestradiol 20 microgram/ Levonorgestrel 100 microgram	Oral
Ethinylestradiol 20 microgram/ Drospirenone 3mg	
Ethinylestradiol 30 microgram/ Desogestrel 150 microgram	
Ethinylestradiol 30 microgram/ Dienogest 2 mg	
Ethinylestradiol 30 microgram/ Drospirenone 3 mg	
Ethinylestradiol 30 microgram/ Gestodene 75 microgram	
Ethinylestradiol 30 microgram/ Levonorgestrel 150 microgram	
Ethinylestradiol 30 microgram/ Levonorgestrel 50 microgram and Ethinylestradiol 40 microgram/ Levonorgestrel 75 microgram	
Ethinylestradiol 30 microgram/ Levonorgestrel 125 microgram	
Ethinylestradiol 35 microgram/ Northisterone 500 microgram	
Ethinylestradiol 35 microgram/ Cyproterone acetate 2 mg	
Ethinylestradiol 35 microgram/ Northisterone 1mg	
Etonogestrel 120mg / Ethinylestradiol 15 microgram e.g. <i>Nuva Ring</i>	
Famciclovir	Oral
Fentanyl #	Intramuscular Intravenous Subcutaneous

Column 1 Medicine	Column 2 Approved route of administration
Flucloxacillin	Oral
Fluconazole	Oral
Flumazenil	Intravenous
Flumethasone pivalate 0.02 % / clioquinol 1 %	Ear drops
Fluoride varnish	Topical to teeth
Funnel web spider anti-venom	Intravenous
Furosemide (frusemide)	Intramuscular Intravenous
Glyceryl Trinitrate Transdermal Patches	Topical
Haloperidol	Intravenous Intramuscular Oral
Hydrocortisone	Intramuscular Intravenous
Imiquimod	Topical
Ipratropium bromide	Nebulised Metered dose inhaler (MDI)
Ketorolac trometamol	Intramuscular
Levonorgestrel 1.5 mg	Oral
Levonorgestrel 30 microgram e.g. <i>Microval</i>	Oral
Lidocaine (lignocaine) 1 %	Subcutaneous - Local infiltration
Lidocaine (lignocaine) 1 % (with ceftriaxone)	Intramuscular
Lidocaine (lignocaine) gel 2%	Topical
Lidocaine (lignocaine) with adrenaline (epinephrine)	Subcutaneous
Lidocaine (lignocaine)-Prilocaine 5%	Topical
Lorazepam #	Oral
Medroxyprogesterone acetate 150 mg/mL depo-injection	Intramuscular
Methoxyflurane	Inhalation
Metoclopramide	Intravenous Intramuscular Oral
Metronidazole	Oral
Miconazole	Vaginal Topical (including dermal and oral)
Midazolam #	Intravenous Intramuscular Intranasal Buccal
Misoprostol	Rectal Sublingual Buccal
Morphine #	Intramuscular Intravenous Subcutaneous
Mupirocin Cream/ointment	Topical
Naloxone	Subcutaneous Intravenous Intramuscular
Nifedipine	Oral
Nitrofurantoin	Oral
Nitrous Oxide	Topical with cryogun
Norethisterone 350 microgram e.g. <i>Noriday</i>	Oral
Nystatin oral drops	Topical
Olanzapine	Oral

Column 1 Medicine	Column 2 Approved route of administration
Ondansetron	Intravenous Oral
Oxybuprocaine eye drop 0.4 % (minim)	Topical to eye
Oxycodone #	Oral
Oxytocin	Intramuscular Intravenous
Phenoxymethylpenicillin	Oral
Podophyllotoxin	Topical
Prednisolone	Oral
Probenecid	Oral
Procaine benzylpenicillin (procaine penicillin)	Intramuscular
Promethazine	Intramuscular Intravenous
Roxithromycin	Oral
Salbutamol	Nebulised
Silver sulfadiazine 1% cream	Topical
Snake polyvalent anti-venom	Intravenous
Tetanus immunoglobulin	Intramuscular
Triamcinolone compound (otic)	Topical to ear
Trimethoprim	Oral
Trimethoprim/ Sulfamethoxazole	Oral
Valaciclovir	Oral

Appendix 3 – Authorised medicines and dealings

Note:

- For **giving a treatment dose** of an S2 or S3 medicine, the maximum quantity is the smallest available manufacturer's pack;
- Monitored medicines are marked with #;
- Hormonal contraceptive or a contraceptive vaginal ring are marked with †
- 'Consult authorised prescriber' means that an oral prescription is required to authorise further administration of the medicine or to give a treatment dose in accordance with the prescription.

Column 1 Medicine	Column 2 Approved route of administration	Restrictions/Conditions for place or circumstance	
		Column 3 Rural & Isolated Practice Areas	Column 4 Sexual Health Program
Aciclovir	Oral	Administer one dose and give a treatment dose as necessary for genital herpes	Administer one dose and give a treatment dose as necessary for genital herpes
Albendazole	Oral	Administer one dose and give a treatment dose as necessary	Not permitted
Amoxicillin	Oral	Administer one dose and give a treatment dose as necessary	Not permitted
Amoxicillin/clavulanic acid	Oral	Administer one dose and give a treatment dose as necessary	Not permitted
Azithromycin	Oral	Administer one dose and give a treatment dose as necessary	Administer one dose and give a treatment dose as necessary
Benzathine penicillin (Bicillin L-A)	Intramuscular	Administer one dose	Administer one dose
Benzatropine	Intramuscular Oral	Maximum 2 mg. Consult authorised prescriber if more than recommended dose required.	Not permitted
Box jellyfish anti-venom	Intravenous Intramuscular	One ampoule (20,000 units). Consult authorised prescriber if more than recommended dose required.	Not permitted
Cefaclor	Oral	Child only Administer one dose and give a treatment dose as necessary	Not permitted
Cefalexin	Oral	Administer one dose and give a treatment dose as necessary	Administer one dose and give a treatment dose as necessary
Ceftriaxone	Intramuscular	Administer one dose reconstituted with Lidocaine (lignocaine) 1 % injection	Administer one dose reconstituted with Lidocaine (lignocaine) 1 % injection
Cefuroxime	Oral	Adult only Administer one dose and give a treatment dose as necessary	Not permitted
Ciprofloxacin	Oral	Single dose only	Single dose only

Column 1 Medicine	Column 2 Approved route of administration	Restrictions/Conditions for place or circumstance	
		Column 3 Rural & Isolated Practice Areas	Column 4 Sexual Health Program
Ciprofloxacin	Ear drops	Aboriginal and Torres Strait Islander persons only - Administer first dose and give remainder of manufacturer's pack	Not permitted
Clindamycin	Oral	Administer one dose and give a treatment dose as necessary	Not permitted
Clindamycin	Intravaginal	Administer first dose and give remainder of manufacturer's pack for a treatment dose	Administer one dose; give remainder of manufacturer's pack for a treatment dose
Clotrimazole	Intravaginal	Authorised under 3.2.7	Give a treatment dose as necessary
Dexamethasone 0.5 mg / framycetin sulfate 5 mg / gramicidin 0.05 mg / mL	Ear drops	Administer first dose and give remainder of manufacturer's pack	Not permitted
Diazepam #	Intravenous Oral Rectal	Administer one dose. Adults: 10 mg Children: Dose adjusted on weight per health management protocol. Consult authorised prescriber if more than recommended dose required.	Not permitted
Dicloxacillin	Oral	Administer one dose and give a treatment dose as necessary	Not permitted
Doxycycline	Oral	Administer one dose and give a treatment dose as necessary	Administer one dose and give a treatment dose as necessary
Erythromycin	Oral	Administer one dose and give a treatment dose as necessary	Not permitted
Estradiol	Intravaginal	Not permitted	Administer one dose; give remainder of manufacturer's pack
Estriol	Intravaginal	Not permitted	Administer one dose; give remainder of manufacturer's pack

Column 1 Medicine	Column 2 Approved route of administration	Restrictions/Conditions for place or circumstance	
		Column 3 Rural & Isolated Practice Areas	Column 4 Sexual Health Program
Ethinylestradiol 20 microgram/ Levonorgestrel 100 microgram †	Oral	Not permitted	Duration must not to exceed end of current prescription or 12- months since last assessment by prescriber. Maximum doses given not to exceed 4-months' supply
Ethinylestradiol 20 microgram/ Drospirinone 3mg †			
Ethinylestradiol 30 microgram/ Desogestrel 150 microgram †			
Ethinylestradiol 30 microgram/ Dienogest 2 mg †			
Ethinylestradiol 30 microgram/ Drospirinone 3 mg †			
Ethinylestradiol 30 microgram/ Gestodene 75 microgram †			
Ethinylestradiol 30 microgram/ Levonorgestrel 150 microgram †	Oral	Not permitted	Duration must not to exceed end of current prescription or 12- months since last assessment by prescriber. Maximum doses given not to exceed 4-months' supply
Ethinylestradiol 30 microgram/ Levonorgestrel 50 microgram and Ethinylestradiol 40 microgram/ Levonorgestrel 75 microgram †			
Ethinylestradiol 30 microgram/ Levonorgestrel 125 microgram †			
Ethinylestradiol 35 microgram/ Norethisterone 500 microgram †			
Ethinylestradiol 35 microgram/ Cyproterone acetate 2 mg			
Ethinylestradiol 35 microgram/ Norethisterone 1mg †			Duration must not to exceed end of current prescription or 12- months since last assessment by prescriber. Maximum doses given not to exceed 4-months' supply

Column 1 Medicine	Column 2 Approved route of administration	Restrictions/Conditions for place or circumstance	
		Column 3 Rural & Isolated Practice Areas	Column 4 Sexual Health Program
Etonogestrel 120mg / Ethinylestradiol 15 microgram † e.g. <i>Nuva Ring</i>	Intravaginal	Not permitted	Duration of must not to exceed end of current prescription or 12- months since last assessment by prescriber. Maximum given not to exceed 3- months' supply
Famciclovir	Oral	Not permitted	Administer one dose and give a treatment dose as necessary
Fentanyl #	Intramuscular Intravenous Subcutaneous	For adults only. IM or Subcutaneous - 1.5 microgram / kg to a max. of 100 microgram IV - Up to a maximum of 100 micrograms in increments, repeated every 5 to 10 minutes if required. Consult authorised prescriber if more than recommended dose required.	Not permitted
Flucloxacillin	Oral	Administer one dose and give a treatment dose as necessary	Not permitted
Fluconazole	Oral	Not permitted	Administer one dose and give a treatment dose as necessary
Flumazenil	Intravenous	Must consult authorised prescriber for an oral prescription unless circumstances do not allow, in which case, notify the authorised prescriber as soon as circumstances allow	Not permitted
Flumethasone pivalate 0.02 % / clioquinol 1 %	Ear drops	Administer one dose and give remainder of manufacturer's pack	Not permitted
Fluoride varnish	Topical to teeth	Administer only.	Not permitted
Funnel web spider anti- venom	Intravenous	Must consult authorised prescriber for an oral prescription unless circumstances do not allow, in which case, notify the authorised prescriber as soon as circumstances allow	Not permitted

Column 1 Medicine	Column 2 Approved route of administration	Restrictions/Conditions for place or circumstance	
		Column 3 Rural & Isolated Practice Areas	Column 4 Sexual Health Program
Furosemide (frusemide)	Intramuscular Intravenous	Adults Only: IM – 40 mg maximum – one dose only OR IV – 40 mg maximum - one dose only Consult authorised prescriber if more than recommended dose required.	Not permitted
Glyceryl Trinitrate Transdermal Patches	Topical	Apply one patch. Consult authorised prescriber if more than recommended dose required.	Not permitted
Haloperidol	Intravenous Intramuscular Oral	Adults only 5 mg Stat with second 5 mg dose if required to a maximum of 10 mg Consult authorised prescriber if more than recommended dose required.	Not permitted
Hydrocortisone	Intramuscular Intravenous	2 – 4 mg / kg to maximum 200 mg Consult authorised prescriber if more than recommended dose required.	Not permitted
Imiquimod	Topical	Not permitted	Apply one dose; give remainder of manufacturer's pack
Ipratropium bromide	Nebulised Metered dose inhaler (MDI)	Administer only	Not permitted
Ketorolac trometamol	Intramuscular	Adult only. Administer single dose up to 30 mg	Not permitted
Levonorgestrel 1.5 mg	Oral	Authorised under 3.2.7	Give one dose
Levonorgestrel 30 microgram † e.g. <i>Microval</i>	Oral	Not permitted	Supply not to exceed end of current prescription or 12- months since last Medical Officer/Nurse Practitioner assessment. Maximum supply not to exceed 4-months
Lidocaine (lignocaine) 1 %	Local infiltration	Administration only	Not permitted
Lidocaine (lignocaine) 1 %	Intramuscular	Administer mixed with ceftriaxone injection	Administer mixed with Ceftriaxone injection
Lidocaine (lignocaine) gel 2%	Topical	Not permitted	Administer and give remainder of manufacturer's pack if necessary

Column 1 Medicine	Column 2 Approved route of administration	Restrictions/Conditions for place or circumstance	
		Column 3 Rural & Isolated Practice Areas	Column 4 Sexual Health Program
Lidocaine (lignocaine) with adrenaline (epinephrine)	Subcutaneous	Administer - Adults and children older than 12 years only	Not permitted
Lidocaine (lignocaine)- Prilocaine 5%	Topical	Not permitted	Administer only
Lorazepam #	Oral	Adult Only 1 mg stat Must consult authorised prescriber for an oral prescription unless circumstances do not allow, in which case notify as soon as circumstances allow	Not permitted
Medroxyprogesterone acetate 150 mg/mL depo-injection †	Intramuscular	Administration not to exceed end of current prescription or 12-month period since last assessment by prescriber.	Administration not to exceed end of current prescription or 12-month period since last assessment by prescriber.
Methoxyflurane	Inhalation	Administration only - Adult and child 6 years or older: 3 mL may be repeated after 20 minutes to a maximum of 6 mL Patient must self- administer	Not permitted
Metoclopramide	Intravenous Intramuscular Oral	Adults Only: Single dose only to maximum 10 mg	Not permitted
Metronidazole	Oral	Administer one dose and give a treatment dose as necessary	Administer one dose and give a treatment dose as necessary
Miconazole	Vaginal Topical (including dermal and oral)	Authorised under 3.2.7	Administer one dose and give a treatment dose as necessary
Midazolam #	Intravenous Intramuscular Intranasal Buccal	Administer one dose. Consult authorised prescriber if more than recommended dose required.	Not permitted
Misoprostol	Rectal Sublingual Buccal	Maximum 1000 micrograms	Not permitted

Column 1 Medicine	Column 2 Approved route of administration	Restrictions/Conditions for place or circumstance	
		Column 3 Rural & Isolated Practice Areas	Column 4 Sexual Health Program
Morphine #	Intramuscular Intravenous Subcutaneous	Adult only IM / Subcutaneous: Up to a maximum of 10 mg IV: Give in increments up to a maximum of 10mg. Consult authorised prescriber if more than recommended dose required.	Not permitted
Mupirocin Cream/ointment	Topical	Administer one dose and give remainder of manufacturer's pack	Not permitted
Naloxone	Subcutaneous Intravenous Intramuscular	Authorised under 3.2.7 for treatment of opioid overdose.	Giving pack not permitted
Nifedipine	Oral	Hypertension in pregnancy – Administer one dose. Must consult authorised prescriber for an oral prescription unless circumstances do not allow, in which case notify as soon as circumstances allow	Not permitted
Nitrofurantoin	Oral	Administer one dose and give a treatment dose as necessary	Not permitted
Nitrous Oxide	Topical with cryogun	Not permitted	Administer only
Norethisterone 350 microgram † e.g. <i>Noriday</i>	Oral	Not permitted	Duration must not to exceed end of current prescription or 12- months since last assessment by prescriber. Maximum doses given not to exceed 4-months' supply
Nystatin oral drops	Topical	Authorised under 3.2.7	Administer one dose and give a treatment dose as necessary
Olanzapine	Oral	Administer one dose. Adults only Must consult authorised prescriber for an oral prescription unless circumstances do not allow, in which case notify as soon as circumstances allow	Not permitted
Ondansetron	Intravenous Oral	Children only: maximum 4 mg IV, 8 mg oral	Not permitted
Oxybuprocaine eye drop 0.4 % (minim)	Topical to eye	Administer only.	Not permitted

Column 1 Medicine	Column 2 Approved route of administration	Restrictions/Conditions for place or circumstance	
		Column 3 Rural & Isolated Practice Areas	Column 4 Sexual Health Program
Oxycodone #	Oral	Administer one dose up to 5mg. Repeat administration after four hours if needed or give one tablet as a treatment dose as necessary. Further doses must consult authorised prescriber.	Not permitted
Oxytocin	Intramuscular Intravenous	Administer only. 5 to 10 units per dose stat or 30 units for IV infusion Consult authorised prescriber if more than recommended dose required	Not permitted
Phenoxymethylpenicillin	Oral	Administer one dose and give a treatment dose as necessary	Not permitted
Podophyllotoxin	Topical	Administer and give a course. Treat with local application twice daily to a maximum of 6 weeks	Administer and give a course. Treat with local application twice daily to a maximum of 6 weeks
Prednisolone	Oral	Administer for treatment of acute asthma for adult and child 6 years and older	Not permitted
Probenecid	Oral	Administer only	Not permitted
Procaine benzylpenicillin (procaine penicillin)	Intramuscular	Administer one dose daily for a maximum of three days	Administer one dose
Promethazine	Intramuscular Intravenous	Adults and children older than 12 years: Maximum 50 mg – one dose only Children older than two years: Maximum 25 mg – one dose only Consult authorised prescriber if more than recommended dose required.	Not permitted
Roxithromycin	Oral	Administer one dose and give a treatment dose as necessary	Administer one dose and give a treatment dose as necessary
Salbutamol	Nebulised	Administer only	Not permitted
Silver sulfadiazine 1% cream	Topical	Administer first dose and give remainder of manufacturer's pack	Not permitted
Snake polyvalent anti-venom	Intravenous	One ampoule (40,000 units) Consult authorised prescriber if more than recommended dose required.	Not permitted

Column 1 Medicine	Column 2 Approved route of administration	Restrictions/Conditions for place or circumstance	
		Column 3 Rural & Isolated Practice Areas	Column 4 Sexual Health Program
Triamcinolone compound Otic	Topical to ear	Administer first dose and give remainder of manufacturer's pack	Not permitted
Trimethoprim	Oral	Administer one dose and give a treatment dose as necessary	Administer one dose and give a treatment dose as necessary
Trimethoprim/ Sulfamethoxazole	Oral	Administer one dose and give a treatment dose as necessary	Not permitted
Valaciclovir	Oral	Administer one dose and give a treatment dose as necessary	Administer one dose and give a treatment dose as necessary

Appendix 4 – Queensland Immunisation Program medicines

Column 1 Antigens	Column 2 Restrictions/Conditions for Sexual Health Program
Diphtheria	Not permitted
<i>Haemophilus influenzae</i> type b	Not permitted
Hepatitis A	Administer only
Hepatitis B	Administer only
Human Papillomavirus	Administer only
Influenza	Not permitted
Measles	Administer only if combined with mumps and rubella antigens
Mumps	Administer only if combined with measles and rubella antigens
Meningococcal (ACWY)	Not permitted
Meningococcal B	Not permitted
Meningococcal C	Not permitted
Pertussis	Not permitted
Pneumococcal	Not permitted
Poliomyelitis	Not permitted
Rotavirus	Not permitted
Rubella	Administer only if combined with measles and mumps antigens
Tetanus	Not permitted
Varicella (chickenpox)	Not permitted
Zoster (herpes zoster)	Not permitted