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Schedule—Consequential Amendments

Notes
An Act relating to therapeutic goods

Chapter 1—Preliminary

1 Short title [see Note 1]

This Act may be cited as the Therapeutic Goods Act 1989.

2 Commencement [see Note 1]

This Act commences on the day after the day on which a House of the Parliament approves regulations made under this Act in the same form as approved by the other House, provided that:

(a) not more than 90 days have elapsed; and
(b) the places of Senators have not become vacant under section 13 of the Constitution; and
(c) a dissolution or expiration of the House of Representatives has not occurred;

between the approval of one House and the approval of the other House.

3 Interpretation

(1) In this Act, unless the contrary intention appears:

accessory means an article that its manufacturer specifically intended to be used together with a medical device to enable the device to be used as the manufacturer of the device intended.

actual or potential tampering has the meaning given by section 42U.

advertisement, in relation to therapeutic goods, includes any statement, pictorial representation or design, however made, that is intended, whether directly or indirectly, to promote the use or supply of the goods.
Section 3

application audit assessment fee means a fee payable under subsection 41LA(3).

assessment fee means:
(a) a conformity assessment fee; or
(b) an application audit assessment fee; payable under Part 4-10.

authorised person means:
(a) in relation to any provision of this Act, a person authorised by the Secretary to exercise powers under that provision; or
(b) in relation to a provision of Part 6-2, a member of the Australian Federal Police, or a Customs officer exercising powers in a Customs place (within the meaning of section 183UA of the Customs Act 1901).

batch means a quantity of a product that is:
(a) uniform in composition, method of manufacture and probability of chemical or microbial contamination; and
(b) made in one cycle of manufacture and, in the case of a product that is sterilised or freeze dried, sterilised or freeze dried in one cycle.

bioburden, in relation to therapeutic goods, means the quantity and characteristics of microorganisms present in the goods or to which the goods may be exposed in a manufacturing environment.

British Pharmacopoeia means the edition of the book of that name, including any additions or amendments, that was in effect for the purposes of the Therapeutic Goods Act 1966 immediately before the commencement of this section and, if additions or amendments of that book are made after that commencement, or new editions of that book are published after that commencement, includes those additions or amendments, or those new editions, from a day specified by the Minister by order published in the Gazette.

British Pharmacopoeia (Veterinary) means the latest edition of the book of that name, including any additions or amendments, published on the recommendation of the Medicines Commission of the United Kingdom immediately before the commencement of this section and, if additions or amendments of that book are made
after that commencement, or new editions of that book are published after that commencement, includes those additions or amendments, or those new editions, from a day specified by the Minister by order published in the Gazette.

**Commonwealth authority** includes:
(a) a body corporate, or an unincorporated body, established for a public purpose by or under an Act; and
(b) a tribunal or authority established by or in accordance with an Act.

**Commonwealth officer** includes:
(a) a Minister; and
(b) a person holding:
   (i) an office established by or under an Act; or
   (ii) an appointment made under an Act; or
   (iii) an appointment made by the Governor-General or a Minister but not under an Act; and
(c) a person who is a member or officer of a Commonwealth authority; and
(d) a person who is in the service or employment of the Commonwealth, or of a Commonwealth authority, or is employed or engaged under an Act or regulations made under an Act.

**composite pack** has the meaning given by subsection 7B(2).

**conformity assessment certificate** means a certificate issued under section 41EE.

**conformity assessment fee** means a fee payable under subsection 41LA(1).

**conformity assessment procedures** has the meaning given by section 41DA.

**conformity assessment standard** means a conformity assessment standard specified in an order under section 41DC.

**container**, in relation to therapeutic goods, means the vessel, bottle, tube, ampoule, syringe, vial, sachet, strip pack, blister pack,
wrapper, cover or other similar article that immediately covers the goods, but does not include an article intended for ingestion.

**corporation** means a body corporate that is:
(a) a foreign corporation; or
(b) a trading corporation formed within the limits of the Commonwealth or a financial corporation so formed.

**corresponding State law** means a State law declared by the regulations to correspond to this Act or the regulations, including such a law as amended from time to time.

**counterfeit** has the meaning given by section 42E.

**current Poisons Standard** has the meaning given by section 52A.

**Customs officer** means an officer of Customs within the meaning of the *Customs Act 1901*.

**data processing device** means any article or material (for example, a disc) from which information is capable of being reproduced with or without the aid of any other article or device.

**device number**, in relation to a medical device, means any combination of numbers, symbols and letters assigned to the device under section 41FL.

**directions for use**, in relation to therapeutic goods, includes information on:
(a) appropriate doses of the goods; and
(b) the method of administration or use of the goods; and
(c) the frequency and duration of treatment for each indication of the goods; and
(d) the use of the goods by persons of particular ages or by persons having particular medical conditions.

**EC/EFTA attestation of conformity** means an attestation of conformity (within the meaning of the EC Mutual Recognition Agreement or the EFTA Mutual Recognition Agreement) issued by an EC/EFTA conformity assessment body that is approved by the Secretary in writing.
EC/EFTA conformity assessment body means a Conformity Assessment Body designated in one of the following Sectoral Annexes to the EC Mutual Recognition Agreement or the EFTA Mutual Recognition Agreement:
(a) Sectoral Annex (Medical Devices);
(b) Sectoral Annex (Medicinal Products GMP Inspection and Batch Certification).

EC Mutual Recognition Agreement means the Agreement on Mutual Recognition in relation to Conformity Assessment, Certificates and Markings between Australia and the European Community, as in force from time to time.

EFTA Mutual Recognition Agreement means the Agreement on Mutual Recognition in relation to Conformity Assessment, Certificates and Markings between Australia and the European Free Trade Association, as in force from time to time.

essential principles has the meaning given by section 41CA.

ethics committee means a committee:
(a) constituted and operating as an ethics committee in accordance with guidelines issued by the National Health and Medical Research Council as in force from time to time; and
(b) which has notified its existence to the Australian Health Ethics Committee established under the National Health and Medical Research Council Act 1992.

exempt device means a medical device that is of a kind that is exempted from Division 3 of Part 4-11 by the regulations.

exempt goods, in relation to a provision of Part 3-2, means therapeutic goods that are exempted from the operation of that Part (except section 31A and sections 31C to 31F) by the regulations.

exempt goods, in relation to a provision of Part 3-3, means therapeutic goods that are exempted from the operation of that Part by the regulations.

exempt person, in relation to therapeutic goods, means a person exempted from the operation of Part 3-3 in relation to those goods by the regulations.
Section 3

**export only medicine** means a medicine that:
(a) is manufactured in Australia for export only, or imported into
Australia for export only; and
(b) is listable goods only because it is so manufactured or
imported (and not for any other reason).

**financial corporation** means a financial corporation within the
meaning of paragraph 51(xx) of the Constitution.

**first Poisons Standard** has the meaning given by section 52A.

**foreign corporation** means a foreign corporation within the
meaning of paragraph 51(xx) of the Constitution.

**gazetted kits group** means a group of kits identified in an order in
force under subsection 16(3A).

**gazetted therapeutic devices group** has the meaning given by
subsection 16(3).

**gazetted therapeutic goods group** has the meaning given by
subsection 16(2).

**Gene Technology Regulator** has the same meaning as in the *Gene
Technology Act 2000*.

**GM product** has the same meaning as in the *Gene Technology Act
2000*.

**grouped therapeutic goods** means therapeutic goods included in:
(a) a gazetted therapeutic goods group; or
(b) a gazetted therapeutic devices group; or
(c) a gazetted kits group.

**included in the Register**, in relation to a medical device to which
Chapter 4 applies, means included in the Register under Chapter 4.

Note: For medical devices to which Chapter 4 applies, see section 41BJ.

**indications**, in relation to therapeutic goods, means the specific
therapeutic uses of the goods.

**international instrument** means:
(a) any treaty, convention, protocol, agreement or other instrument that is binding in international law; and
(b) a part of such a treaty, convention, protocol, agreement or other instrument.

*kind*, in relation to a medical device, has the meaning given by section 41BE.

*label*, in relation to therapeutic goods, means a display of printed information:
(a) on or attached to the goods; or
(b) on or attached to a container or primary pack in which the goods are supplied; or
(c) supplied with such a container or pack.

*licence* means a licence under Part 3-3.

*lizable devices* means therapeutic devices that are required to be included in the part of the Register for listed goods.

*lizable goods* means therapeutic goods that are required:
(a) under the regulations; or
(b) by a notice published in the *Gazette* under subsection 17(5); to be included in the part of the Register relating to listed goods.

*listed goods* means therapeutic goods that are included in the Part of the Register for goods known as listed goods.

*listing number*, in relation to listed goods, means any combination of numbers, symbols and letters assigned to the goods under section 27.

*manufacture*, in relation to therapeutic goods that are not medical devices, means:
(a) to produce the goods; or
(b) to engage in any part of the process of producing the goods or of bringing the goods to their final state, including engaging in the processing, assembling, packaging, labelling, storage, sterilising, testing or releasing for supply of the goods or of any component or ingredient of the goods as part of that process.
Section 3

**manufacturer.** of a medical device, has the meaning given by section 41BG.

**manufacturing premises** means premises (including premises that comprise 2 or more sites):

(a) that are for use in the manufacture of a particular kind of therapeutic goods; and

(b) at which the same persons have control of the management of the production of the goods and the procedures for quality control.

**manufacturing principles** means the principles for the time being having effect under section 36.

**medical device** has the meaning given by section 41BD.

**medical device classification** means a classification specified in the regulations made for the purposes of section 41DB.

**medical device standard,** in relation to a kind of medical device, means a medical device standard, specified in an order under section 41CB, that is applicable to that kind of medical device.

**medicine** means:

(a) therapeutic goods that are represented to achieve, or are likely to achieve, their principal intended action by pharmacological, chemical, immunological or metabolic means in or on the body of a human or animal; and

(b) any other therapeutic goods declared by the Secretary, for the purpose of the definition of **therapeutic device,** not to be therapeutic devices.

**member of EFTA** means a country declared by the Minister under section 3A to be a member of the European Free Trade Association.

**member of the European Community** means a country declared by the Minister under section 3A to be a member of the European Community.

National Manager of the Therapeutic Goods Administration means:
(a) the person holding the position of National Manager of the Therapeutic Goods Administration; or
(b) if the position of National Manager of the Therapeutic Goods Administration ceases to exist, or ceases to be referred to by that name—the person holding a position determined in writing by the Secretary.

non-EC/EFTA attestation of conformity, for a non-EC/EFTA MRA, means an attestation of conformity issued, after the non-EC/EFTA MRA has come into force, by a conformity assessment body that is designated in the non-EC/EFTA MRA and approved by the Secretary in writing for the non-EC/EFTA MRA.

non-EC/EFTA MRA means an international instrument that Australia is bound by, or is a party to, if:
(a) a purpose of the instrument is the recognition of attestations of conformity; and
(b) the instrument satisfies the requirements (if any) set out in regulations made for the purposes of this paragraph; but does not include:
(c) the EC Mutual Recognition Agreement; or
(d) the EFTA Mutual Recognition Agreement.

poison means an ingredient, compound, material or preparation which, or the use of which, may cause death, illness or injury and includes any ingredient, compound, material or preparation referred to in a schedule to the current Poisons Standard.

premises includes:
(a) a structure, building, aircraft, vehicle or vessel; and
(b) a place (whether enclosed or built upon or not); and
(c) a part of a thing referred to in paragraph (a) or (b).

presentation, in relation to therapeutic goods, means the way in which the goods are presented for supply, and includes matters relating to the name of the goods, the labelling and packaging of the goods and any advertising or other informational material associated with the goods.
Section 3

primary pack, in relation to therapeutic goods, means the complete pack in which the goods, or the goods and their container, are to be supplied to consumers.

protected information, in relation to therapeutic goods, has the meaning given by section 25A.

quality, in relation to therapeutic goods, includes the composition, strength, potency, stability, sterility, purity, bioburden, design, construction and performance characteristics of the goods.

refurbishment has the meaning given by the regulations.

Register means the Australian Register of Therapeutic Goods maintained under section 9A.

registered goods means therapeutic goods included in the part of the Register for goods known as registered goods.

registration number, in relation to registered goods, means any combination of numbers, symbols and letters assigned to the goods under section 27.

restricted goods means medicines (including progesterone antagonists and vaccines against human chorionic gonadotrophin) intended for use in women as abortifacients.

scheduling has the meaning given by section 52A.

Secretary means the Secretary to the Department.

sponsor, in relation to therapeutic goods, means:

(a) a person who exports, or arranges the exportation of, the goods from Australia; or
(b) a person who imports, or arranges the importation of, the goods into Australia; or
(c) a person who, in Australia, manufactures the goods, or arranges for another person to manufacture the goods, for supply (whether in Australia or elsewhere);

but does not include a person who:

(d) exports, imports or manufactures the goods; or
(e) arranges the exportation, importation or manufacture of the goods;
on behalf of another person who, at the time of the exportation, importation, manufacture or arrangements, is a resident of, or is carrying on business in, Australia.

**standard**. in relation to therapeutic goods, means a standard that:
(a) is specified in an order under section 10 that is applicable to the goods; or
(b) if no such order is applicable to the goods but the goods are the subject of a monograph in:
   (i) in the case of goods for use in humans—the British Pharmacopoeia; or
   (ii) in the case of goods for use in animals—the British Pharmacopoeia (Veterinary);
   is constituted by the statements in that monograph.

**State** includes the Australian Capital Territory and the Northern Territory.

**State law** means a law of a State, of the Australian Capital Territory or of the Northern Territory.

**supply** includes:
(a) supply by way of sale, exchange, gift, lease, loan, hire or hire-purchase; and
(b) supply, whether free of charge or otherwise, by way of sample or advertisement; and
(c) supply, whether free of charge or otherwise, in the course of testing the safety or efficacy of therapeutic goods in persons or animals; and
(d) supply by way of administration to, or application in the treatment of, a person or animal.

**system or procedure pack** has the meaning given by section 41BF.

**tamper**: therapeutic goods are tampered with if:
(a) they are interfered with in a way that affects, or could affect, the quality, safety or efficacy of the goods; and
(b) the interference has the potential to cause, or is done for the purpose of causing, injury or harm to any person.
Chapter 1  Preliminary

Section 3

*therapeutic device* means therapeutic goods consisting of an instrument, apparatus, appliance, material or other article (whether for use alone or in combination), together with any accessories or software required for its proper functioning, which does not achieve its principal intended action by pharmacological, chemical, immunological or metabolic means though it may be assisted in its function by such means, but the expression does not include therapeutic goods declared by the Secretary, by order published in the *Gazette*, not to be therapeutic devices.

*therapeutic goods* means goods:

(a) that are represented in any way to be, or that are, whether because of the way in which the goods are presented or for any other reason, likely to be taken to be:

   (i) for therapeutic use; or

   (ii) for use as an ingredient or component in the manufacture of therapeutic goods; or

   (iii) for use as a container or part of a container for goods of the kind referred to in subparagraph (i) or (ii); or

(b) included in a class of goods the sole or principal use of which is, or ordinarily is, a therapeutic use or a use of a kind referred to in subparagraph (a)(ii) or (iii);

and includes medical devices and goods declared to be therapeutic goods under an order in force under section 7, but does not include:

(c) goods declared not to be therapeutic goods under an order in force under section 7; or

(d) goods in respect of which such an order is in force, being an order that declares the goods not to be therapeutic goods when used, advertised, or presented for supply in the way specified in the order where the goods are used, advertised, or presented for supply in that way; or

(e) goods (other than goods declared to be therapeutic goods under an order in force under section 7) for which there is a prescribed standard in the Australia New Zealand Food Standards Code as defined in subsection 3(1) of the *Australia New Zealand Food Authority Act 1991*; or

(f) goods which, in Australia or New Zealand, have a tradition of use as foods for humans in the form in which they are presented.
Therapeutic Goods Advertising Code means the Code known as the Therapeutic Goods Advertising Code notified in the Gazette with effect from the date of commencement of Schedule 1 to the Therapeutic Goods Amendment Act (No. 1) 2003 together with any amendments of the Code published by the Minister in the Gazette from time to time.

therapeutic use means use in or in connection with:
(a) preventing, diagnosing, curing or alleviating a disease, ailment, defect or injury in persons or animals; or
(b) influencing, inhibiting or modifying a physiological process in persons or animals; or
(c) testing the susceptibility of persons or animals to a disease or ailment; or
(d) influencing, controlling or preventing conception in persons; or
(e) testing for pregnancy in persons; or
(f) the replacement or modification of parts of the anatomy in persons or animals.

trading corporation means a trading corporation within the meaning of paragraph 51(xx) of the Constitution.

working day, for a person, means any day except:
(a) Saturday or Sunday; or
(b) a day that is a public holiday in the State or Territory in which the person is located.

(2) For the purposes of this Act:
(a) therapeutic goods are to be taken to be for use in animals if:
   (i) the goods bear a name or description that indicates, or is likely to give the impression, that the goods are intended for use in animals and are not intended for use in humans; or
   (ii) the goods are otherwise represented, or otherwise purport, to be intended for use in animals and not intended for use in humans; and
(b) therapeutic goods are to be taken to be for use in humans if they are not solely for use in animals.
Section 3

(3) The Secretary must, at least once in each year, cause to be published in the Gazette a list of the names of all persons, other than members of the Australian Federal Police, who are, at the time of publication, authorised persons.

(4) The provisions of this Act are in addition to, and not in substitution for, the provisions of any other Act that relate to therapeutic goods.

(5) For the purposes of this Act, the presentation of therapeutic goods is unacceptable if it is capable of being misleading or confusing as to the content or proper use or identification of the goods and, without limiting the previous words in this subsection, the presentation of therapeutic goods is unacceptable:

(a) if it states or suggests that the goods have ingredients, components or characteristics that they do not have; or

(b) if a name applied to the goods is the same as the name applied to other therapeutic goods that are supplied in Australia where those other goods contain additional or different therapeutically active ingredients; or

(c) if the label of the goods does not declare the presence of a therapeutically active ingredient; or

(d) if a form of presentation of the goods may lead to unsafe use of the goods or suggests a purpose that is not in accordance with conditions applicable to the supply of the goods in Australia; or

(e) in prescribed cases.

(6) A reference in this Act to an annual registration charge, an annual listing charge, an annual charge for inclusion in the Register or an annual licensing charge is a reference to such a charge imposed under the Therapeutic Goods (Charges) Act 1989.

(7) A reference to an offence against this Act includes a reference to:

(a) an offence against the regulations; and

(b) an offence against section 6 of the Crimes Act 1914, or section 11.1, 11.4 or 11.5 of the Criminal Code, in relation to an offence against this Act or the regulations; and

(c) an offence against section 136.1, 137.1 or 137.2 of the Criminal Code in relation to this Act or the regulations.
(7A) For the purposes of this Act, a corresponding State law imposes a duty on a Commonwealth officer or Commonwealth authority if:
   (a) the corresponding State law confers a function or power on the officer or authority; and
   (b) the circumstances in which the function or power is conferred give rise to an obligation on the officer or authority to perform the function or to exercise the power.

(8) A maximum penalty specified:
   (a) at the foot of a section of this Act (other than a section that is divided into subsections); or
   (b) at the foot of a subsection of this Act;
indicates that a person who contravenes the section or subsection is guilty of an offence against the section or subsection and is punishable, on conviction, by a penalty up to that maximum.

3A Declaration—member of European Community

(1) The Minister may declare, in writing, that a country specified in the declaration is a member of:
   (a) the European Community; or
   (b) the European Free Trade Association.

(2) A declaration under subsection (1) must be published in the Gazette.

3B Declaration—country covered by non-EC/EFTA MRA

(1) The Minister may declare, in writing, that a country specified in the declaration is covered by the non-EC/EFTA MRA specified in the declaration.

(2) A declaration under subsection (1) must be published in the Gazette.

4 Objects of Act

(1) The objects of this Act are to do the following, so far as the Constitution permits:
Section 5

(a) provide for the establishment and maintenance of a national system of controls relating to the quality, safety, efficacy and timely availability of therapeutic goods that are:
   (i) used in Australia, whether produced in Australia or elsewhere; or
   (ii) exported from Australia;
(b) to provide a framework for the States and Territories to adopt a uniform approach to control the availability and accessibility, and ensure the safe handling, of poisons in Australia.

(1A) The reference in paragraph (1)(a) to the efficacy of therapeutic goods is a reference, if the goods are medical devices, to the performance of the devices as the manufacturer intended.

(2) This Act is therefore not intended to apply to the exclusion of a law of a State, of the Australian Capital Territory or of the Northern Territory to the extent that the law is capable of operating concurrently with this Act.

5 Act to bind Crown

This Act binds the Crown in right of the Commonwealth, of each of the States, of the Australian Capital Territory and of the Northern Territory, but nothing in this Act renders the Crown liable to be prosecuted for an offence.

5A Application of the Criminal Code

Chapter 2 (other than Part 2.5) of the Criminal Code applies to all offences against this Act.

Note: Chapter 2 of the Criminal Code sets out the general principles of criminal responsibility.

6 Operation of Act

(1) This Act applies to:
   (a) things done by corporations; and

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Section 6AA

(b) things done by natural persons or corporations in so far as those things are done:
   (i) in the course of, or in preparation for, trade or commerce between Australia and a place outside Australia, among the States, between a State and a Territory or between 2 Territories; or
   (ii) under a law of the Commonwealth relating to the provision of pharmaceutical or repatriation benefits; or
   (iii) in relation to the Commonwealth or in relation to an authority of the Commonwealth.

(2) Without limiting the effect of this Act apart from this subsection, this Act also has the effect it would have if the reference in paragraph (1)(a) to things done by corporations were confined to things done by trading corporations for the purposes of their trading activities.

6AA Importation of restricted goods

(1) In spite of any other provision of this Act, a person must not, without the written approval of the Minister, import any restricted goods into Australia.

Penalty: 300 penalty units.

(2) A written approval may be given:
   (a) unconditionally or subject to conditions; or
   (b) in respect of particular restricted goods or classes of restricted goods.

(3) A person is guilty of an offence if:
   (a) the person engages in conduct; and
   (b) the conduct contravenes a condition of an approval.

Penalty: 200 penalty units.

(3A) In subsection (3):

**engage in conduct** means:
   (a) do an act; or
   (b) omit to perform an act.
Section 6AB

(4) A written approval shall be laid before each House of the Parliament by the Minister within 5 sitting days of being given.

(5) Unless:
   (a) a written approval is in effect; and
   (b) the Minister has notified the Chief Executive Officer of Customs in writing of the approval;

restricted goods are for the purposes of the *Customs Act 1901* taken to be prohibited imports.

6AB Exempt goods

Regulations exempting restricted goods from the operation of a Part of this Act must not take effect before the expiration of the time within which a House of the Parliament may disallow the regulations.

6AAA Commonwealth consent to conferral of functions etc. on its officers and authorities by corresponding State laws

(1) A corresponding State law may confer functions or powers, or impose duties, on:
   (a) a Commonwealth officer; or
   (b) a Commonwealth authority.

(2) Subsection (1) does not authorise the conferral of a function or power, or the imposition of a duty, by a corresponding State law to the extent to which:
   (a) the conferral or imposition, or the authorisation, would contravene any constitutional doctrines restricting the duties that may be imposed on Commonwealth officers or Commonwealth authorities; or
   (b) the authorisation would otherwise exceed the legislative power of the Commonwealth.

(3) Subsection (1) does not extend to a function, power or duty of a kind specified in regulations made for the purposes of this subsection.

(4) This Act is not intended to exclude or limit the operation of a corresponding State law that confers any functions or powers, or
imposes any duties, on a Commonwealth officer or Commonwealth authority to the extent to which that law:
(a) is consistent with subsections (1) to (3); and
(b) is capable of operating concurrently with this Act.

6AAB When duty imposed

Application

(1) This section applies if a corresponding State law purports to impose a duty on a Commonwealth officer or Commonwealth authority.

State legislative power sufficient to support duty

(2) The duty is taken not to be imposed by this Act (or any other law of the Commonwealth) to the extent to which:
(a) imposing the duty is within the legislative powers of the State concerned; and
(b) imposing the duty by the corresponding State law is consistent with the constitutional doctrines restricting the duties that may be imposed on a Commonwealth officer or Commonwealth authority.

Note: If this subsection applies, the duty will be taken to be imposed by force of the corresponding State law (the Commonwealth having consented under section 6AAA to the imposition of the duty by the corresponding State law).

Commonwealth legislative power sufficient to support duty but State legislative powers are not

(3) If, to ensure the validity of the purported imposition of the duty, it is necessary that the duty be imposed by a law of the Commonwealth (rather than by force of the corresponding State law), the duty is taken to be imposed by this Act to the extent necessary to ensure that validity.

(4) If, because of subsection (3), this Act is taken to impose the duty, it is the intention of the Parliament to rely on all powers available to it under the Constitution to support the imposition of the duty by this Act.
(5) The duty is taken to be imposed by this Act in accordance with subsection (3) only to the extent to which imposing the duty:
(a) is within the legislative powers of the Commonwealth; and
(b) is consistent with the constitutional doctrines restricting the duties that may be imposed on a Commonwealth officer or Commonwealth authority.

(6) To avoid doubt, neither this Act (nor any other law of the Commonwealth) imposes a duty on the Commonwealth officer or Commonwealth authority to the extent to which imposing such a duty would:
(a) contravene any constitutional doctrine restricting the duties that may be imposed on a Commonwealth officer or Commonwealth authority; or
(b) otherwise exceed the legislative power of the Commonwealth.

(7) Subsections (1) to (6) do not limit section 6AAA.

6AAC Imposing duty under State law

(1) This section:
(a) applies only for the purposes of the application of the provisions of this Act or another law of the Commonwealth (with or without modification) as a law of a State by a provision of a corresponding State law; and
(b) does not apply for those purposes if the corresponding State law otherwise provides.

(2) If the corresponding State law purports to impose a duty on a Commonwealth officer or Commonwealth authority to do a particular thing, the duty is taken to be imposed by the corresponding State law to the extent to which imposing the duty:
(a) is within the legislative powers of the State; and
(b) is consistent with the constitutional doctrines restricting the duties that may be imposed on a Commonwealth officer or Commonwealth authority.

(3) To avoid doubt, the corresponding State law does not impose the duty on the Commonwealth officer or Commonwealth authority to the extent to which imposing the duty would:
(a) contravene any constitutional doctrine restricting the duties that may be imposed on a Commonwealth officer or Commonwealth authority; or
(b) otherwise exceed the legislative powers of the State.

(4) If imposing on the Commonwealth officer or Commonwealth authority the duty to do that thing would:
   (a) contravene any constitutional doctrine restricting the duties that may be imposed on a Commonwealth officer or Commonwealth authority; or
   (b) otherwise exceed the legislative powers of both the State and the Commonwealth;
the corresponding State law is taken instead to confer on the officer or authority a power to do that thing at the discretion of the officer or authority.

6AAD Conferral of jurisdiction on federal courts

If:
   (a) a provision of a corresponding State law purports to apply a provision of a law of the Commonwealth (the applied provision) as a law of the State; and
   (b) the applied provision purports to confer jurisdiction in relation to a matter on a federal court;
the jurisdiction in relation to that matter is taken to be conferred on the court by this section.

6AAE Consequences of State law conferring duty, function or power on Commonwealth officer or Commonwealth authority

(1) If a corresponding State law confers on a Commonwealth officer or Commonwealth authority:
   (a) the function of including goods in the Register; or
   (b) the power to include goods in the Register;
the officer or authority may include the goods in the Register in accordance with the corresponding State law.

(2) If a corresponding State law authorises or requires a Commonwealth officer or Commonwealth authority to cancel the inclusion of goods in the Register, the officer or authority may
Section 6B

cancel the inclusion of the goods in the Register in accordance with the corresponding State law.

(3) The inclusion of goods in the Register under subsection (1) does not subject any person to any liability whatever under this Act, except a liability under Part 6-1.

(4) A Commonwealth officer or Commonwealth authority may make any notations in the Register that the officer or authority considers necessary to identify entries that relate to goods included in the Register under subsection (1).

(5) Goods may be included in the Register under subsection (1) even though the same goods have already been included in the Register under another provision of this Act.

(6) A reference in this section to the inclusion of goods in the Register is a reference to the inclusion of the goods:
   (a) in the part of the Register for goods known as registered goods; or
   (b) in the part of the Register for goods known as listed goods; or
   (c) in the part of the Register for medical devices included under Chapter 4.

6B Review of certain decisions under State laws

(1) Application may be made to the Administrative Appeals Tribunal for review of a reviewable State decision.

(2) A decision made by the Secretary in the performance of a function, or the exercise of a power, conferred by a corresponding State law is a reviewable State decision for the purpose of this section if:
   (a) the law under which the decision was made provides for review by the Administrative Appeals Tribunal; and
   (b) the decision is declared by the regulations to be a reviewable decision for the purposes of this section.

(3) For the purposes of subsection (1), the Administrative Appeals Tribunal Act 1975 has effect as if a corresponding State law were an enactment.
Section 6C

6C Fees payable to Commonwealth under State laws

(1) This section applies to fees payable to the Commonwealth under a State law in respect of the performance or exercise of functions or powers conferred by that law on the Secretary.

(2) The Secretary may make arrangements with the appropriate authority of a State, of the Australian Capital Territory or of the Northern Territory in relation to the payment to the Commonwealth of fees to which this section applies.

7 Declaration that goods are/are not therapeutic goods

(1) Where the Secretary is satisfied that particular goods or classes of goods:
   (a) are or are not therapeutic goods; or
   (b) when used, advertised, or presented for supply in a particular way, are or are not therapeutic goods;
the Secretary may, by order published in the Gazette, declare that the goods, or the goods when used, advertised, or presented for supply in that way, are or are not, for the purposes of this Act, therapeutic goods.

(2) The Secretary may exercise his or her powers under this section of his or her own motion or following an application made in writing to the Secretary.

(3) A declaration under this section takes effect on the day on which the declaration is published in the Gazette or on such later day as is specified in the order.

(4) If a declaration under this section:
   (a) is a declaration that particular goods or classes of goods are not therapeutic goods; and
   (b) applies wholly or partly to goods that, apart from this section, would be medical devices;
the goods are not medical devices, or are not medical devices when used, advertised, or presented for supply in the way specified in the declaration.
Section 7A

7A Authorised persons

The Secretary may, in writing, authorise any of the following persons to exercise powers under a specified provision of this Act:

(a) an officer of the Department, of another Department or of an authority of the Commonwealth;

(b) an officer of:

(i) a Department of State of a State; or

(ii) a Department or administrative unit of the Public Service of a Territory; or

(iii) an authority of a State or of a Territory;

being a Department, unit or authority that has functions relating to health matters or law enforcement matters.

7B Kits

(1) A package and therapeutic goods in the package together constitute a kit for the purposes of this Act if:

(a) the package and the therapeutic goods are for use as a unit;

and

(b) each item of the therapeutic goods consists of goods that are registered or listed or are exempt goods in relation to Part 3-2; and

(c) the package and therapeutic goods do not constitute a composite pack or a system or procedure pack.

(2) A package and therapeutic goods in the package together constitute a composite pack if:

(a) the therapeutic goods are of 2 or more kinds; and

(b) the package does not contain any medical devices or therapeutic devices; and

(c) the therapeutic goods are for administration as a single treatment or as a single course of treatment; and

(d) it is necessary that the therapeutic goods be combined before administration or that they be administered in a particular sequence.

(3) To avoid doubt, it is declared that a kit constitutes therapeutic goods.
8 Power to obtain information with respect to therapeutic goods

(1) The Secretary may, by notice in writing given to a person who has imported into Australia or has supplied in Australia:
   (a) therapeutic goods; or
   (b) goods in relation to which the Secretary is considering making a declaration under section 7;
request the person to give to an officer of the Department identified in the notice, within such reasonable period as is specified in the notice, information required by the notice concerning the composition, indications, directions for use or labelling of the goods or concerning advertising material relating to the goods.

(1A) A notice under subsection (1) may require the information to be given:
   (a) in writing; or
   (b) in accordance with specified software requirements:
      (i) on a specified kind of data processing device; or
      (ii) by way of a specified kind of electronic transmission.

(2) A person must not fail to comply with a notice given to the person under this section.
   Maximum penalty: 60 penalty units.

(3) Subsection (2) does not apply if the person has a reasonable excuse.

Note: The defendant bears an evidential burden in relation to the matter in subsection (3). See subsection 13.3(3) of the Criminal Code.

(4) An offence under subsection (2) is an offence of strict liability.

Note: For strict liability, see section 6.1 of the Criminal Code.

9 Arrangements with States etc.

(1) The Minister may make arrangements with the appropriate Minister of a State, of the Australian Capital Territory or of the Northern Territory for the carrying out by that State or Territory, on behalf of the Commonwealth, of:
   (a) the evaluation of therapeutic goods for registration; or
   (b) the inspection of manufacturers of therapeutic goods; or
Section 9

(c) other functions under this Act or the regulations.

(2) An arrangement under this section may provide for the payment to a State or Territory of amounts in respect of the performance of functions under the arrangement.
Chapter 2—Australian Register of Therapeutic Goods

9A Australian Register of Therapeutic Goods

(1) The Secretary is to cause to be maintained a register, to be known as the Australian Register of Therapeutic Goods, for the purpose of compiling information in relation to, and providing for evaluation of, therapeutic goods for use in humans.

(2) Subject to subsection (3), the Register is to be kept in such form as the Secretary determines.

(3) The Register is to contain these 3 parts:

(a) a part for goods to be known as registered goods; and
(b) a part for goods to be known as listed goods; and
(c) a part for medical devices included in the Register under Chapter 4.

(4) The regulations may prescribe:

(a) the therapeutic goods, or the classes of therapeutic goods, that are required to be included in each part of the Register; and

(b) the ways in which:

(i) goods that are included in the part of the Register relating to registered goods may be transferred, or may be required to be transferred, to the part of the Register for listed goods; and

(ii) goods that are included in the part of the Register relating to listed goods may be transferred, or may be required to be transferred, to the part of the Register for registered goods; and

(c) the ways in which goods that have been assigned a registration or listing number may be assigned a different registration or listing number; and

(d) the ways in which medical devices that have been assigned a device number may be assigned a different device number.
Section 9B

(5) The Minister may, by notice published in the Gazette:
   (a) require that specified therapeutic goods be included in the part of the Register for listed goods; and
   (b) specify the conditions subject to which such goods may be included in that part of the Register.

(6) If the regulations are amended to require any of those goods to be included in the part of the Register for listed or registered goods, then the Gazette notice ceases to have effect in respect of the goods included in the regulations.

9B When registrations or listings of medical devices are taken to be cancelled

(1) The registration or listing of a medical device to which subsection 15A(5) applied is taken to be cancelled:
   (a) on the second anniversary of the day on which Chapter 4 commences; or
   (b) if the medical device is of a kind included in the Register under Chapter 4 before that second anniversary—when that inclusion takes effect.

(2) The registration or listing of any other medical device is taken to be cancelled:
   (a) on the fifth anniversary of the day on which Chapter 4 commences; or
   (b) if the medical device is of a kind included in the Register under Chapter 4 before that fifth anniversary—when that inclusion takes effect.

(3) This section does not prevent the Secretary from taking action under section 30.

9C Inspection of entries in Register

(1) The Register is not open for public inspection, but a person in relation to whom therapeutic goods are entered on the Register may make a written request to the Secretary for a copy of the entry in the Register in relation to the goods.
Section 9D

(2) If the person makes such a request, the Secretary must send to the person a copy of so much (if any) of that entry as is contained in any computer database maintained by the Department for purposes connected with the administration of this Act (other than any part of that entry that was supplied in confidence by another person).

(3) If the person makes such a request, then, instead of providing a copy of an entry to the person, the Secretary may, if the request is for the provision of an electronic copy, provide the information contained in the entry:

(a) on a data processing device; or

(b) by way of electronic transmission.

9D Variation of entries in Register

(1) The Secretary may:

(a) following a request by a person in relation to whom therapeutic goods are entered on the Register; or

(b) on the Secretary’s own initiative;

vary the entry in the Register in relation to the goods if the entry contains information that is incomplete or incorrect.

(2) If:

(a) the person in relation to whom therapeutic goods are registered or listed has requested the Secretary to vary product information included in the entry in the Register that relates to the goods; and

(b) the only effect of the variation would be:

   (i) to reduce the class of persons for whom the goods are suitable; or

   (ii) to add a warning, or precaution, that does not include any comparison of the goods with any other therapeutic goods by reference to quality, safety or efficacy;

the Secretary must vary the entry in accordance with the request.

(3) If:

(a) the person in relation to whom therapeutic goods are registered or listed has requested the Secretary to vary information included in the entry in the Register that relates to the goods; and
Section 9E

(b) subsection (2) does not apply to the request; and
(c) the Secretary is satisfied that the variation requested does not
indicate any reduction in the quality, safety or efficacy of the
goods for the purposes for which they are to be used;
the Secretary may vary the entry in accordance with the request.

(4) If:
(a) particular therapeutic goods cease to be medical devices
because of a declaration under subsection 41BD(3); and
(b) those goods are included in the Register under Chapter 4 as a
kind of medical device;
the Secretary must move the entry relating to the goods from the
part of the Register for medical devices to the part for goods to be
known as registered goods or to the part for goods to be known as
listed goods (whichever is applicable).

(5) In this section:

*product information*, in relation to therapeutic goods, means
information relating to the safe and effective use of the goods,
including information regarding the usefulness and limitations of
the goods.

Note: Variations to the Register also occur to give effect to limited
cancellations of entries of kinds of medical devices from the Register:
see subsection 41GO(2).

9E Publication of list of goods on Register

The Secretary must, at least once every 12 months, publish a list of
the therapeutic goods included in the Register.
Chapter 3—Medicines and other therapeutic goods that are not medical devices

Note: For 5 years following the commencement of Chapter 4 (Medical devices), this Chapter will still apply to medical devices that are registered or listed goods.

Part 3-1—Standards

10 Determination of standards

(1) The Minister may, by order published in the Gazette, determine that matters specified in the order constitute a standard for therapeutic goods or a class of therapeutic goods identified in the order (whether or not those goods are the subject of a monograph in the British Pharmacopoeia or the British Pharmacopoeia (Veterinary)).

(2) Without limiting the generality of subsection (1), an order establishing a standard for therapeutic goods may:
   (a) be specified by reference to:
      (i) the quality of the goods; or
      (ii) the quantity of the goods when contained in specified containers; or
      (iii) procedures to be carried out in the manufacture of the goods; or
      (iv) a monograph in the British Pharmacopoeia or the British Pharmacopoeia (Veterinary); or
      (v) a monograph in another publication approved by the Minister for the purposes of this subsection; or
      (vi) such a monograph as modified in a manner specified in the order establishing the standard; or
      (vii) a standard published by the Standards Australia International Limited; or
      (viii) such other matters as the Minister thinks fit; or
   (b) require that a matter relating to the standard be determined in accordance with a particular test; or
Section 10A

(c) require that therapeutic goods or a class of therapeutic goods identified in the order be labelled or packaged in a manner, or kept in containers that comply with requirements, specified in the order.

(3) Without limiting the generality of paragraph (2)(c), the Minister may, in an order establishing a standard, direct that there be set out, in a manner specified in the order, on:
   (a) therapeutic goods or a class of therapeutic goods identified in the order; or
   (b) a container or package containing therapeutic goods or a class of therapeutic goods identified in the order; or
   (c) a label of therapeutic goods or a class of therapeutic goods identified in the order;
   such particulars as are required by the order.

(4) The Minister must not determine a standard or amend or revoke a standard unless the Minister has consulted with respect to the proposed action with a committee established by the regulations to advise the Minister on standards.

10A Application of standards to medical devices

A standard under section 10 does not apply to a medical device unless Part 3-2 applies to the device.

Note: Section 15A sets out when Part 3-2 applies to a medical device.

11 Date of effect of standards

A standard under section 10 takes effect on the day on which the order establishing the standard is published in the Gazette or on such later day as is specified in the order.

12 Standards to be disallowable

Standards under section 10 and orders revoking, varying or modifying standards of that kind are disallowable instruments for the purposes of section 46A of the Acts Interpretation Act 1901.
Section 13

13 Special provisions relating to standards

(1) Unless the contrary intention appears in a standard, the standard applies to therapeutic goods for use in humans and therapeutic goods for use in animals.

(2) For the purposes of this Part, where a statement in a monograph in the British Pharmacopoeia or the British Pharmacopoeia (Veterinary) refers to a statement in a monograph in another publication, the first-mentioned statement is to be taken to include the other statement.

(3) Subject to subsection (4), where:
(a) a standard applicable to therapeutic goods is constituted by statements in a monograph in the British Pharmacopoeia or the British Pharmacopoeia (Veterinary); and
(b) requirements applicable to the labelling or packaging of the goods are specified in the British Pharmacopoeia or the British Pharmacopoeia (Veterinary); and
(c) the goods are not labelled or packaged in accordance with those requirements;
the goods are to be taken not to comply with that standard.

(4) Where:
(a) a standard under section 10 applies to therapeutic goods; and
(b) requirements applicable to the goods are specified in the British Pharmacopoeia or the British Pharmacopoeia (Veterinary); and
(c) those requirements are inconsistent with the requirements specified in the standard;
the requirements referred to in paragraph (b) are, so far as they are inconsistent, to be disregarded for the purposes of this Act.

(5) Where:
(a) a standard applies to a class of therapeutic goods; and
(b) another standard applies to some only of the therapeutic goods within that class; and
Section 14

(c) those standards are inconsistent;
the standard referred to in paragraph (a) is, to the extent of the inconsistency, of no effect in relation to the goods referred to in paragraph (b).

(6) Where:
(a) therapeutic goods consist, or are represented to consist, of a mixture of ingredients or of a combination of component parts; and
(b) a standard is applicable to the mixture or the combination;
that standard takes precedence over any standard that is applicable to the ingredients or the component parts.

(7) Where:
(a) therapeutic goods consist, or are represented to consist, of a mixture of ingredients or of a combination of component parts; and
(b) there is no standard applicable to the goods but a standard is applicable to at least one of the ingredients or component parts; and
(c) the Minister has, by order published in the Gazette, determined that the standard does not apply to the goods;
the standard is to be disregarded in so far as it would otherwise apply to the goods.

14 Compliance with standards

(1) Except with the consent in writing of the Secretary, a person must not:
(a) import therapeutic goods into Australia; or
(b) supply therapeutic goods for use in Australia;
if the goods do not conform with a standard applicable to the goods.

Maximum penalty: Imprisonment for 12 months or 1,000 penalty units, or both.

(2) Paragraph (1)(a) does not apply to goods that do not conform with a standard applicable to the goods by reason only of matters relating to labelling or packaging.
Section 15

(3) Except in exceptional circumstances and with the consent in writing of the Secretary, a person must not export therapeutic goods from Australia if the goods do not conform with a standard applicable to the goods (other than a standard relating to the labelling of the goods for supply in Australia).

Maximum penalty:  Imprisonment for 12 months or 1,000 penalty units, or both.

(4) Where:

(a) the importation or exportation of goods is prohibited under subsection (1) or (3); and

(b) the Secretary notifies the Comptroller-General in writing that the Secretary wishes the Customs Act 1901 to apply to that importation or exportation;

the Customs Act 1901 has effect as if the goods included in that importation or exportation were goods described as forfeited to the Crown under section 229 of that Act because they were:

(c) prohibited imports within the meaning of that Act; or

(d) prohibited exports within the meaning of that Act; as the case requires.

(5) The Secretary must, as soon as practicable after making a decision to give a consent under this section, cause particulars of the decision to be published in the Gazette.

(6) The Secretary must, within 28 days after making a decision to refuse to give a consent under this section, notify the applicant in writing of the decision and of the reasons for the decision.

15 Consent may be subject to conditions etc.

(1) The consent of the Secretary under section 14 may be given:

(a) unconditionally or subject to conditions; or

(b) in respect of particular goods or classes of goods.

(2) A person is guilty of an offence if:

(a) the person engages in conduct; and

(b) the conduct breaches a condition of such a consent.

Penalty:  120 penalty units.
Section 15

(2A) In subsection (2):

*engage in conduct* means:

(a) do an act; or

(b) omit to perform an act.
Part 3-2—Registration and listing of therapeutic goods

Division 1—Preliminary

15A Application of this Part to medical devices

The general rule

(1) This Part does not apply to a medical device unless this section provides otherwise.

Previously registered or listed devices

(2) If a medical device is registered goods or listed goods before the commencement of this section, this Part continues to apply to the device unless the registration or listing is cancelled.

Note: A registration or listing can be cancelled under section 30, or can be taken to be cancelled under section 9B.

Pending applications

(3) This Part continues to apply to a medical device if:
   (a) before the commencement of this section, an application was made under Part 3 for registration or listing of therapeutic goods that include that medical device; and
   (b) immediately before that commencement, the application was not yet finally determined; and
   (c) the application has not been, and is not, withdrawn either before or after that commencement.

However, this Part ceases to apply to the device if, having been registered goods or listed goods, the registration or listing is cancelled.

(4) For the purposes of paragraph (3)(b), an application is finally determined when the application, and any applications for review or appeals arising out of it, have been finally determined or otherwise disposed of.
Chapter 3  Medicines and other therapeutic goods that are not medical devices
Part 3-2  Registration and listing of therapeutic goods
Division 1  Preliminary

Section 15A

Applications made within 2 years relating to certain medical devices

(5) This Part applies to a medical device if:

(a) regulations made for the purposes of this section specified either:

(i) the medical device classification applying to the kind of medical device that includes that device; or
(ii) medical devices of that kind; and

(b) during the period of 2 years after the commencement of this section, an application was made under Part 3-2 for registration or listing of therapeutic goods that include that medical device.

However, this Part ceases to apply to the device if, having been registered goods or listed goods, the registration or listing is cancelled.

Note: Medical devices that are registered or listed because of this subsection are taken to be cancelled 2 years after Chapter 4 commences, or before then if medical devices of that kind are included in the Register under Chapter 4: see subsection 9B(1).

Medical devices that are exempt goods

(6) This Part applies to a medical device, during the period of 2 years after the commencement of this section, if the device is exempt goods.

Existing approvals under section 19

(7) This Part continues to apply to a medical device if:

(a) an approval or authorisation in force under section 19 applies to the device; and

(b) that approval or authorisation was in force immediately before the commencement of this section.

New approvals under section 19

(8) This Part applies to a medical device if:

(a) subsection (7) does not apply to the device; and
(b) during the period of 2 years after the commencement of this section, an approval is granted or an authorisation is given under section 19 that applies to the device. However, this subsection does not apply after the end of that period.

16 Therapeutic goods and gazetted groups

(1) For the purposes of this Part, therapeutic goods (other than medicine of the kind to which subsection (1A) applies) are to be taken to be separate and distinct from other therapeutic goods if they have:

(a) a different formulation, composition or design specification; or
(b) a different strength or size (disregarding pack size); or
(c) a different dosage form or model; or
(d) a different name; or
(e) different indications; or
(f) different directions for use; or
(g) a different type of container (disregarding container size).

(1A) Medicines that are listable goods (other than export only medicines) are taken to be separate and distinct from other therapeutic goods if the medicines have:

(a) different active ingredients; or
(b) different quantities of active ingredients; or
(c) a different dosage form; or
(d) such other different characteristics as the regulations prescribe; from the therapeutic goods.

(2) The Secretary may, by order published in the Gazette, determine that a group of therapeutic goods (not being medical devices or therapeutic devices) identified in the order is a gazetted therapeutic goods group because the goods within the group have common characteristics.

(3) The Secretary may, by order published in the Gazette, determine that a group of therapeutic goods (being therapeutic devices)
identified in the order is a gazetted therapeutic devices group because the goods within the group:
   (a) have common characteristics; and
   (b) have been produced by the same manufacturer.

(3A) The Secretary may, by order published in the *Gazette*, determine that a group of kits identified in the order is a gazetted kits group.

(4) An order under subsection (2), (3) or (3A) may make provision for or in relation to a matter by applying, adopting or incorporating, with or without modification, a document as in force from time to time, if the document is:
   (a) published by the Department (whether in electronic form or otherwise); and
   (b) available for sale to the public; and
   (c) available for inspection (whether by using a visual display unit or otherwise) by the public at offices of the Department specified by the Secretary.

### 18 Exempt goods

(1) The regulations may, subject to such conditions (if any) as are specified in the regulations, exempt:
   (a) all therapeutic goods, except those included in a class of goods prescribed for the purposes of this paragraph; or
   (b) specified therapeutic goods; or
   (c) a specified class of therapeutic goods;

from the operation of this Part (except section 31A and sections 31C to 31F).

(2) An exemption in terms of paragraph (1)(a) has effect only in relation to such classes of persons as are prescribed for the purposes of this subsection.

(3) Where the regulations revoke an exemption, the revocation takes effect on the day, not being earlier than 28 days after the day on which the regulations are made, specified in the regulations.
18A Exemption because of emergency

Minister’s power

(1) The Minister may exempt from the operation of Division 2 of this Part:
   (a) specified therapeutic goods; or
   (b) therapeutic goods in a specified class.

The exemption must be made in writing.

(2) The Minister may exempt goods under subsection (1) only if the Minister is satisfied that, in the national interest:
   (a) the exemption should be made so that the goods may be stockpiled as quickly as possible in order to create a preparedness to deal with a potential threat to public health that may be caused by a possible future emergency; or
   (b) the exemption should be made so that the goods can be made available urgently in Australia in order to deal with an actual threat to public health caused by an emergency that has occurred.

When the exemption has effect

(3) The exemption takes effect:
   (a) on the day on which the exemption is made; or
   (b) on a later day that is specified in the exemption.

(4) The exemption ceases to have effect:
   (a) at the end of the period specified by the Minister in the exemption as the period for which the exemption is to have effect; or
   (b) when the exemption is revoked; whichever first occurs.

(5) The exemption ceases to have effect in relation to particular therapeutic goods:
   (a) when those goods become registered or listed goods; or
   (b) when the Minister varies the exemption by removing those goods from the exemption; whichever first occurs.
(6) If the Minister revokes the exemption as mentioned in paragraph (4)(b), or varies the exemption as mentioned in paragraph (5)(b), the revocation or variation takes effect:
   (a) if the Minister states in the revocation or variation that the revocation or variation is necessary to prevent imminent risk of death, serious illness or serious injury—on the day on which the revocation or variation is made; or
   (b) in any other case—on the day specified by the Minister in the revocation or variation.

The day specified under paragraph (b) of this subsection must not be earlier than 28 days after the day on which the revocation or variation is made.

Note: The revocation or variation must be made in writing, see subsection 33(3) of the Acts Interpretation Act 1901.

Conditions for the exemption

(7) The exemption is subject to conditions specified in the exemption about any of the following:
   (a) the period for which the exemption is to have effect;
   (b) the quantity of goods that are exempt;
   (c) the source of those goods;
   (d) the persons or class of persons who may import, manufacture, supply or export those goods;
   (e) the supply of those goods (including the persons or class of persons to whom goods may be supplied for use and the circumstances under which a stockpile of goods may be supplied for use);
   (f) the storage and security of those goods;
   (g) the keeping and disclosure of, and access to, records about those goods;
   (h) the disposal of those goods;
   (i) the manner in which any of those goods are to be dealt with if a condition of the exemption is breached;
   (j) any other matters that the Minister thinks appropriate.

Whether or not goods are exempt under this section is not affected by whether or not there is a breach of a condition of an exemption under this section in relation to those goods.
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Note: A person may commit an offence by breaching a condition of an exemption under this section, see subsections 20(2A) and (2C), 22(7AB) and (7AD), and 30H(1) and (3).

(8) The Minister may revoke or vary the conditions (including by imposing new conditions) after the exemption is made. The revocation or variation must be made in writing.

(9) A revocation or variation under subsection (8) takes effect:
   (a) if the Minister states in the revocation or variation that the revocation or variation is necessary to prevent imminent risk of death, serious illness or serious injury—on the day on which the revocation or variation is made; or
   (b) in any other case—on the day specified by the Minister in the revocation or variation.

The day specified under paragraph (b) must not be earlier than 28 days after the day on which the revocation or variation is made.

Exemption etc. to be disallowable

(9A) An exemption covered by paragraph (2)(a), and a revocation or variation under subsection (8) of an exemption covered by paragraph (2)(a), are disallowable instruments for the purposes of section 46A of the Acts Interpretation Act 1901.

Notification

(10) The Secretary must cause a document setting out particulars of:
   (a) an exemption covered by paragraph (2)(b); and
   (b) a revocation or variation under subsection (8) of an exemption covered by paragraph (2)(b);

to be published in the Gazette within 5 working days after the day on which the Minister makes the exemption, revocation or variation. However, an exemption, or a revocation or variation, is not invalid merely because of a failure to comply with this subsection.

Tabling

(11) The Minister must cause a document setting out particulars of:
   (a) an exemption covered by paragraph (2)(b); and
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(b) a revocation or variation under subsection (8) of an exemption covered by paragraph (2)(b); to be tabled before each House of the Parliament within 5 sitting days of that House after the day on which the Minister makes the exemption, revocation or variation. However, an exemption, or a revocation or variation, is not invalid merely because of a failure to comply with this subsection.

Exclusion of liability of the Commonwealth etc.

(12) An exemption under this section does not render the Commonwealth, the Minister or a delegate of the Minister liable to a person for loss, damage or injury of any kind suffered by the person as a result of, or arising out of, the use of therapeutic goods by that person or another person.

Note: There are other requirements in other parts of this Act about goods exempt under this section:

- sections 20 and 22 (breach of a condition of the exemption);
- section 30F (goods not conforming to standards etc.);
- section 30G (disposal of unused goods);
- section 30H (record keeping);
- section 31AA (providing information to the Secretary);
- sections 35, 39 and 41 (manufacturing goods that are exempt under this section);
- section 46A (search of premises).

19 Exemptions for special and experimental uses

(1) The Secretary may, by notice in writing, grant an approval to a person for the importation into, or the exportation from, Australia or the supply in Australia of specified therapeutic goods that are not registered goods, listed goods or exempt goods:

- for use in the treatment of another person; or
- for use solely for experimental purposes in humans;

and such an approval may be given subject to such conditions as are specified in the notice of approval.

(1A) An approval for the purpose mentioned in paragraph (1)(b) is subject to the conditions (if any) specified in the regulations. Those
(2) An application for an approval must be made to the Secretary and must:

(a) in the case of an application for use of the kind referred to in paragraph (1)(a)—be accompanied by such information relating to the goods the subject of the application as is required by the Secretary; and

(b) in the case of an application for use of the kind referred to in paragraph (1)(b):
   (i) be made in writing; and
   (ii) be accompanied by such information relating to the goods the subject of the application as is required by the Secretary; and
   (iii) be accompanied by the prescribed evaluation fee.

(3) Without limiting the conditions to which an approval under subsection (1) may be made subject, those conditions may include a condition relating to the charges that may be made for the therapeutic goods to which the approval relates.

(4) Where an application for an approval is made, the Secretary must, after having considered the application and, in the case of an application for the use of therapeutic goods for experimental purposes in humans, after having evaluated the information submitted with the application, notify the applicant of the decision on the application within 28 days of making the decision and, in the case of a decision not to grant the approval, of the reasons for the decision.

(4A) The use by a person for experimental purposes in humans of specified therapeutic goods that are the subject of an approval granted to someone else under paragraph (1)(b) is subject to the conditions (if any) specified in the regulations relating to one or more of the following:

(a) the preconditions on the use of the goods for those purposes;
(b) the principles to be followed in the use of the goods for those purposes;
(c) the monitoring of the use, and the results of the use, of the goods for those purposes;
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(d) the circumstances in which the person must cease the use of the goods for those purposes.

(5) The Secretary may, in writing, authorise a specified medical practitioner to supply:

(a) specified therapeutic goods for use in the treatment of humans; or
(b) a specified class of such goods;

to the class or classes of recipients specified in the authority.

(5A) An authority may be given subject to the conditions (if any) specified in the authority.

(5B) The Secretary may impose conditions (or further conditions) on an authority given to a person under subsection (5) by giving to the person written notice of the conditions (or further conditions).

(6) An authority under subsection (5) may only be given:

(a) to a medical practitioner included in a class of medical practitioners prescribed by the regulations for the purposes of this paragraph; and

(aa) to a medical practitioner who has the approval of an ethics committee to supply the specified therapeutic goods or the specified class of such goods; and

(b) in relation to a class or classes of recipients prescribed by the regulations for the purposes of this paragraph.

Paragraph (aa) does not apply in the exceptional circumstances (if any) prescribed by the regulations for the purposes of this subsection.

(7) The regulations may prescribe the circumstances in which therapeutic goods may be supplied under an authority under subsection (5).

(8) The giving of an authority under subsection (5) does not render the Commonwealth, the Secretary or a delegate of the Secretary liable to a person in respect of loss, damage or injury of any kind suffered by the person as a result of, or arising out of, the use of therapeutic goods by that person or another person.

(9) In this section, medical practitioner means a person who is registered, in a State or internal Territory, as a medical practitioner.
19A Exemptions where unavailability etc. of therapeutic goods

(1) The Secretary may, by notice in writing, grant an approval to a person for the importation into Australia, or the supply in Australia, of specified therapeutic goods if the Secretary is satisfied that:

(a) registered goods that could act as a substitute for the goods are unavailable or are in short supply; and

(b) either:

(i) the goods that are the subject of the application are registered or approved for general marketing in at least one foreign country specified by the Secretary in a determination under subsection (3); or

(ii) an application that complies with section 23 has been made under that section for registration of the goods; and

(c) the goods are of a kind:

(i) included in Schedule 10 of the Therapeutic Goods Regulations; or

(ii) specified by the Secretary in a determination under subsection (4); and

(d) the approval is necessary in the interests of public health.

(2) The Secretary may, by notice in writing, grant an approval to a person for the importation into Australia, or the supply in Australia, of specified therapeutic goods if the Secretary is satisfied that:

(a) registered goods that could act as a substitute for the goods do not exist; and

(b) an application that complies with section 23 has been made under that section for registration of the goods; and

(c) the goods are of a kind:

(i) included in Schedule 10 of the Therapeutic Goods Regulations; or

(ii) specified by the Secretary in a determination under subsection (4); and

(d) the approval is necessary in the interests of public health.
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(3) The Secretary may, for the purposes of subparagraph (1)(b)(i), make written determinations specifying the foreign countries in which registration or approval for general marketing of the goods is a prerequisite for approval by the Secretary under this section.

(4) The Secretary may make written determinations specifying the kinds of goods that can be the subject of an approval under this section.

(5) Determinations under subsections (3) and (4) are disallowable instruments for the purposes of section 46A of the Acts Interpretation Act 1901.

(6) The Secretary may grant the approval subject to any conditions that are specified in the notice of approval.

(7) The Secretary may grant the approval for such period as is specified in the notice of approval.

(8) The approval lapses if:
   (a) the period specified in the notice of approval expires; or
   (b) a decision has been made under section 25 in relation to the goods.

(9) The approval lapses if:
   (a) the Secretary is satisfied that paragraph (1)(a), (b), (c) or (d), or paragraph (2)(a), (b), (c) or (d), as the case requires, no longer applies in relation to the goods, or that a condition of the approval has been contravened; and
   (b) the Secretary has given to the person to whom the approval was granted a notice stating that the Secretary is so satisfied.

(10) The lapsing of the approval on the expiry of the period specified in the notice of approval does not prevent another approval being granted under this section in relation to the goods before the lapsing of the first-mentioned approval. The other approval may be expressed to take effect on the expiry of that period.

20 Offences relating to importation, exportation, manufacture and supply of therapeutic goods

(1) A person is guilty of an offence if:
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(a) the person:
   (i) imports into Australia therapeutic goods for use in humans; or
   (ii) exports from Australia therapeutic goods for use in humans; or
   (iii) manufactures in Australia therapeutic goods for use in humans; or
   (iv) supplies in Australia therapeutic goods for use in humans; and
(b) none of the following subparagraphs applies in relation to the goods:
   (i) the goods are registered goods or listed goods in relation to the person;
   (ii) the goods are exempt goods;
   (iia) the goods are exempt under section 18A;
   (iii) the goods are the subject of an approval or authority under section 19;
   (iv) the goods are the subject of an approval under section 19A.

Note: A person may commit an offence against subsection (2A) or (2C) by importing into Australia therapeutic goods that are exempt under section 18A.

(1AA) An offence against subsection (1) is punishable on conviction by imprisonment for 12 months or a fine not more than 1,000 penalty units, or both.

(1A) It is a defence to a prosecution under subsection (1) if the defendant proves that the defendant was not the sponsor of the goods at the time of the importation, export, manufacture or supply, as the case may be.

Note: The defendant bears a legal burden in relation to the matter in subsection (1A). See section 13.4 of the Criminal Code.

(1B) A person is guilty of an offence if:
   (a) the person is the sponsor of therapeutic goods for use in humans; and
   (b) the person:
      (i) imports the goods into Australia; or
      (ii) exports the goods from Australia; or
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(iii) manufactures the goods in Australia; or
(iv) supplies the goods in Australia; and
(c) the person has not, at the time of the importation, export, manufacture or supply, properly notified to the Secretary either or both of the following:
   (i) the manufacturer of the goods;
   (ii) premises used in the manufacture of the goods.

Maximum penalty: Imprisonment for 12 months or 1,000 penalty units, or both.

(1C) For the purposes of paragraph (1B)(c):
   (a) a manufacturer is properly notified to the Secretary if:
      (i) the manufacturer was nominated, as a manufacturer of the goods, in an application for the registration or listing of the goods; or
      (ii) the Secretary was subsequently informed in writing that the manufacturer is a manufacturer of the goods; and
   (b) premises are properly notified to the Secretary if:
      (i) the premises were nominated, as premises used in the manufacture of the goods, in an application for the registration or listing of the goods; or
      (ii) the Secretary was subsequently informed in writing that the premises are used in the manufacture of the goods.

(2) A person in relation to whom therapeutic goods are registered or listed must not import those goods into Australia, or supply those goods in Australia, unless:
   (a) the registration number or listing number of the goods is set out on the label of the goods in the prescribed manner or, in the case of an importation, that number is so set out, or is to be so set out before the goods are supplied in Australia; or
   (b) the goods are devices that are listed goods.

Maximum penalty: 60 penalty units.

(2A) A person commits an offence if:
   (a) the person imports therapeutic goods into Australia; and
   (b) the goods are exempt under section 18A; and
   (c) the importation breaches a condition of the exemption.
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Maximum penalty: Imprisonment for 4 years or 240 penalty units, or both.

(2B) Strict liability applies to paragraph (2A)(b).

Note: For strict liability, see section 6.1 of the *Criminal Code*.

(2C) A person commits an offence if:

(a) the person imports therapeutic goods into Australia; and

(b) the goods are exempt under section 18A; and

(c) the importation breaches a condition of the exemption.

Maximum penalty: 60 penalty units.

(2D) An offence under subsection (2C) is an offence of strict liability.

Note: For strict liability, see section 6.1 of the *Criminal Code*.

(3) Where:

(a) the importation or exportation of goods is prohibited under subsection (1); and

(b) the Secretary notifies the Comptroller-General in writing that the Secretary wishes the *Customs Act 1901* to apply to that importation or exportation;

the *Customs Act 1901* has effect as if the goods included in that importation or exportation were goods described as forfeited to the Crown under section 229 of that Act because they were:

(c) prohibited imports within the meaning of that Act; or

(d) prohibited exports within the meaning of that Act; as the case requires.

21 Offence relating to wholesale supply

A person must not supply in Australia therapeutic goods for use in humans (other than listable devices), being goods of which the person is not a sponsor, to another person who is not the ultimate consumer of the goods unless:

(a) the goods are registered goods or listed goods; or

(b) the goods are exempt goods; or

(ba) the goods are exempt under section 18A; or

(c) the goods are the subject of an approval or authority under section 19; or

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(d) the goods are the subject of an approval under section 19A.

Maximum penalty: 120 penalty units.

22 General offences relating to this Part

(1) A person must not set out or cause to be set out, on a container or package that contains therapeutic goods or on a label of goods of that kind, a number that purports to be the registration number or listing number of the goods in relation to a particular person if the number is not that number.

Maximum penalty: 60 penalty units.

(2A) A person must not, in or in connection with a certification of any matter under subsection 26A(2), make a statement that is false or misleading in a material particular.

Maximum penalty: Imprisonment for 12 months or 1,000 penalty units, or both.

(3) A person is guilty of an offence if:
   (a) therapeutic goods are registered or listed in relation to the person; and
   (b) the person engages in conduct; and
   (c) the conduct breaches a condition of the registration or listing of the goods.

Penalty: 60 penalty units.

(3A) In subsection (3):

engage in conduct means:
   (a) do an act; or
   (b) omit to perform an act.

(4) A person must not:
   (a) represent therapeutic goods that are not included in the Register as being so included; or
   (b) represent therapeutic goods that are not exempt goods as being exempt goods; or
(ba) represent therapeutic goods that are not goods exempt under section 18A as being goods exempt under that section; or
(c) represent therapeutic goods that are included in one part of the Register as being included in the other part of the Register; or
(d) represent therapeutic goods that are not the subject of an approval or authority under section 19 as being the subject of such an approval or authority; or
(e) represent therapeutic goods that are not the subject of an approval under section 19A as being the subject of such an approval.

Maximum penalty: 60 penalty units.

(5) A person, being the sponsor of therapeutic goods that are included in the Register, must not, by any means, advertise the goods for an indication other than those accepted in relation to the inclusion of the goods in the Register.

Maximum penalty: 60 penalty units.

(6) A person must not make a claim, by any means, that the person or another person can arrange the supply of therapeutic goods (not being exempt goods or goods exempt under section 18A) that are not registered goods or listed goods.

Maximum penalty: 60 penalty units.

(7) A person is guilty of an offence if:
(a) the person does an act or omits to do an act; and
(b) the act or omission results in the breach of:
(i) a condition of an exemption applicable under regulations made for the purposes of subsection 18(1); or
(ii) a condition of an approval under section 19; or
(iii) a condition applicable under regulations made for the purposes of subsection 19(4A); or
(iv) a condition of an approval under section 19A.

(7AA) An offence against subsection (7) is punishable on conviction by a fine of not more than 60 penalty units.
(7AB) A person commits an offence if:
(a) the person does an act or omits to do an act in relation to therapeutic goods; and
(b) the goods are exempt under section 18A; and
(c) the act or omission results in the breach of a condition of the exemption; and
(d) the act or omission is likely to cause a serious risk to public health.

Maximum penalty: Imprisonment for 5 years or 300 penalty units, or both.

Note 1: A person may commit an offence against subsection 20(2A) or (2C) by breaching a condition of an exemption of therapeutic goods under section 18A that relates to the importation of the goods.

Note 2: A person may commit an offence against subsection 30H(1) or (3) by breaching a condition of an exemption of therapeutic goods under section 18A that relates to records about the goods.

(7AC) Strict liability applies to paragraph (7AB)(b).

Note: For strict liability, see section 6.1 of the **Criminal Code**.

(7AD) A person commits an offence if:
(a) the person does an act or omits to do an act in relation to therapeutic goods; and
(b) the goods are exempt under section 18A; and
(c) the act or omission results in the breach of a condition of the exemption.

Maximum penalty: Imprisonment for 4 years or 240 penalty units, or both.

(7AE) Strict liability applies to paragraph (7AD)(b).

Note: For strict liability, see section 6.1 of the **Criminal Code**.

(7A) A person to whom an authority under subsection 19(5) has been granted must not supply the therapeutic goods to which the authority relates except in accordance with:
(a) the authority; and
(aa) the conditions (if any) to which the authority is subject; and
(b) any regulations made for the purpose of subsection 19(7).

Maximum penalty: 60 penalty units.

(8) A person must not use therapeutic goods, other than exempt goods, listed goods, registered goods, goods exempt under section 18A or goods that are the subject of an approval under section 19A:

(a) for use in the treatment of another person; or

(b) for use solely for experimental purposes in humans; except in accordance with an approval or authority under section 19 or a condition applicable under regulations made for the purposes of subsection 19(4A).

Maximum penalty: 60 penalty units.

22A False statements in applications for registration

A person must not, in or in connection with an application for registration of therapeutic goods, make a statement that is false or misleading in a material particular.

Maximum penalty: Imprisonment for 12 months or 1,000 penalty units, or both.
Chapter 3  Medicines and other therapeutic goods that are not medical devices
Part 3-2  Registration and listing of therapeutic goods
Division 2  Registration and listing

Section 23AA

Division 2—Registration and listing

23AA  Ministerial approval of evaluation, registration or listing of restricted goods

(1) In spite of any provision of this Division, restricted goods must not be evaluated or registered or listed without the written approval of the Minister.

(2) A written approval shall be laid before each House of the Parliament by the Minister within 5 sitting days of being given.

23  Applications generally

(1) An application for registration or listing of therapeutic goods must:
   (a) be made in accordance with a form approved, in writing, by the Secretary or in such other manner as is approved, in writing, by the Secretary; and
   (b) be delivered to an office of the Department specified by the Secretary.

(2) An application is not effective unless:
   (a) the prescribed application fee has been paid; and
   (b) the applicant has delivered to the office to which the application was made such information, in a form approved, in writing, by the Secretary, as will allow the determination of the application; and
   (c) if the Secretary so requires—the applicant has delivered to the office to which the application was made a reasonable number of samples of the goods.

(3) An approval of a form may require or permit an application or information to be given in accordance with specified software requirements:
   (a) on a specified kind of data processing device; or
   (b) by way of a specified kind of electronic transmission.

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24 Applications for registration

(1) Where an application is made for the registration of therapeutic goods in accordance with section 23 and the goods are goods that are required to be registered, a fee specified in or determined in accordance with the regulations is payable by the applicant in respect of the evaluation of the goods for registration, and the Secretary must notify each such applicant of the amount of the evaluation fee.

(2) Subject to section 24D, an application for registration of therapeutic goods lapses if:

(a) any part of the evaluation fee payable in respect of those goods remains unpaid at the end of the period of 2 months after the day on which the amount became due and payable; or

(b) the application contains information that is inaccurate or misleading in a material particular; or

(c) information given to the Secretary by, or on behalf of, the applicant in connection with the application, including information given for the purpose of a requirement under section 31, is inaccurate or misleading in a material particular; or

(d) the applicant fails to comply with a requirement under section 31 to give information consisting of individual patient data in relation to the goods.

(3) In this section, individual patient data, in relation to therapeutic goods, means information, derived from clinical trials, relating to individuals before, during and after the administration of the goods to those individuals, including, but not limited to, demographic, biochemical and haematological information.

24A When evaluation fee due for payment

Subject to sections 24B and 24D, an evaluation fee under section 24 payable by an applicant is due and payable on the day on which the applicant is notified of the amount of the evaluation fee.
Chapter 3 Medicines and other therapeutic goods that are not medical devices
Part 3-2 Registration and listing of therapeutic goods
Division 2 Registration and listing

Section 24B

24B Payment of evaluation fee by instalments

(1) The regulations may provide for the payment of an evaluation fee under section 24 to be made by such instalments and at such times as are ascertained in accordance with the regulations, and the evaluation fee is due and payable accordingly.

(2) Regulations made for the purposes of subsection (1) may provide that a person is not allowed to pay an evaluation fee under section 24 by instalments if any part of an instalment of:
   (a) that or any other evaluation fee under section 24 payable by the person; or
   (b) any assessment fee under section 41LA payable by the person;
was unpaid immediately after the time when it became due for payment.

(3) Subsection (2) does not limit the generality of subsection (1).

24C Recovery of evaluation fee

An evaluation fee under section 24 may be recovered by the Commonwealth as a debt due to the Commonwealth.

24D Reduction of evaluation fee where evaluation not completed within prescribed period

(1) This section applies to an application under section 23 in relation to therapeutic goods for the evaluation of which a period is prescribed under paragraph 63(2)(da).

(2) Nothing in section 24, 24A or 24B requires the applicant to pay more than three-quarters of the evaluation fee before the completion of the evaluation of the goods.

(3) If the evaluation is not completed within the period referred to in subsection (1), this Act has effect as if the evaluation fee were reduced to three-quarters of the fee that, under the regulations, would have been the evaluation fee.

(4) If:

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(a) the evaluation is completed within the period referred to in subsection (1); and
(b) part of the evaluation fee under section 24 is unpaid when the evaluation is completed;
that part becomes due and payable on the completion of the evaluation.

(5) For the purposes of subsections (2), (3) and (4), the evaluation is to be taken to be completed when the applicant is notified according to subsection 25(3) of the Secretary’s decision on the application.

(6) Despite subsection (5), if:
(a) the Secretary has given the applicant all evaluation reports relating to the application; and
(b) the Secretary has given those reports, or proposes to give those reports, to a committee established under the regulations to advise the Secretary on applications to register therapeutic goods to which subsection (1) applies; and
(c) the applicant withdraws the application after being given the reports, and before the end of the period mentioned in subsection (1);
the evaluation is taken for the purposes of subsections (2), (3) and (4) to be completed immediately before the time of withdrawal.

24E Deemed refusal of application

(1) This section applies in the case of an application under section 23 in relation to therapeutic goods for the evaluation of which a period is prescribed under paragraph 63(2)(da).

(2) If, at the end of the period referred to in subsection (1), the evaluation has not been completed, the applicant may give the Secretary written notice that the applicant wishes to treat the application as having been refused.

(3) A notice under subsection (2) may be given at any time before the evaluation is completed.

(4) Where a notice has been given, this Act (except for subsection 60(5)) has effect as if:
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(a) the Secretary had decided not to register the goods the subject of the application; and
(b) the Minister had made a decision under subsection 60(3) confirming the decision of the Secretary; and
(c) the Minister’s decision had been made on the day on which notice was given to the Secretary under subsection (2).

25 Evaluation and registration of therapeutic goods

(1) Where:
(a) an application is made for the registration of therapeutic goods in relation to a person in accordance with section 23; and
(b) there is no part of an evaluation fee under section 24 in respect of those goods that:
   (i) is due and payable by the person; and
   (ii) remains unpaid; and
(c) the person has complied with any requirements made by the Secretary under section 31 in relation to the goods; the goods are to be evaluated for registration having regard to:
(d) whether the quality, safety and efficacy of the goods for the purposes for which they are to be used have been satisfactorily established; and
(e) whether the presentation of the goods is acceptable; and
(f) whether the goods conform to any standard applicable to the goods, or any requirements relating to advertising applicable under Part 5-1 or under the regulations; and
(g) if a step in the manufacture of the goods has been carried out outside Australia—whether the manufacturing and quality control procedures used in the manufacture of the goods are acceptable; and
(h) if the goods have been manufactured in Australia—whether the goods have been manufactured in accordance with Part 3-3; and
(j) whether the goods contain substances that are prohibited imports for the purposes of the Customs Act 1901; and
(ja) whether all of the manufacturers of the goods are nominated as manufacturers of the goods in the application; and
(k) such other matters (if any) as the Secretary considers relevant.

Note: The Secretary must not use protected information when evaluating therapeutic goods for registration: see section 25A.

(2) In making a decision for the purposes of paragraph (1)(g), the matters that may be taken into account include:

(a) whether the applicant has provided:

(i) if the goods are not therapeutic devices and a step in the manufacture of the goods has been carried out in a country that is a member of the European Community or a member of EFTA—an EC/EFTA attestation of conformity in relation to the goods; or

(ii) in any other case—an acceptable form of evidence from a relevant overseas authority establishing that the manufacture of the goods is of an acceptable standard; and

(b) whether the applicant has agreed to provide, where the Secretary considers inspection of the manufacturing procedures used in the manufacture of the goods to be necessary:

(i) funds for the carrying out of that inspection by the Department; and

(ii) evidence that the manufacturer has agreed to such an inspection.

(2A) An evaluation under this section of goods in relation to which a period has been prescribed under paragraph 63(2)(da) must be completed within that period.

(2B) If therapeutic goods are exempt from the operation of Part 3-3 or a person is exempt from the operation of that Part in relation to the manufacture of the goods, subsection (1) has effect, in relation to the goods, as if paragraph (h) were omitted.
(2C) If a person is exempt from the operation of Part 3-3 in relation to a step in the manufacture of therapeutic goods, subsection (1) has effect, in relation to the goods, as if the reference in paragraph (h) to Part 3-3 were a reference to that Part to the extent that it applies to that person in relation to the manufacture of the goods.

(2D) If:
   (a) therapeutic goods were made outside Australia; and
   (b) had the goods been made in Australia, they would have been exempt from the operation of Part 3-3;
subsection (1) has effect, in relation to the goods, as if paragraph (g) were omitted.

(2E) A decision for the purposes of paragraph (1)(g) may also take into account any information provided to the Secretary by a health authority of a Convention country and relating to:
   (a) the general standards of manufacturing practice of a particular manufacturer; or
   (b) the specific standards of manufacture or control adopted by a particular manufacturer in relation to particular goods.

(2F) For the purposes of subsection (2E), a Convention country is a country that is a party to the Mutual Recognition Convention.

(2G) Information referred to in subsection (2E) and provided in accordance with the Mutual Recognition Convention is to be treated as equivalent to information obtained as a result of an inspection under Part 3-3 of this Act.

(3) If:
   (a) the therapeutic goods are therapeutic devices; and
   (b) the evaluation of the goods for registration has been completed;
the Secretary must:
   (c) notify the applicant in writing of his or her decision on the evaluation within 28 days of the making of the decision and, in the case of a decision not to register the goods, of the reasons for the decision; and
   (d) if the decision is to register the goods—include the goods in the Register and give the applicant a certificate of registration.
(4) If:

(a) the therapeutic goods are not therapeutic devices; and
(b) the evaluation of the goods for registration has been completed;

the Secretary must:

(c) notify the applicant in writing of his or her decision on the evaluation within 28 days of the making of the decision and, in the case of a decision not to register the goods, of the reasons for the decision; and

(d) if the decision is to register the goods:

(i) notify the applicant in writing that the goods will be included in the Register if the applicant gives the Secretary the certificate required under subsection 26B(1); and

(ii) include the goods in the Register and give the applicant a certificate of registration if the applicant gives the Secretary the certificate required under subsection 26B(1).

To avoid doubt, if the applicant gives the Secretary the certificate required under subsection 26B(1), the Secretary must include the goods in the Register under subparagraph (d)(ii) without inquiring into the correctness of the certificate.

(4A) Civil proceedings do not lie against the Secretary (or a delegate of the Secretary) in respect of loss, damage or injury of any kind suffered by another person as a result of the Secretary (or the delegate) including therapeutic goods in the Register in reliance on a certificate required under subsection 26B(1).

(5) The registration of therapeutic goods commences on the day specified for the purpose in the certificate of registration.

(6) The failure to complete an evaluation within the period mentioned in subsection (2A) does not render the Commonwealth, the Secretary or a delegate of the Secretary liable to a person in respect of loss, damage or injury of any kind caused by, or arising out of, the failure.
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25A When the Secretary must not use protected information

(1) When evaluating therapeutic goods for registration, the Secretary must not use information about other therapeutic goods that is protected information.

(2) Information is protected information if:
   (a) the information was given to the Secretary in relation to an application to register therapeutic goods (the new goods):
      (i) not being therapeutic devices; and
      (ii) consisting of, or containing, an active component; and
   (b) the information is about the active component and is not available to the public; and
   (c) when the application to register the new goods was lodged:
      (i) no other therapeutic goods consisting of, or containing, that active component were included in the Register; and
      (ii) no such therapeutic goods had been included in the Register at any time before then; and
   (d) the new goods became registered on or after the commencement of this subsection; and
   (e) 5 years have not passed since the day the new goods became registered; and
   (f) the person in relation to whom the new goods are registered has not given the Secretary permission in writing for the Secretary to use the information.

(3) For the purposes of subsection (2), an active component, in relation to therapeutic goods, is a substance that is, or one of the substances that together are, primarily responsible for the biological or other effect identifying the goods as therapeutic goods.

(4) The use of protected information contrary to subsection (1) does not render the Commonwealth, the Secretary or a delegate of the Secretary liable to a person in respect of loss, damage or injury of any kind suffered by the person as a result of, or arising out of, the use of that information.
25B  Registration of therapeutic device to which EC/EFTA attestation of conformity applies

(1) If:
   (a) an application is made in accordance with section 23 for the registration of a therapeutic device in relation to a person; and
   (b) the applicant gives to the Secretary an EC/EFTA attestation of conformity as to the matters that would require evaluation under subsection 25(1) if that subsection applied in relation to the device;

   the Secretary must register the device unless the Secretary considers that the device may compromise the health or safety of users.

(2) The Secretary must notify the applicant in writing of his or her decision on the application within 28 days of the making of the decision. If the Secretary decides not to register the device, the notice must contain the reasons for that decision.

(3) If the Secretary decides to register the device, the Secretary must:
   (a) include the device in the Register; and
   (b) give to the applicant a certificate of registration.

(4) The registration of the device commences on the day specified for the purpose in the certificate of registration.

26  Listing of therapeutic goods

(1) Where:
   (a) an application is made for the listing of therapeutic goods in relation to a person in accordance with section 23; and
   (aa) if goods are not therapeutic devices—the application is accompanied by the certificate required under subsection 26B(1); and
   (b) the person has complied with any requirements made by the Secretary under section 31 in relation to the goods; and
(ba) the goods are not goods which may be listed under section 26A;

then, subject to section 26AA, the Secretary is not to refuse to list the goods in relation to the person except where the Secretary is satisfied that:

(c) the goods are not eligible for listing; or

(d) the goods are not safe for the purposes for which they are to be used; or

(e) the presentation of the goods is unacceptable; or

(f) the goods do not conform to a standard applicable to the goods or to a requirement relating to advertising applicable under Part 5-1 or under the regulations; or

(g) if a step in the manufacture of the goods (not being therapeutic devices other than devices prescribed for the purposes of this paragraph) has been carried out outside Australia—the manufacturing and quality control procedures used in the manufacture of the goods are not acceptable; or

(h) if the goods have been manufactured in Australia—the goods have been manufactured contrary to Part 3-3; or

(j) if the goods have been manufactured in Australia, or imported into Australia, solely for export—a relevant authority of the country to which the goods are to be exported has not confirmed its willingness to accept the goods and:

(i) the goods have been refused registration or listing for supply in Australia; or

(ii) the Secretary requires such a confirmation for a reason other than because the goods have been refused registration or listing; or

(k) the goods do not comply with prescribed quality or safety criteria; or

(m) the goods contain substances that are prohibited imports for the purposes of the Customs Act 1901; or

(n) one or more of the manufacturers of the goods are not nominated as manufacturers of the goods in the application.

(1A) To avoid doubt, if:

(a) an application is made for the listing of therapeutic goods in relation to a person in accordance with section 23; and
(b) the application is accompanied by the certificate required under subsection 26B(1); and
(c) the other requirements in subsection (1) are met;
the Secretary must list the goods under subsection (1) without inquiring into the correctness of the certificate.

(1B) Civil proceedings do not lie against the Secretary (or a delegate of the Secretary) in respect of loss, damage or injury of any kind suffered by another person as a result of the Secretary (or the delegate) listing therapeutic goods in relation to a person in reliance on a certificate required under subsection 26B(1).

(2) In making a decision for the purposes of paragraph (1)(g), the matters that may be taken into account include:
(a) whether the applicant has provided:
   (i) if the goods are not therapeutic devices and a step in the manufacture of the goods has been carried out in a country that is a member of the European Community or a member of EFTA—an EC/EFTA attestation of conformity in relation to the goods; or
   (ia) if the goods are not therapeutic devices and a step in the manufacture of the goods has been carried out in a country declared by the Minister under section 3B to be covered by a non-EC/EFTA MRA—a non-EC/EFTA attestation of conformity, for the non-EC/EFTA MRA, in relation to the goods; or
   (ii) in any other case—an acceptable form of evidence from a relevant overseas authority establishing that the manufacture of the goods is of an acceptable standard; and
(b) whether the applicant has agreed to provide, where the Secretary considers inspection of the manufacturing procedures used in the manufacture of the goods to be necessary:
   (i) funds for the carrying out of that inspection by the Department; and
   (ii) evidence that the manufacturer has agreed to such an inspection.
(2A) If therapeutic goods are exempt from the operation of Part 3-3 or a
person is exempt from the operation of that Part in relation to the
manufacture of the goods, subsection (1) has effect, in relation to
the goods, as if paragraph (h) were omitted.

(2B) If a person is exempt from the operation of Part 3-3 in relation to a
step in the manufacture of therapeutic goods, subsection (1) has
effect, in relation to the goods, as if the reference in paragraph (h)
to Part 3-3 were a reference to that Part to the extent that it applies
to that person in relation to the manufacture of the goods.

(2C) If:
(a)  therapeutic goods were made outside Australia; and
(b)  had the goods been made in Australia, they would have been
     exempt from the operation of Part 3-3;
subsection (1) has effect, in relation to the goods, as if
paragraph (g) were omitted.

(2D) A decision for the purposes of paragraph (1)(g) may also take into
account any information provided to the Secretary by a health
authority of a Convention country and relating to:
(a)  the general standards of manufacturing practice of a
     particular manufacturer; or
(b)  the specific standards of manufacture or control adopted by a
     particular manufacturer in relation to particular goods.

(2E) For the purposes of subsection (2D), a Convention country is a
country that is a party to the Mutual Recognition Convention.

(2F) Information referred to in subsection (2D) and provided in
accordance with the Mutual Recognition Convention is to be
treated as equivalent to information obtained as a result of an
inspection under Part 3-3 of this Act.

(3) Where an application is made, the Secretary must notify the
applicant in writing of his or her decision on the application within
28 days of the making of the decision and, in the case of a decision
not to list the goods, of the reasons for the decision.
(4) As soon as practicable after an applicant has been informed that therapeutic goods in respect of which an application was made are acceptable for listing, the Secretary must give to the applicant a certificate of listing of the goods, and the listing of the goods commences on the day specified for the purpose in the certificate.

26AA Listing of therapeutic device to which EC/EFTA attestation of conformity applies

(1) If:
   (a) an application is made in accordance with section 23 for the listing of a therapeutic device in relation to a person; and
   (b) the applicant gives to the Secretary an EC/EFTA attestation of conformity as to the matters specified in paragraphs 26(1)(c) to (m) in relation to the device;

the Secretary must list the device in relation to the person unless the Secretary considers that the device may compromise the health or safety of users.

(2) The Secretary must notify the applicant in writing of his or her decision within 28 days of the making of the decision. If the Secretary decides not to list the device, the notice must contain the reasons for that decision.

(3) If the Secretary decides to list the device, the Secretary must:
   (a) include the device in the Register; and
   (b) give to the applicant a certificate of listing.

(4) The listing of the device commences on the day specified for the purpose in the certificate of listing.

26A Listing of certain medicines

(1) If:
   (a) an application is made for the listing of medicine in relation to a person in accordance with section 23; and
   (b) the application is accompanied by the certificate required under subsection 26B(1); and
   (c) the requirements of subsection (2) and (where applicable) subsection (3) have been complied with; and
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(d) the medicine is not export only medicine; and
(e) the medicine is not one that has previously had its registration or listing cancelled;
the Secretary must list the medicine in relation to the person.

(1A) To avoid doubt, if:
(a) an application is made for the listing of a medicine in relation to a person in accordance with section 23; and
(b) the application is accompanied by the certificate required under subsection 26B(1); and
(c) the other requirements in subsection (1) are met;
the Secretary must list the medicine under subsection (1) without inquiring into the correctness of the certificate.

(1B) Civil proceedings do not lie against the Secretary (or a delegate of the Secretary) in respect of loss, damage or injury of any kind suffered by another person as a result of the Secretary (or the delegate) listing a medicine in relation to a person in reliance on a certificate required under subsection 26B(1).

(2) The applicant must certify that:
(a) the medicine is eligible for listing; and
(b) the medicine is safe for the purposes for which it is to be used; and
(c) the presentation of the medicine is not unacceptable; and
(d) the medicine conforms to every standard (if any) applicable to the medicine and to every requirement (if any) relating to advertising applicable under Part 5-1 or under the regulations; and
(e) if the medicine has been manufactured in Australia—each step in the manufacture of the medicine has been carried out by a person who is the holder of a licence to carry out that step granted under section 38; and
(f) the medicine complies with all prescribed quality or safety criteria; and
(g) the medicine does not contain substances that are prohibited imports for the purposes of the *Customs Act 1901*; and
(h) all the manufacturers of the medicine are nominated as manufacturers in the application; and
(i) the applicant has, with manufacturers of the medicine who are manufacturers of the prescribed kind, written agreements containing such matters as are prescribed; and
(j) the applicant holds information or evidence to support any claim that the applicant makes relating to the medicine; and
(k) the information included in or with the application is correct.

(3) Subject to subsection (7), if a step in the manufacture of the medicine has been carried out outside Australia, the Secretary must have certified, prior to the application being made, that the manufacturing and quality control procedures used in each such step are acceptable.

(4) In deciding whether so to certify for the purposes of subsection (3), the matters that may be taken into account include:
(a) whether the applicant has provided:
   (i) if a step in the manufacture of the medicine has been carried out in a country that is a member of the European Community or a member of EFTA—an EC/EFTA attestation of conformity in relation to the medicine; or
   (ia) if a step in the manufacture of the medicine has been carried out in a country declared by the Minister under section 3B to be covered by a non-EC/EFTA MRA—a non-EC/EFTA attestation of conformity, for the non-EC/EFTA MRA, in relation to the medicine; or
   (ii) in any other case—an acceptable form of evidence from a relevant overseas authority establishing that the manufacture of the medicine is of an acceptable standard; and
(b) whether the applicant has agreed to provide, if the Secretary considers inspection of the manufacturing procedures used in the manufacture of the medicine to be necessary:
   (i) funds for the carrying out of that inspection by the Department; and
   (ii) evidence that the manufacturer has agreed to such an inspection; and
(c) whether the applicant has complied with any requirements made by the Secretary under section 31 in relation to the manufacture or preparation of the medicine.
(5) If a medicine is exempt from the operation of Part 3-3 or a person is exempt from the operation of that Part in relation to the manufacture of the medicine, subsection (2) has effect, in relation to the medicine, as if paragraph (2)(e) were omitted.

(6) If a person (the manufacturer) is exempt from the operation of Part 3-3 in relation to a step in the manufacture of a medicine, subsection (2) has effect, in relation to the medicine, as if the reference in paragraph (2)(e) to a person who is the holder of a licence were a reference to the manufacturer to the extent that Part 3-3 applies to the manufacturer in relation to the manufacture of the medicine.

(7) If:
   (a) a medicine was made outside Australia; and
   (b) had the medicine been made in Australia, it would have been exempt from the operation of Part 3-3;
   subsection (3) does not apply in relation to the medicine.

(9) As soon as practicable after a medicine has been listed under this section, the Secretary must give to the applicant a certificate of listing of the medicine. The listing of the medicine commences on the day specified for the purpose in the certificate.

26B Certificates required in relation to patents

(1) The certificate required by this subsection is either:
   (a) a certificate to the effect that the applicant, acting in good faith, believes on reasonable grounds that it is not marketing, and does not propose to market, the therapeutic goods in a manner, or in circumstances, that would infringe a valid claim of a patent that has been granted in relation to the therapeutic goods; or
   (b) a certificate to the effect that:
      (i) a patent has been granted in relation to the therapeutic goods; and
      (ii) the applicant proposes to market the therapeutic goods before the end of the term of the patent; and
(iii) the applicant has given the patentee notice of the application for registration or listing of the therapeutic goods under section 23.

The certificate must be signed by, or on behalf of, the applicant and must be in a form approved by the Secretary.

(2) A person is guilty of an offence if:
(a) the person gives a certificate required under subsection (1); and
(b) the certificate is false or misleading in a material particular.

Maximum penalty: 1,000 penalty units.

(3) For the purposes of this section, a patent is taken to have been granted in relation to therapeutic goods if marketing the goods without the authority of the patentee would constitute an infringement of the patent.

(4) In this section:

patent has the same meaning as in the Patents Act 1990.

26C Certificates required in relation to patent infringement proceedings

(1) This section applies if:
(a) a person gives a certificate required under subsection 26B(1) in relation to therapeutic goods; and
(b) another person (the second person) intends to commence proceedings under the Patents Act 1990 against the person referred to in paragraph (1)(a) for infringement of a patent that has been granted in relation to the therapeutic goods (the proceedings).

(2) The second person, before the date upon which the proceedings are commenced, must give to the Secretary and to the person referred to in paragraph (1)(a) the certificate required by subsection (3).

(3) The certificate required by this subsection is a certificate to the effect that the proceedings:
(a) are to be commenced in good faith; and
(b) have reasonable prospects of success; and
(c) will be conducted without unreasonable delay.

The certificate must be signed by, or on behalf of, the second person and must be in a form approved by the Secretary.

(4) For the purpose of paragraph (3)(b), proceedings have reasonable prospects of success if:

(a) the second person had reasonable grounds in all the circumstances known to the second person, or which ought reasonably to have been known to the second person (in addition to the fact of grant of the patent), for believing that he or she would be entitled to be granted final relief by the court against the person referred to in paragraph (1)(a) for infringement by that person of the patent; and

(b) the second person had reasonable grounds in all the circumstances known to the second person, or which ought reasonably to have been known to the second person (in addition to the fact of grant of the patent), for believing that each of the claims, in respect of which infringement is alleged, is valid; and

(c) the proceedings are not otherwise vexatious or unreasonably pursued.

(5) The person referred to in paragraph (1)(a), with leave of the court, or the Attorney-General, may apply to a prescribed court for an order that the second person pay to the Commonwealth a pecuniary penalty if the second person gives a certificate required under subsection (3) and:

(a) the certificate is false or misleading in a material particular; or

(b) the second person breaches an undertaking given in the certificate.

Maximum penalty: $10,000,000.

(6) When determining the extent of a pecuniary penalty to be ordered pursuant to subsection (5), the court must take into account:

(a) any profit obtained by the second person; and

(b) any loss or damage suffered by any person;

by reason of the second person exploiting the patent during the proceedings.
(7) For the avoidance of doubt, subsection (6) does not limit the matters the court may take into account when determining a pecuniary penalty ordered pursuant to subsection (5).

(8) If:
   (a) the second person has sought and obtained in the proceedings an interlocutory injunction restraining the person referred to in paragraph (1)(a) from infringing a patent; and
   (b) section 26D does not apply; and
   (c) a prescribed court declares that the second person has given a certificate required under subsection (3); and
   (d) a prescribed court declares that:
      (i) the certificate is false or misleading in a material particular; or
      (ii) the second person has breached an undertaking given in the certificate;

the prescribed court may, pursuant to this section, order that the second person pay to the Commonwealth, a State or a Territory compensation for any damages sustained or costs incurred by the Commonwealth, a State or a Territory as a result of the grant of the interlocutory injunction.

(9) In this section:

   prescribed court has the same meaning as in the Patents Act 1990.

26D Requirements for interlocutory injunction

(1) This section applies where:
   (a) an applicant gives notice to a patentee in accordance with subparagraph 26B(1)(b)(iii); and
   (b) the patentee and/or its exclusive licensee (in this section the party or parties is or are referred to as the patentee) applies to a prescribed court for an interlocutory injunction to restrain the applicant from marketing the therapeutic goods the subject of the application on the ground that such conduct will constitute an infringement of its patent.

(2) An application for interlocutory relief in accordance with subsection (1) may not be instituted unless the patentee has first
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notified the Attorney-General of the Commonwealth, or of a State or of a Territory, in writing of the application.

(3) The Attorney-General of the Commonwealth shall be deemed to be a party to any proceedings commenced in accordance with subsection (1) unless the Attorney-General gives written notice to the prescribed court that he or she does not desire to be a party.

(4) If an interlocutory injunction is granted pursuant to an application made as described in subsection (1) and:

(a) the patentee subsequently discontinues the principal proceedings without the consent of the other parties thereto; or

(b) the principal proceedings are dismissed; and

(c) in either case, the prescribed court declares that:

(i) the patentee did not have reasonable grounds, in all the circumstances known to the patentee or which ought reasonably have been known to the patentee:

(A) to believe that it would be granted final relief by the prescribed court against the applicant referred to in paragraph (1)(a) for infringement by that person of the patent; or

(B) (in addition to the fact of grant of the patent), for believing that each of the claims, in respect of which infringement is alleged in the proceedings, would have a reasonable prospect of being held to be valid if challenged by the applicant referred to in paragraph (1)(a); or

(ii) the application for the interlocutory injunction was otherwise vexatious or not reasonably made or pursued;

the prescribed court may, in addition to any other relief which it believes should be granted to any person, make any of the orders described in subsection (5).

(5) If the prescribed court makes a declaration pursuant to paragraph (4)(c), the prescribed court may, pursuant to the usual undertaking as to damages given by the patentee to the prescribed court to obtain the interlocutory injunction:

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(a) assess and award compensation to the applicant referred to in paragraph (1)(a) against whom the interlocutory injunction was made:

(i) on the basis of an account of the gross profits of the patentee arising from the sale by it in Australia of the therapeutic goods the subject of the interlocutory injunction, during the period of the interlocutory injunction, without requiring the said applicant to establish or quantify its actual loss; or

(ii) on such other basis as the court determines to be appropriate; and

(b) award to the Commonwealth compensation for any damages sustained, or costs incurred, by it as a result of the grant of the interlocutory injunction; and

(c) award to a State or a Territory compensation for any damages sustained, or costs incurred, by it as a result of the grant of the interlocutory injunction.

(6) In this section:

prescribed court has the same meaning as in the Patents Act 1990.

27 Registration or listing number  

(1) Where the Secretary includes therapeutic goods (other than grouped therapeutic goods) in the Register, the Secretary is to assign a unique registration or listing number to the goods.

(2) Where the Secretary includes grouped therapeutic goods in the Register, the Secretary is to assign a single, unique registration or listing number to the grouped therapeutic goods.

28 Conditions of registration or listing

(1) Where the Secretary includes therapeutic goods in the Register in relation to a person the Secretary may, in writing, impose conditions on the registration or listing of those goods.

(2) Conditions referred to in subsection (1) may relate to:

(a) the manufacture of the goods; or
(b) the custody, use, supply, disposal or destruction of the goods; or
(c) the keeping of records relating to the goods; or
(d) matters dealt with in, or matters additional to matters dealt with in, standards applicable to the goods; or
(e) such other matters relating to the goods as the Secretary thinks appropriate.

(3) The Secretary may, by notice in writing given to the person in relation to whom therapeutic goods are registered or listed, impose new conditions on the registration or listing or vary or remove existing conditions.

(3A) The Secretary’s power under subsection (3) may be exercised at the request of the person concerned or of the Secretary’s own motion.

(4) The imposition or variation of a condition under subsection (3) takes effect:
(a) if the notice states that the action is necessary to prevent imminent risk of death, serious illness or serious injury—on the day on which the notice is given to the person; or
(b) in any other case—on the day specified for the purpose in the notice, being a day not earlier than 28 days after the notice is given to the person.

(5) In addition to any conditions imposed under subsection (1) or (3), the registration or listing of therapeutic goods is subject to the conditions that the person in relation to whom the goods are registered or listed will:
(a) allow an authorised person:
   (i) to enter, at any reasonable time, premises at which the person deals with the goods; and
   (ii) while on those premises, to inspect those premises and therapeutic goods at those premises and to take samples of goods of that kind; and
(b) if requested to do so by an authorised person, produce to the person such documents relating to the goods as the person requires and allow the person to copy the documents; and
(c) in relation to each batch of the goods—keep a record, at least until the end of the period of 12 months after the expiry date.
for the goods, of all of the manufacturers involved in the manufacture of that batch; and

(d) if requested to do so by an authorised person, make any such record available to the authorised person for inspection:
   (i) at or before the time the authorised person requests, or (if the authorised person requests) immediately; and
   (ii) either in electronic form or in paper form, as the authorised person requests; and

(e) comply, in relation to the goods, with any reporting requirements that are prescribed; and

(f) if a manufacturer who was not nominated as a manufacturer of the goods in the application for the registration or listing of the goods becomes a manufacturer of the goods—inform the Secretary in writing of that fact, no later than 10 working days after the manufacturer becomes a manufacturer of the goods; and

(g) if premises that were not nominated as premises to be used in the manufacture of the goods in the application become premises used in the manufacture of the goods—inform the Secretary in writing of that fact, no later than 10 working days after the premises are first used for that purpose.

(5A) In addition to any conditions imposed under subsection (1), (3) or (5), the listing of a medicine under section 26A is subject to a condition that the person in relation to whom the medicine is listed will deliver a reasonable number of samples of the medicine if the Secretary so requests:
   (a) within the period specified in the request; and
   (b) in accordance with any other requirements specified in the request.

The period specified in the request must include at least 10 working days.

(6) If:
   (a) in, or in connection with, an application for the listing of therapeutic goods, a claim is made by the applicant in relation to the goods; and
   (b) the claim is included in the Register in respect of the goods;

the listing of the goods is subject to the following conditions:
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(c) a condition that the sponsor of the goods had, at the time when the claim was made, information or evidence that supported the claim and complied with the requirements (if any) of the regulations;
(d) a condition that the sponsor retains the information or evidence at all times while the goods remain listed;
(e) a condition that, at any time while the goods remain listed, the sponsor will, if asked to do so by the Secretary, give the information or evidence to the Secretary.

(7) The regulations may prescribe the amount, standard or type of information or evidence required for the purposes of paragraph (6)(c).

29  Duration of registration or listing

Where goods are included in the Register in relation to a person, the goods remain so included until their registration or listing is cancelled under this Part.

29A  Notification of adverse effects etc. of goods

(1) As soon as a person in relation to whom therapeutic goods are registered or listed becomes aware of information of a kind mentioned in subsection (2) relating to the goods, the person must give the information to the Secretary in writing.

Maximum penalty: 400 penalty units.

(2) The information with which subsection (1) is concerned is information of the following kinds:
   (a) information that contradicts information already furnished by the person under this Act;
   (b) information that indicates that the use of the goods in accordance with the recommendations for their use may have an unintended harmful effect;
   (c) information that indicates that the goods, when used in accordance with the recommendations for their use, may not be as effective as the application for registration or listing of the goods or information already furnished by the person under this Act suggests;

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(d) information that indicates that the quality, safety or efficacy of the goods is unacceptable.

**29B Notification of adverse effects etc. where application withdrawn or lapses**

(1) If an application for registration or listing of goods is withdrawn or lapses, the Secretary may give the applicant written notice requiring the applicant:

(a) to inform the Secretary in writing whether the applicant is aware of any information of a kind mentioned in subsection 29A(2) relating to the goods; and

(b) if the applicant is aware of such information, to give the information to the Secretary in writing.

(2) Notice under subsection (1) may be given within 14 days after an application is withdrawn or lapses.

(3) A person must comply with the requirements of a notice under subsection (1) within 30 days after the notice is given to the person.

Maximum penalty: 400 penalty units.

(4) A person must not, in purported compliance with a notice under subsection (1), give information that is false or misleading in a material particular.

Maximum penalty: 400 penalty units.

**30 Cancellation of registration or listing**

(1) The Secretary may, by notice in writing given to a person in relation to whom therapeutic goods are included in the Register, cancel the registration or listing of the goods if:

(a) it appears to the Secretary that failure to cancel the registration or listing would create an imminent risk of death, serious illness or serious injury; or

(b) the goods become exempt goods; or

(c) the person requests in writing the cancellation of the registration or listing; or
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(d) the goods contain substances that are prohibited imports for the purposes of the *Customs Act 1901*; or
(da) the person has refused or failed to comply with the condition to which the inclusion of the goods is subject under paragraph 28(5)(d):
   (i) if the person was requested under that paragraph to make the record in question available at or before a requested time—before the end of the period of 24 hours after that time; or
   (ii) if the person was requested under that paragraph to make the record in question available immediately—within 24 hours after the request was made; or
(e) in the case of a medicine listed under section 26A, it appears to the Secretary that any of the certifications under paragraph 26A(2)(a), (e) or (g) are incorrect or (if applicable) the requirements under subsection 26A(3) are not fulfilled; or
(f) both of the following apply:
   (i) under the regulations, an authority constituted by or under the regulations gives a direction to, or makes a requirement of, the person in relation to an advertisement of the goods to ensure that advertising complies with the Therapeutic Goods Advertising Code;
   (ii) the person does not comply with the direction or requirement.

(1A) The Secretary may, by notice in writing given to a person in relation to whom a medicine is listed under section 26A, cancel the listing of the medicine if:
   (a) the medicine is not eligible for listing; or
   (b) the medicine is exempt; or
   (c) there is a serious breach, involving the medicine, of the requirements relating to advertising applicable under Part 5-1 or under the regulations, and the Secretary is satisfied that:
      (i) the breach is significant; and
      (ii) as a result of the breach, the presentation of the medicine is misleading to a significant extent.

(1B) However, paragraph (1A)(c) does not apply to medicines that are manufactured in Australia for export only, or are imported into Australia for export only.
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(1C) The Secretary may, by notice in writing given to a person in relation to whom a medicine is listed under section 26A, cancel the listing of the medicine if:

(a) the Secretary, under section 31, gives to the person a notice requiring the person to give to the Secretary information or documents relating to the medicine; and

(b) the notice is given for the purposes of ascertaining whether the medicine should have been listed; and

(c) the person fails to comply with the notice within 20 working days after the notice is given.

(2) Subject to subsection (3), the Secretary may, by notice in writing given to a person in relation to whom therapeutic goods are included in the Register, cancel the registration or listing of the goods if:

(a) it appears to the Secretary that the quality, safety or efficacy of the goods is unacceptable; or

(b) the goods have changed so that they have become separate and distinct from the goods as so included; or

(ba) in the case of a medicine listed under section 26A, it appears to the Secretary that any of the certifications under paragraph 26A(2)(b), (c), (d), (f), (h), (i), (j) or (k) are incorrect; or

(c) the sponsor has refused or failed to comply with a condition to which the inclusion of the goods is subject (other than the condition under paragraph 28(5)(d)); or

(ca) the person has contravened subsection 29A(1) in relation to the goods; or

(d) the goods become required to be included in the other part of the Register; or

(e) the goods do not conform to a standard applicable to the goods or to a requirement relating to advertising applicable to the goods under Part 5-1 or under the regulations; or

(f) the annual registration or listing charge is not paid within 28 days after it becomes payable.

(3) Where the Secretary proposes to cancel the registration or listing of goods in relation to a person under subsection (2) otherwise than as a result of a failure to pay the annual registration or listing charge, the Secretary must:
Section 30C

(a) inform the person in writing that the Secretary proposes to cancel that registration or listing and set out the reasons for that proposed action; and
(b) give the person a reasonable opportunity to make submissions to the Secretary in relation to the proposed action.

(4) Where a person makes submissions in accordance with paragraph (3)(b), the Secretary is not to make a decision relating to the cancellation until the Secretary has taken the submissions into account.

(4A) The Secretary must, by notice in writing given to a person in relation to whom therapeutic goods are included in the Register, cancel the registration of the goods if the Secretary becomes aware that protected information was used when evaluating the goods for registration.

(5) Where the Secretary cancels the registration or listing of goods in relation to a person, the goods cease to be registered or listed:
(a) if the cancellation is effected under subsection (1), (1A) or (1C)—on the day on which the notice of cancellation is given to the person; or
(b) in any other case—on such later day as is specified in the notice.

30C Consultation with Gene Technology Regulator

(1) This section applies to an application for listing or registration of a therapeutic good under section 23 if the therapeutic good is, or contains, a GM product.

(2) Subject to subsection (5), the Secretary must give written notice to the Gene Technology Regulator:
(a) stating that the application has been made; and
(b) requesting the Gene Technology Regulator to give advice about the application.

(3) If the Secretary gives the Gene Technology Regulator a notice under subsection (2), the Gene Technology Regulator may give written advice to the Secretary about the application.

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(4) The advice is to be given within the period specified in the notice.

(5) If an advice from the Gene Technology Regulator is in force under section 30D in relation to a class of therapeutic goods, the Secretary is not required to notify the Regulator under this section in relation to an application for listing or registration of a therapeutic good belonging to that class.

30D Secretary may seek advice about classes of GM products

(1) The Secretary may request advice from the Gene Technology Regulator in relation to therapeutic goods that consist of, or that contain, a GM product belonging to a class of GM products specified in the request.

(2) A request for advice under subsection (1) must specify the matters to which the advice is to relate.

(3) If the Secretary requests advice from the Gene Technology Regulator under subsection (1), the Gene Technology Regulator may provide written advice in relation to the matters specified in the request.

(4) If theGene Technology Regulator gives advice to the Secretary under subsection (3), the advice remains in force until it is withdrawn by the Gene Technology Regulator by written notice given to the Secretary.

30E Secretary to take advice into account

If the Secretary receives advice from the Gene Technology Regulator:

(a) in response to a notice under section 30C within the period specified in the notice; or

(b) under section 30D;

the Secretary must:

(c) ensure that the advice is taken into account in making a decision on the application to which the notice relates, or on an application to which the advice under section 30D relates, as the case requires; and
Section 30E

(d) inform the Gene Technology Regulator of the decision on the application.

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Division 2A—Public notification and recovery of therapeutic goods

30EA  Public notification and recovery of therapeutic goods

(1) The Secretary may, in writing, impose requirements, relating to therapeutic goods, on a person if:

(a) any of the circumstances referred to in the second column of an item in the following table occur in relation to the goods; and

(b) the person is referred to in the third column of that item.

<table>
<thead>
<tr>
<th>Item</th>
<th>Circumstance relating to therapeutic goods</th>
<th>Person subject to requirements</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>The goods are supplied while they are registered goods or listed goods, but they do not conform with a standard applicable to the goods</td>
<td>The person in relation to whom the goods are included in the Register</td>
</tr>
<tr>
<td>2.</td>
<td>The goods are supplied while they are registered goods or listed goods, but the manufacturing principles have not been observed in the manufacture of the goods</td>
<td>The person in relation to whom the goods are included in the Register</td>
</tr>
<tr>
<td>3.</td>
<td>The goods are supplied while: (a) they are exempt goods; or (b) they are exempt under section 18A; or (c) they are the subject of an approval or authority under section 19; or (d) they are the subject of an approval under section 19A; but they do not conform with a standard applicable to the goods</td>
<td>The person supplying the goods</td>
</tr>
</tbody>
</table>
Circumstances in which requirements may be imposed

<table>
<thead>
<tr>
<th>Item</th>
<th>Circumstance relating to therapeutic goods</th>
<th>Person subject to requirements</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.</td>
<td>The goods are supplied while:</td>
<td>The person supplying the goods</td>
</tr>
<tr>
<td></td>
<td>(a) they are exempt goods; or</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(b) they are exempt under section 18A;</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(c) they are the subject of an approval</td>
<td></td>
</tr>
<tr>
<td></td>
<td>or authority under section 19; or</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(d) they are the subject of an approval</td>
<td></td>
</tr>
<tr>
<td></td>
<td>under section 19A; but the manufacturing</td>
<td></td>
</tr>
<tr>
<td></td>
<td>principles have not been observed in</td>
<td></td>
</tr>
<tr>
<td></td>
<td>the manufacture of the goods</td>
<td></td>
</tr>
<tr>
<td>5.</td>
<td>The goods are supplied in contravention</td>
<td>The person supplying the goods</td>
</tr>
<tr>
<td></td>
<td>of subsection 20(1) or 42E(1)</td>
<td></td>
</tr>
<tr>
<td>6.</td>
<td>The goods are supplied while they are</td>
<td>The person in relation to whom</td>
</tr>
<tr>
<td></td>
<td>registered goods or listed goods, but</td>
<td>the goods are included in the</td>
</tr>
<tr>
<td></td>
<td>one or more steps in the manufacture of</td>
<td>Register</td>
</tr>
<tr>
<td></td>
<td>the goods has been carried out by a</td>
<td></td>
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<tr>
<td></td>
<td>manufacturer while the manufacturer</td>
<td></td>
</tr>
<tr>
<td></td>
<td>did not hold a licence that was in force</td>
<td></td>
</tr>
<tr>
<td>7.</td>
<td>The registration or listing of the goods</td>
<td>The person in relation to whom</td>
</tr>
<tr>
<td></td>
<td>has been cancelled under this Part</td>
<td>the goods were included in the</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Register</td>
</tr>
</tbody>
</table>

(2) The requirements may be one or more of the following:

(a) to take specified steps, in the specified manner and within such reasonable period as is specified, to recover therapeutic goods that have been distributed;

(b) to inform the public or a specified class of persons, in the specified manner and within such reasonable period as is specified, to the effect that the circumstances referred to in paragraph (1)(a) have occurred in relation to therapeutic goods;

(c) to publish, in the specified manner and within such reasonable period as is specified, specified information, or information of a specified kind, relating to the manufacture or distribution of therapeutic goods.

(3) If the circumstances referred to in paragraph (1)(a) apply only to a batch of therapeutic goods, the Secretary may limit the imposition of the requirements to the therapeutic goods included in that batch.
(4) A requirement to recover therapeutic goods under this section does not apply to therapeutic goods that cannot be recovered because they have been administered to, or applied in the treatment of, a person.

30EB Publication of requirements

The Secretary must cause to be published in the Gazette, as soon as practicable after imposing a requirement under section 30EA, a notice setting out particulars of the requirement.

30EC Non-compliance with requirements

A person is guilty of an offence if:
(a) the person does an act, or omits to do an act; and
(b) the act or omission constitutes a contravention of a requirement imposed on the person under section 30EA.

Maximum penalty: 60 penalty units.

30ED Power of cancellation unaffected

Imposition of a requirement under section 30EA does not affect the Secretary’s power to cancel the registration or listing of therapeutic goods under this Part.
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30F  Goods exempt under section 18A not conforming to standards etc.

(1) This section applies if:
   (a) therapeutic goods of a particular kind are exempt under section 18A; and
   (b) a person supplies a batch of goods of that kind; and
   (c) the Secretary is satisfied that the goods included in that batch:
      (i) do not conform to a standard applicable to goods of that kind; or
      (ii) are otherwise not fit to be used for their intended purposes.

(2) The Secretary may, by written notice given to the person, require the person to take steps to recover the goods included in that batch (except any of those goods that cannot be recovered because they have been administered to, or applied in the treatment of, a person or animal).

(3) The notice may specify one or more of the following requirements:
   (a) the steps to be taken to recover the goods;
   (b) the manner in which the steps are to be taken;
   (c) a reasonable period within which the steps are to be taken.

(4) The Secretary must, as soon as practicable after giving the notice, cause particulars of it to be published in the Gazette.

(5) A person commits an offence if:
   (a) the Secretary gives a notice to the person under subsection (2); and
   (b) the notice specifies a particular requirement mentioned in subsection (3); and
   (c) the person fails to comply with that requirement.

   Maximum penalty: Imprisonment for 12 months or 60 penalty units, or both.
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(6) For the purposes of an offence against subsection (5), strict liability applies to the following physical elements of circumstances:
   (a) that the notice concerned is given under subsection (2);
   (b) that the particular requirement concerned is a requirement mentioned in subsection (3).

Note: For strict liability, see section 6.1 of the Criminal Code.

30G Disposal of unused goods exempt under section 18A

(1) This section applies to particular therapeutic goods if:
   (a) an exemption in relation to those goods under section 18A ceases to have effect otherwise than because those goods have become registered goods or listed goods (see paragraph 18A(5)(a)); and
   (b) those goods have not been used before the exemption so ceases to have effect.

(2) The Secretary may arrange for the disposal of any of those goods in accordance with the regulations.

(3) Regulations made for the purposes of subsection (2) may set out the methods by which those goods are to be stored, supplied, destroyed, exported or otherwise disposed of.

(4) A method set out in the regulations under subsection (3) must not enable or permit any benefit to be conferred on a person (including the Commonwealth) other than the owner of those goods.

30H Record for goods exempt under section 18A

(1) A person commits an offence if:
   (a) there are therapeutic goods that are exempt under section 18A; and
   (b) a condition of the exemption:
      (i) requires the person to keep a record about those goods; or
      (ii) specifies the manner in which the person must keep the record; and
   (c) the person does an act or omits to do an act in relation to those goods; and

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(d) the act or omission results in the breach of that condition of
the exemption.

Maximum penalty:  240 penalty units.

(2) Strict liability applies to paragraph (1)(b).

Note: For strict liability, see section 6.1 of the Criminal Code.

(3) A person commits an offence if:
    (a) there are therapeutic goods that are exempt under
        section 18A; and
    (b) a condition of the exemption:
        (i) requires the person to keep a record about those goods;
        or
        (ii) specifies the manner in which the person must keep the
            record; and
    (c) the person does an act or omits to do an act in relation to
        those goods; and
    (d) the act or omission results in the breach of that condition of
        the exemption.

Maximum penalty:  60 penalty units.

(4) An offence under subsection (3) is an offence of strict liability.

Note: For strict liability, see section 6.1 of the Criminal Code.

31 Secretary may require information

(1) The Secretary may, by notice in writing given to a person who is
    an applicant for the registration of therapeutic goods or in relation
    to whom therapeutic goods are registered, require the person to
give to the Secretary, within such reasonable time as is specified in
the notice and in such form as is specified in the notice,
information or documents relating to one or more of the following:
    (a) the formulation of the goods;
    (b) the composition of the goods;
    (c) the design specifications of the goods;
    (d) the quality of the goods;
    (e) the method and place of manufacture or preparation of the
        goods and the procedures employed to ensure that proper
standards are maintained in the manufacture and handling of the goods;
(f) the presentation of the goods;
(g) the safety and efficacy of the goods for the purposes for which they are to be used;
(h) the conformity of the goods to a requirement relating to advertising applicable under Part 5-1 or under the regulations;
(j) the regulatory history of the goods in another country;
(k) any other matter prescribed by the regulations for the purposes of this paragraph in relation to goods of that kind.

(2) The Secretary may, by notice in writing given to a person who is an applicant for the listing of therapeutic goods or in relation to whom therapeutic goods are listed, require the person to give to the Secretary, within such reasonable time as is specified in the notice, information or documents relating to one or more of the following:
(a) the formulation of the goods;
(b) the composition of the goods;
(c) the design specifications of the goods;
(d) the method and place of manufacture or preparation of the goods and the procedures employed to ensure that proper standards are maintained in the manufacture and handling of the goods;
(e) the presentation of the goods;
(f) the safety of the goods for the purposes for which they are to be used;
(g) the conformity of the goods to a standard applicable to the goods, or to a requirement relating to advertising applicable to the goods under Part 5-1 or under the regulations;
(h) any other matter prescribed by the regulations for the purposes of this paragraph in relation to goods of that kind.

(3) An approval of a form may require or permit information to be given in accordance with specified software requirements:
(a) on a specified kind of data processing device; or
(b) by way of a specified kind of electronic transmission.
(4) A person in relation to whom therapeutic goods are registered or listed must not fail to comply with a notice given to the person under this section.

Maximum penalty: 60 penalty units.

(4A) Subsection (4) does not apply if the person has a reasonable excuse.

Note: The defendant bears an evidential burden in relation to the matter in subsection (4A). See subsection 13.3(3) of the Criminal Code.

(5) An offence under subsection (4) is an offence of strict liability.

Note: For strict liability, see section 6.1 of the Criminal Code.

(6) A person in relation to whom a medicine is listed under section 26A must not, in purported compliance with a notice under this section relating to the medicine, provide information that is false or misleading in a material particular.

Maximum penalty: 400 penalty units.

31A Secretary may require information etc. about goods exempt under section 18

Exempt goods for use for experimental purposes in humans

(1) If therapeutic goods are exempt under subsection 18(1) from the operation of this Part (except this section and sections 31C to 31F) to allow for their use for experimental purposes in humans, the Secretary may give the sponsor of the goods a written notice requiring the sponsor to give to the Secretary specified information or documents relating to one or more of the following:

(a) the supply of the goods;

(b) the handling of the goods;

(c) the monitoring of the supply of the goods;

(d) the results of the supply of the goods;

(e) any other matter prescribed by the regulations for the purposes of this paragraph in relation to goods of that kind.
Statement by medical practitioner about medicine

(2) If a medicine is exempt under subsection 18(1) from the operation of this Part (except this section and sections 31C to 31F) because a medical practitioner has signed a statement in accordance with regulation 12A of the Therapeutic Goods Regulations 1990, the Secretary may give the medical practitioner a written notice requiring the medical practitioner to give to the Secretary specified information or documents relating to one or more of the following:
   (a) the condition of the person to whom the medicine is to be given or is given;
   (b) the supply of the medicine;
   (c) the handling of the medicine;
   (d) the monitoring of the supply of the medicine;
   (e) the results of the supply of the medicine;
   (f) any other matter prescribed by the regulations for the purposes of this paragraph in relation to medicines of that kind.

Compliance period

(3) A notice under subsection (1) or (2) must specify a reasonable period within which the person to whom the notice is given must comply with it. The period must be at least 14 days starting on the day on which the notice is given.

31AA Secretary may require information etc. about goods exempt under section 18A

(1) This section applies to a person who is required to comply with a condition of an exemption of therapeutic goods under section 18A.

(2) The Secretary may, by written notice given to the person, require the person to give to the Secretary specified information or documents relating to one or more of the following:
   (a) the supply of any of those goods;
   (b) the handling of any of those goods;
   (c) the monitoring of the supply of any of those goods;
   (d) the results of the supply of any of those goods;
(e) any other matter prescribed by the regulations for the purposes of this paragraph.

**Compliance period**

(3) The notice must specify a reasonable period within which the person must comply with it. The period must be at least 14 days starting on the day on which the notice is given.

### 31B Secretary may require information relating to approvals and authorities under section 19

**Approval under subsection 19(1)**

(1) The Secretary may give to a person who is granted an approval under subsection 19(1) in relation to specified therapeutic goods a written notice requiring the person to give to the Secretary specified information or documents relating to one or more of the following:

(a) the supply of the goods;
(b) the handling of the goods;
(c) the monitoring of the supply of the goods;
(d) the results of the supply of the goods;
(e) any other matter prescribed by the regulations for the purposes of this paragraph in relation to goods of that kind.

**Approval under subsection 19(1)—use by another person**

(2) The Secretary may give to a person using specified therapeutic goods that are the subject of an approval granted to someone else under paragraph 19(1)(b) a written notice requiring the person to give to the Secretary specified information or documents relating to either of both of the following:

(a) the use of the goods;
(b) any other matter prescribed by the regulations for the purposes of this paragraph in relation to goods of that kind.

**Authority under subsection 19(5)**

(3) The Secretary may give to a person who is granted an authority under subsection 19(5) in relation to specified therapeutic goods, or
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a specified class of therapeutic goods, a written notice requiring the person to give to the Secretary specified information or documents relating to one or more of the following:

(a) the supply of the goods;
(b) the handling of the goods;
(c) the monitoring of the supply of the goods;
(d) the results of the supply of the goods;
(e) any other matter prescribed by the regulations for the purposes of this paragraph in relation to goods of that kind.

Compliance period

(4) A notice under subsection (1), (2) or (3) must specify a reasonable period within which the person to whom the notice is given must comply with it. The period must be at least 14 days starting on the day on which the notice is given.

31C Requirements in relation to information or documents sought under section 31A, 31AA or 31B

When information or documents must be given etc.

(1) A person to whom a notice is given under section 31A, 31AA or 31B must give the Secretary:

(a) the information or documents specified in the notice within the period specified in the notice; and
(b) the information specified in the notice in the form (if any) specified in the notice.

Way in which information given

(2) A notice mentioned in subsection (1) may require information to be given in accordance with specified software requirements:

(a) on a specified kind of data processing device; or
(b) by way of a specified kind of electronic transmission.
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Offence

(3) A person mentioned in subsection (1) is guilty of an offence if the person fails to comply with that subsection.

Note: The privilege against self-incrimination is not a reasonable excuse for the purposes of subsection (3). However, the information given, and the fact that a document was given under this section (and other information, documents or things obtained because of giving the information or document) generally cannot be used in a prosecution (see section 31F).

Penalty

(4) An offence against subsection (3) is punishable on conviction by a fine of not more than 60 penalty units.

31D False or misleading information

(1) A person to whom a notice is given under section 31A, 31AA or 31B is guilty of an offence if:

(a) the person gives information to the Secretary in compliance or purported compliance with subsection 31C(1); and

(b) the person does so knowing that the information:

(i) is false or misleading; or

(ii) omits any matter or thing without which the information is misleading.

Maximum penalty: Imprisonment for 12 months.

(2) Subsection (1) does not apply as a result of subparagraph (1)(b)(i) if the information is not false or misleading in a material particular.

Note: A defendant bears an evidential burden in relation to the matter in subsection (2).

(3) Subsection (1) does not apply as a result of subparagraph (1)(b)(ii) if the information did not omit any matter or thing without which the information is misleading in a material particular.

Note: A defendant bears an evidential burden in relation to the matter in subsection (3).
31E False or misleading documents

(1) A person is guilty of an offence if:
   (a) the person produces a document to the Secretary; and
   (b) the person does so knowing that the document is false or misleading; and
   (c) the document is produced in compliance or purported compliance with subsection 31C(1).

   Maximum penalty: Imprisonment for 12 months.

(2) Subsection (1) does not apply if the document is not false or misleading in a material particular.

   Note: A defendant bears an evidential burden in relation to the matter in subsection (2).

(3) Subsection (1) does not apply to a person who produces a document if the document is accompanied by a written statement signed by the person or, in the case of a body corporate, by a competent officer of the body corporate:
   (a) stating that the document is, to the knowledge of the first-mentioned person, false or misleading in a material particular; and
   (b) setting out, or referring to, the material particular in which the document is, to the knowledge of the first-mentioned person, false or misleading.

   Note: A defendant bears an evidential burden in relation to the matter in subsection (3).

31F Self-incrimination

(1) A person is not excused from giving information or a document under section 31C on the ground that the giving of the information or document would tend to incriminate the person or expose the person to a penalty.

(2) However, in the case of an individual:
   (a) the information given; or
   (b) the giving of the document; or
   (c) any information, document or thing obtained as a direct or indirect consequence of giving the information or document;
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is not admissible in evidence in criminal proceedings against the individual (except proceedings under, or arising out of, section 31D or 31E).
Part 3-3—Manufacturing of therapeutic goods

33A Application of this Part to medical devices

This Part does not apply to a medical device unless Part 3-2 applies to the device.

Note: Section 15A sets out when Part 3-2 applies to a medical device.

34 Exempt goods and exempt persons

(1) The regulations may exempt therapeutic goods or a class of therapeutic goods identified in the regulations from the operation of this Part.

(2) The regulations may exempt a person identified in the regulations from the operation of this Part in relation to the manufacture or a step in the manufacture of therapeutic goods or a class of therapeutic goods identified in the regulations.

(3) Where the regulations revoke an exemption, the revocation takes effect on the day, not being earlier than 28 days after the day on which the regulations are made, as is specified in the regulations.

35 Offences relating to manufacturing and licences

(1) A person must not, at premises in Australia, carry out a step in the manufacture of therapeutic goods (other than goods exempt under section 18A) for supply for use in humans unless:

(a) the goods are exempt goods or the person is an exempt person in relation to the manufacture of the goods; or

(b) the person is the holder of a licence that is in force that authorises the carrying out of that step in relation to the goods at those premises.

Maximum penalty: Imprisonment for 12 months or 1,000 penalty units, or both.

(2) A person is guilty of an offence if:

(a) the person holds a licence; and
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(b) the person engages in conduct; and
(c) the conduct breaches a condition of the licence.

Maximum penalty: Imprisonment for 12 months or 1,000 penalty units, or both.

(3) In subsection (2):

*engage in conduct* means:
(a) do an act; or
(b) omit to perform an act.

(4) A person commits an offence if:
(a) the person, at premises in Australia, carries out a step in the manufacture of therapeutic goods for supply for use in humans; and
(b) the goods are exempt under section 18A; and
(c) the person is not the holder of a licence that:
   (i) is in force; and
   (ii) authorises the carrying out of that step in relation to the goods at those premises.

Maximum penalty: Imprisonment for 12 months or 1,000 penalty units, or both.

(5) Strict liability applies to paragraph (4)(b).

Note: For strict liability, see section 6.1 of the *Criminal Code*.

36 Manufacturing principles

(1) The Minister may, from time to time, determine written principles to be observed in the manufacture of therapeutic goods for use in humans.

(2) The manufacturing principles may relate to:
(a) the standards to be maintained, and the equipment to be used, at premises used for the manufacturing of therapeutic goods for use in humans; or
(b) procedures for quality assurance and quality control to be employed in the manufacturing of therapeutic goods for use in humans; or
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(c) the qualifications and experience required of persons employed in the manufacture of therapeutic goods for use in humans; or

(d) the manufacturing practices to be employed in the manufacturing of therapeutic goods for use in humans; or

(e) other matters relevant to the quality, safety and efficacy of therapeutic goods for use in humans that are manufactured in Australia; and may include codes of good manufacturing practice.

(3) The Minister may, before taking action under subsection (1) in relation to the manufacturing principles, obtain advice from a committee established by the regulations on the action that should be taken under that subsection as to the principles to be observed in the manufacture of therapeutic goods for use in humans.

(4) Manufacturing principles are disallowable instruments for the purposes of section 46A of the Acts Interpretation Act 1901.

37 Application for licence

(1) An application for a licence must:

(a) be made in writing in accordance with a form approved by the Secretary; and

(b) identify the therapeutic goods or classes of therapeutic goods that the applicant proposes to manufacture; and

(c) identify the manufacturing premises that will be used in the manufacture of those goods; and

(d) identify the steps in the manufacture of those goods that the applicant proposes to carry out under the licence; and

(da) if the applicant proposes to carry out steps in the manufacture of blood or blood components under the licence—contain information relating to those steps set out in regulations made for the purposes of this paragraph; and

(e) state the names, qualifications and experience of the persons who are to have control of the production of the goods and of the quality control measures that are to be employed; and

(f) be delivered to an office of the Department specified in the form; and

(g) be accompanied by the prescribed application fee.
Section 38

(2) The Secretary may, by notice in writing given to an applicant for a licence, require the applicant:

(a) to give to the Secretary, within such reasonable time as is specified in the notice, such further information concerning the application as is specified in the notice; or

(b) to allow an authorised person, at any reasonable time specified in the notice, to inspect the premises, equipment, processes and facilities that will be used in the manufacture of the goods, or other goods on those premises.

38 Grant of licence

(1) Where:

(a) a person has made an application to carry out steps in the manufacture of therapeutic goods at particular manufacturing premises; and

(b) the prescribed application fee has been paid; and

(c) any applicable prescribed inspection fees have been paid; and

(d) the applicant has complied with any requirements made by the Secretary under subsection 37(2) in relation to the application;

the Secretary must grant the applicant a licence to carry out those steps at those premises unless the Secretary is satisfied that:

(e) the applicant will be unable to comply with the manufacturing principles; or

(f) the premises are not satisfactory for the manufacture of the goods; or

(g) the applicant is not a fit and proper person to hold a licence; or

(h) a person who is participating in, or is likely to participate in, managing the applicant’s affairs is not a fit and proper person to participate in the management of the affairs of a holder of a licence; or

(i) a person who has, or is likely to have, effective control over the applicant is not a fit and proper person to have effective control over a holder of a licence.

(1A) Without limiting the matters to which the Secretary may have regard in considering whether the applicant or person is not a fit

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and proper person for the purposes of paragraph (1)(g), (h) or (i), the Secretary must have regard to:

(a) any suspension or revocation of a manufacturing licence granted to:
   (i) the applicant or person; or
   (ii) another person who controls the applicant or person (whether directly, or indirectly through one or more interposed entities); or
   (iii) another person whom the applicant or person controlled (whether directly, or indirectly through one or more interposed entities) at the time of the suspension or revocation; or

(b) any conviction, for an offence against a law of the Commonwealth or a law of a State or Territory, against:
   (i) the applicant or person; or
   (ii) another person who controls the applicant or person (whether directly, or indirectly through one or more interposed entities); or
   (iii) another person whom the applicant or person controlled (whether directly, or indirectly through one or more interposed entities) at the time the offence was committed or the time of the conviction; or

(c) any failure to comply with a condition of a manufacturing licence by:
   (i) the applicant or person; or
   (ii) another person who controls the applicant or person (whether directly, or indirectly through one or more interposed entities); or
   (iii) another person whom the applicant or person controlled (whether directly, or indirectly through one or more interposed entities) at the time of the failure.

(1B) In subsection (1A):

*manufacturing licence* means:

(a) a licence granted under this Part; or

(b) a licence, granted under a law of a State or Territory relating to therapeutic goods, relating to manufacturing therapeutic goods.
Chapter 3  Medicines and other therapeutic goods that are not medical devices
Part 3-3  Manufacturing of therapeutic goods

Section 39

(2) Notwithstanding paragraphs (1)(g), (h) and (i), the Secretary may grant a licence to an applicant who, apart from this subsection, could not be granted a licence because of one or more of those paragraphs if, in the opinion of the Secretary, special circumstances make it appropriate to do so.

(3) Where the Secretary grants or refuses to grant a licence to an applicant, the Secretary must:
   (a) give the applicant written notice of the decision; and
   (b) in the case of a refusal—include in the notice the reasons for the refusal.

(4) Where the Secretary grants a licence, the Secretary must cause particulars of the decision to be published in the Gazette as soon as is practicable after the decision is made.

39  Term of licence

(1) A licence commences on the day specified in the licence and remains in force until it is revoked or suspended.

(2) If:
   (a) the licence covers therapeutic goods that are exempt under section 18A; and
   (b) those goods cease to be exempt under that section before the licence is revoked;

   the licence ceases to be in force in relation to those goods when those goods cease to be exempt under that section.

   Note: An exemption under section 18A may cease to have effect only in relation to some of the goods covered by the exemption, see subsection 18A(5).

40  Conditions of licences

(1) A licence may be granted subject to such conditions relating to the manufacture of the goods as the Secretary thinks appropriate.

(2) The Secretary may, by notice in writing given to the holder of a licence, impose new conditions on the licence or vary or remove existing conditions.

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(3) The imposition or variation of a condition under subsection (2) takes effect:

(a) if the notice states that the action is necessary to prevent imminent risk of death, serious illness or serious injury—on the day on which the notice is given to the person; or

(b) in any other case—on the day specified for the purpose in the notice, being a day not earlier than 28 days after the notice is given to the person.

(4) In addition to any conditions imposed under subsection (1) or (2), each licence is, except as otherwise specified in the licence, subject to the conditions that the holder of the licence will:

(a) ensure that:

(i) the goods conform to any standard applicable to the goods; and

(ii) the holder of the licence observes the manufacturing principles in carrying out any steps in the manufacture of the goods under the licence; and

(aa) if:

(i) the holder of the licence carries out, or proposes to carry out, steps in the manufacture of blood or blood components under the licence; and

(ii) regulations made for the purposes of this paragraph set out particular information relating to those steps;

comply with a request by the Secretary to provide such information, in accordance with those regulations; and

(ab) as soon as the holder of the licence becomes aware of information of a kind mentioned in subsection (5), give the information to the Secretary in writing; and

(b) allow an authorised person:

(i) to enter, at any reasonable time, the manufacturing premises to which the licence relates; and

(ii) while on those premises, to inspect those premises, any therapeutic goods manufactured at those premises and processes relating to that manufacture, and to take samples of goods of that kind and to take photographs of those premises, goods or processes; and

(c) where an authorised person enters premises as mentioned in subparagraph (b)(i), require the holder or his or her
employees at those premises to answer questions relating to procedures carried out at the premises; and
(d) if requested to do so by an authorised person:
   (i) produce to the person such documents relating to the manufacture of therapeutic goods manufactured at those premises as the person requires and allow the person to copy the documents; or
   (ii) produce to the person for examination any batch samples kept by the holder; and
(e) comply with such other conditions (if any) as are specified in the regulations for the purposes of this section.

(5) The information with which paragraph (4)(ab) is concerned is information of the following kinds:

(a) information that indicates that the use of the goods in accordance with the recommendations for their use may have an unintended harmful effect;
(b) information that indicates that the goods, when used in accordance with the recommendations for their use, may not be as effective as was suggested by:
   (i) the application for registration or listing of the goods; or
   (ii) information already furnished by the holder of the licence under this Act; or
   (iii) if the holder of the licence is not the sponsor of the goods—information already furnished by the sponsor of the goods under this Act;
(c) information that indicates that the quality, safety or efficacy of the goods is unacceptable.

41 Revocation and suspension of licences

(1) Subject to subsection (2), the Secretary may, by notice in writing given to the holder of a licence, revoke the licence, or suspend the licence for a period specified in the notice, if:
   (a) the holder has been convicted of an offence against this Act; or
   (aa) the holder controls another person (whether directly, or indirectly through one or more interposed entities) that has
been convicted of an offence against this Act or a law of a State or Territory relating to therapeutic goods; or

(ab) the holder controlled another person (whether directly, or indirectly through one or more interposed entities) when the other person committed an offence against this Act or a law of a State or Territory relating to therapeutic goods, and the other person has been convicted of that offence; or

(ac) the holder is controlled by another person (whether directly, or indirectly through one or more interposed entities) and that other person has been convicted of an offence against this Act or a law of a State or Territory relating to therapeutic goods; or

(b) the holder has breached a condition of the licence; or

(c) the holder is controlled by another person (whether directly, or indirectly through one or more interposed entities) and that other person has breached a condition of a licence; or

(ca) the holder controls another person (whether directly, or indirectly through one or more interposed entities) and that other person has, while controlled by the holder, breached a condition of a licence; or

(cb) the holder is not a fit and proper person to hold a licence; or

(cc) a person who is participating in managing the holder’s affairs is not a fit and proper person to participate in the management of the affairs of a holder of a licence; or

(cd) a person who has effective control over the holder is not a fit and proper person to have effective control over a holder of a licence; or

(d) the holder requests in writing that the licence be revoked or suspended, as the case may be; or

(e) the holder ceases to carry on the business of manufacturing the goods to which the licence relates; or

(f) the annual licensing charge, or any applicable prescribed inspection fees, have not been paid within 28 days after they become payable; or

(g) the goods are exempt under section 18A and the holder has breached a condition of the exemption in relation to those goods.
Section 41A

(1A) Without limiting the matters to which the Secretary may have regard in considering whether the holder or another person is not a fit and proper person for the purposes of paragraph (1)(cb), (cc) or (cd), the Secretary must have regard to the matters set out in paragraphs 38(1A)(a), (b) and (c).

(2) Where the Secretary proposes to revoke a licence or suspend a licence otherwise than at the request of the holder of the licence, the Secretary must, unless the Secretary considers that failure to revoke or suspend the licence immediately would create an imminent risk of death, serious illness or serious injury:

(a) by notice in writing given to the holder, inform the holder of the action that the Secretary proposes to take and of the reasons for that proposed action; and

(b) except where the proposed action is to be taken as a result of a failure to pay the annual licensing charge or an applicable prescribed inspection fee—give the holder an opportunity to make, within such reasonable time as is specified in the notice, submissions to the Secretary in relation to the proposed action.

(3) Where the holder makes submissions in accordance with paragraph (2)(b), the Secretary is not to make a decision relating to the revocation or suspension of the licence before taking into account the submissions.

(4) A licence may be revoked notwithstanding that the licence is suspended.

(5) Where a licence is suspended, the Secretary may, by notice in writing given to the holder of the licence, revoke the suspension.

(6) Where the Secretary revokes or suspends a licence, the Secretary must cause particulars of the decision to be published in the Gazette as soon as is practicable after the decision is made.

41A Publication of list of manufacturers etc.

The Secretary may, from time to time and in such manner as the Secretary determines, publish a list of the persons who are licensed under this Part, the classes of goods to which the licences relate, the steps of manufacture that the licences authorise and the
addresses of the manufacturing premises to which the licences relate.
Chapter 4—Medical devices

Note: For 5 years following the commencement of this Chapter, this Chapter will not apply, and Chapter 3 (Medicines and other therapeutic goods that are not medical devices) will still apply, to medical devices that are registered or listed goods.

Part 4-1—Introduction

Division 1—Overview of this Chapter

41B General

The purpose of this Chapter is to ensure the safety and satisfactory performance of medical devices. It does this by:

(a) setting out particular requirements for medical devices; and
(b) establishing administrative processes principally aimed at ensuring those requirements are met; and
(c) providing for enforcement through a series of offences.

41BA Requirements for medical devices (Parts 4-2 and 4-3)

The requirements for medical devices are:

(a) essential principles (that are about the safety and performance characteristics of medical devices); and
(b) conformity assessment procedures (that are mainly about the application of quality management systems).

Note: Medical device standards may be made under Division 2 of Part 4-2, and conformity assessment standards may be made under Division 2 of Part 4-3, but they are not requirements.

41BB Administrative processes (Parts 4-4 to 4-10)

The administrative processes under this Chapter are:

(a) issuing conformity assessment certificates for some manufacturers of medical devices; and
(b) including medical devices in the Register; and
(c) suspending or cancelling entries of medical devices from the Register;
(d) exempting medical devices from the requirement to be included in the Register; and
(e) obtaining information about medical devices; and
(f) requiring public notification of problems with medical devices, and recovery of such devices.

Note: Part 4-10 provides for assessment fees to be payable in some circumstances.

41BC Enforcement (Part 4-11)

Part 4-11 contains offences that are aimed at ensuring that:
(a) the requirements for medical devices are complied with; and
(b) the administrative processes under this Chapter (particularly the inclusion of medical devices in the Register) are followed.

Note: There are some offences in Parts 4-4 to 4-9. They generally relate to matters ancillary to administrative processes in those Parts (e.g. false or misleading statements in applications).
Division 2—Interpretation

41BD What is a medical device

(1) A medical device is:
   (a) any instrument, apparatus, appliance, material or other article
       (whether used alone or in combination, and including the
        software necessary for its proper application) intended, by
        the person under whose name it is or is to be supplied, to be
        used for human beings for the purpose of one or more of the
        following:
           (i) diagnosis, prevention, monitoring, treatment or
               alleviation of disease;
           (ii) diagnosis, monitoring, treatment, alleviation of or
                compensation for an injury or handicap;
           (iii) investigation, replacement or modification of the
                 anatomy or of a physiological process;
           (iv) control of conception;
       and that does not achieve its principal intended action in or
       on the human body by pharmacological, immunological or
       metabolic means, but that may be assisted in its function by
       such means; or
   (b) an accessory to such an instrument, apparatus, appliance,
       material or other article.

Note: Declarations under subsection (3) exclude articles from the scope of
this definition. Declarations under section 7 can also have this effect:
see subsection 7(4).

(2) For the purposes of paragraph (1)(a), the purpose for which an
article is to be used is to be ascertained from the information
supplied, by the person under whose name the article is or is to be
supplied, on or in any one or more of the following:
   (a) the labelling on the article;
   (b) the instructions for using the article;
   (c) any advertising material relating to the article.

(3) The Secretary may, by order published in the Gazette, declare that
a particular instrument, apparatus, appliance, material or other
article, or that a particular class of instruments, apparatus,
appliances, materials or other articles, are not, for the purposes of this Act, medical devices.

Note: A declaration under this section does not stop articles from being therapeutic goods.

(4) A declaration under this section takes effect on the day on which the declaration is published in the Gazette or on such later day as is specified in the order.

41BE Kinds of medical devices

General

(1) For the purposes of this Chapter, a medical device is taken to be of the same kind as another medical device if they:
   (a) have the same sponsor; and
   (b) have the same manufacturer; and
   (c) have the same device nomenclature system code (see subsection (3)); and
   (d) have the same medical device classification; and
   (e) are the same in relation to such other characteristics as the regulations prescribe, either generally or in relation to medical devices of the kind in question.

Unique medical devices

(2) If a medical device is not of the same kind as any other medical device:
   (a) this Chapter applies in relation to the device as if it were a kind of medical device; and
   (b) references in this Chapter to delivering a reasonable number of samples of the kind of device are taken to be references to delivering the device.

Device nomenclature codes

(3) The regulations may specify device nomenclature codes for medical devices.
Section 41BF

41BF System or procedure packs

(1) A package and therapeutic goods in the package are a system or procedure pack if:
   (a) the package and the therapeutic goods are for use as a unit, either in combination as a system or in a medical or surgical procedure; and
   (b) the package contains at least one medical device; and
   (c) the package and the therapeutic goods do not constitute a composite pack.

(2) To avoid doubt, a system or procedure pack is a medical device.

41BG Manufacturers of medical devices

(1) The manufacturer of a medical device is the person who is responsible for the design, production, packaging and labelling of the device before it is supplied under the person’s name, whether or not it is the person, or another person acting on the person’s behalf, who carries out those operations.

(2) If subsection (1) does not apply to a medical device, the manufacturer of the device is the person who, with a view to supplying the device under the person’s name, does one or more of the following using ready-made products:
   (a) assembles the device;
   (b) packages the device;
   (c) processes the device;
   (d) fully refurbishes the device;
   (e) labels the device;
   (f) assigns to the device its purpose by means of information supplied, by the person, on or in any one or more of the following:
      (i) the labelling on the device;
      (ii) the instructions for using the device;
      (iii) any advertising material relating to the device.

(3) However, a person is not the manufacturer of a medical device if:
   (a) the person assembles or adapts the device for an individual patient; and
(b) the device has already been supplied by another person; and
(c) the assembly or adaptation does not change the purpose intended for the device by means of information supplied by that other person, on or in any one or more of the following:
   (i) the labelling on the device;
   (ii) the instructions for using the device;
   (iii) any advertising material relating to the device.

41BH Meaning of compliance with essential principles

(1) A medical device complies, for the purposes of this Chapter (including Part 4-11), with the essential principles if and only if it does not contravene any of the essential principles.

(2) However, a medical device is also taken, for the purposes of this Chapter (other than Part 4-11), to comply with the essential principles if:
   (a) the medical device complies with one or more medical device standards that apply to it; and
   (b) the medical device contravenes the essential principles only in respect of a part or parts of the essential principles to which that medical device standard, or one or more of those medical device standards, relate.

(3) For the purposes of this section, a medical device standard relates to a part or parts of the essential principles only if the standard specifies that part or parts.

41BI Meaning of non-application of conformity assessment procedures

(1) A conformity assessment procedure is taken, for the purposes of this Chapter, not to have been applied to a medical device if:
   (a) there has been a contravention of the conformity assessment procedures; and
   (b) the contravention relates, wholly or partly, to that device or its manufacture.

(2) However, for the purposes of this Chapter (other than Part 4-11), subsection (1) does not apply if:
Section 41BI

(a) the quality management system applied in the manufacture of the medical device complies with one or more conformity assessment standards that apply to it; and

(b) the contravention is only in respect of a part or parts of the conformity assessment procedures to which that conformity assessment standard, or one or more of those conformity assessment standards, relate.

(3) For the purposes of this section, a conformity assessment standard relates to a part or parts of the conformity assessment procedures only if the standard specifies that part or parts.

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Division 3—Application provisions

41BJ Application of this Chapter to medical devices covered by Part 3-2

(1) This Chapter does not apply to a medical device if section 15A applies to the device, except for purposes connected with:
   (a) applications for including the medical device in the Register under this Chapter; or
   (b) including the medical device in the Register under this Chapter.

Note: Section 15A sets out the circumstances in which Part 3-2 applies or continues to apply to medical devices.

(2) However, if an exemption under section 34 applied to a medical device, or the manufacturer of the device, immediately before the commencement of this Chapter:
   (a) Parts 4-3 and 4-4, and Division 2 of Part 4-11, apply in relation to the device after the end of the period of 2 years after that commencement; and
   (b) Parts 4-8, 4-9 and 4-10, and Divisions 3 and 4 of Part 4-11, apply in relation to the device, to the extent that they relate to any of the provisions referred to in paragraph (a), after the end of that period.

41BK Application of the Criminal Code

Chapter 2 of the Criminal Code applies to all offences against this Chapter.

Note: Chapter 2 of the Criminal Code sets out the general principles of criminal responsibility.
Part 4-2—Essential principles and medical device standards

41C What this Part is about

The essential principles set out the requirements relating to the safety and performance characteristics of medical devices. Compliance with applicable medical device standards is not required, but it is one way to establish compliance with essential principles.

Note: Dealing in medical devices that do not comply with the essential principles may be an offence: see Division 1 of Part 4-11.

Division 1—Essential principles

41CA Essential principles

(1) The regulations may set out requirements for medical devices.

(2) These requirements are to be known as the essential principles.
Division 2—Medical device standards

41CB Medical device standards

(1) The Minister may, by order published in the Gazette, determine that:

(a) matters specified in the order constitute a medical device standard for kinds of medical devices identified in the order; and

(b) medical devices of those kinds that comply with the standard are to be treated as complying with those parts of the essential principles specified in the standard.

(2) A medical device standard takes effect on the day on which the order establishing the medical device standard is published in the Gazette or on such later day as is specified in the order.

(3) A medical device standard is a disallowable instrument for the purposes of section 46A of the Acts Interpretation Act 1901.

41CC Content of medical device standards

(1) Without limiting the scope of section 41CB, an order establishing a medical device standard for kinds of medical devices may be specified by reference to:

(a) the safety or performance characteristics of the devices; or

(b) a monograph in the British Pharmacopoeia, the European Pharmacopoeia or the United States Pharmacopoeia; or

(c) a monograph in a publication approved by the Minister for the purposes of this subsection; or

(d) such a monograph as modified in a manner specified in the order; or

(e) a standard published by a standards organisation; or

(f) such other matters as the Minister thinks fit.

(2) For the purposes of paragraph (1)(e), these are standards organisations:

(a) Standards Australia International Limited;

(b) the International Organisation for Standardization;
(c) the International Electrotechnical Commission;
(d) the European Committee for Standardization;
(e) the European Committee for Electrotechnical Standardization;
(f) any other organisation declared by the Minister by notice published in the Gazette.

41CD  Inconsistencies between medical device standards

(1) A medical device standard that:
   (a) applies to a kind of medical device; and
   (b) is inconsistent with another medical device standard that applies only to some of the devices of that kind;
   is, to the extent of the inconsistency, of no effect in relation to the devices referred to in paragraph (b).

(2) A medical device standard that applies to a kind of medical device that consists of a combination of component parts takes precedence over any medical device standard that applies to the component parts.
Part 4-3—Conformity assessment procedures

41D What this Part is about

The conformity assessment procedures set out the requirements relating to the application of quality management systems for medical devices, and other requirements imposed on manufacturers.

Compliance with applicable conformity assessment standards is not required, but it is one way to establish that one or more parts of the conformity assessment procedures have been applied to medical devices.

Note 1: Dealing in medical devices that have not had the conformity assessment procedures applied may be an offence: see Division 2 of Part 4-11.

Note 2: See section 41BI on applying the conformity assessment procedures.

Division 1—Conformity assessment procedures

41DA Conformity assessment procedures

(1) The regulations may set out requirements relating to the obligations of manufacturers of medical devices.

(2) These requirements are to be known as the conformity assessment procedures.

(3) The conformity assessment procedures, or any part of the conformity assessment procedures, may:
   (a) be limited in their application to one or more medical device classifications; or
   (b) apply differently to different medical device classifications, different kinds of medical devices or different manufacturers.

(4) Without limiting subsection (1), the regulations may relate to all or any of the following:
(a) application of quality management systems for the manufacture of medical devices;
(b) certification of compliance with the essential principles, or the quality management systems for the manufacture of medical devices;
(c) notification of, and assessment of, changes to a manufacturer’s product range, product design or quality management systems;
(d) declarations to be made by manufacturers of medical devices that conformity assessment procedures have been applied to the devices;
(e) marks to be affixed to medical devices indicating the application of the conformity assessment procedures to the devices;
(f) monitoring and inspecting the design of medical devices or the manufacturing processes for medical devices;
(g) monitoring the performance of medical devices;
(h) corrective action required in relation to the design, manufacture, packaging, labelling and supply of medical devices;
(i) keeping records of the manufacture of medical devices, the design of medical devices or the manufacturing processes for medical devices.

41DB Medical device classifications

The regulations may specify:
(a) classifications, to be known as medical device classifications, applying to medical devices or kinds of medical devices; and
(b) matters in relation to the classification of medical devices or kinds of medical devices.
Division 2—Conformity assessment standards

41DC Conformity assessment standards

(1) The Minister may, by order published in the Gazette, determine that:
   (a) matters specified in the order constitute a conformity assessment standard for quality management systems identified in the order; and
   (b) a quality management system that complies with the standard is to be treated as having had applied to it those parts of the conformity assessment procedures specified in the standard.

(2) A conformity assessment standard may be limited to particular kinds of medical devices.

(3) A conformity assessment standard takes effect on the day on which the order establishing the conformity assessment standard is published in the Gazette or on such later day as is specified in the order.

(4) A conformity assessment standard is a disallowable instrument for the purposes of section 46A of the Acts Interpretation Act 1901.

41DD Content of conformity assessment standards

(1) Without limiting the scope of section 41DC, an order establishing a conformity assessment standard for a kind of medical device may be specified by reference to:
   (a) procedures to be carried out under the quality management systems for the design, manufacture and final inspection of the devices; or
   (b) a standard published by a standards organisation; or
   (c) such other matters as the Minister thinks fit.

(2) For the purposes of paragraph (1)(b), these are standards organisations:
   (a) Standards Australia International Limited;
   (b) the International Organisation for Standardization;
Section 41DE

(c) the European Committee for Standardization;
(d) any other organisation declared by the Minister by notice published in the Gazette.

41DE  Inconsistencies between conformity assessment standards

A conformity assessment standard that:
(a) identifies quality management systems to which it applies; and
(b) is inconsistent with another conformity assessment standard that applies only to particular kinds of medical devices;
is, to the extent of the inconsistency, of no effect in relation to the devices referred to in paragraph (b).
Part 4-4—Conformity assessment certificates

41E What this Part is about

The Secretary can issue a conformity assessment certificate (which may be limited to some medical devices) in respect of a manufacturer of medical devices, signifying one or more of these:

(a) that relevant quality management systems have been applied to the device;
(b) the essential principles for the device have been complied with;
(c) other certification requirements of the conformity assessment procedures have been met.

Note: A conformity assessment certificate may be required before a valid application can be made for including a kind of medical device in the Register under this Chapter: see subsection 41FC(2).

Division 1—Issuing conformity assessment certificates

41EA When conformity assessment certificates are required

The regulations may prescribe:

(a) kinds of manufacturers in respect of whom a conformity assessment certificate must be issued before valid applications can be made for kinds of medical devices, manufactured by those manufacturers, to be included in the Register; or
(b) kinds of medical devices in respect of which a conformity assessment certificate must be issued before valid applications can be made for those kinds of medical devices to be included in the Register.

Note: The regulations may prescribe different levels of fees for different kinds of manufacturers and medical devices: see subsection 41LA(2).
Chapter 4  Medical devices
Part 4-4  Conformity assessment certificates
Division 1  Issuing conformity assessment certificates

Section 41EB

41EB  Applications

(1) An application for a conformity assessment certificate must:
   (a) be made in accordance with a form approved, in writing, by
       the Secretary or in such other manner as is approved, in
       writing, by the Secretary; and
   (b) be delivered to an office of the Department specified by the
       Secretary.

   Note: A conformity assessment fee is payable under section 41LA for
   consideration of the application.

(2) An application is not effective if:
   (a) the prescribed application fee has not been paid; or
   (b) the application contains information that is false or
       misleading in a material particular.

   Note: A person might also be guilty of an offence if the person makes a
   statement in an application that is false or misleading in a material
   particular: see section 41EI.

(3) An approval of a form may require or permit an application or
    information to be given in accordance with specified software
    requirements:
    (a) on a specified kind of data processing device; or
    (b) by way of a specified kind of electronic transmission.

(4) The Secretary may, by written notice given to an applicant for a
    conformity assessment certificate, require the applicant to allow an
    authorised person, at any reasonable time specified in the notice, to
    inspect:
    (a) the premises (including premises outside Australia) and
        equipment, processes and facilities that are being or will be
        used to manufacture medical devices of the kind in question;
        and
    (b) any other kinds of medical devices on those premises.

41EC  Considering applications

(1) If the application is made in accordance with section 41EB, the
    Secretary must decide whether to issue the conformity assessment
    certificate.

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(2) In deciding whether to issue the certificate, the Secretary must consider some or all aspects of whether the conformity assessment procedures relating to one or more of the following have been applied to the medical device:

(a) the application of quality management systems for the manufacture of medical devices;
(b) the certification of compliance with the essential principles;
(c) any other requirement of the conformity assessment procedures specified in regulations made for the purposes of this subsection.

(3) In deciding whether to issue the certificate, the Secretary must also consider:

(a) whether the applicant is a fit and proper person to hold a conformity assessment certificate; and
(b) whether the persons who participate in, or who are likely to participate in, managing the applicant’s affairs are fit and proper persons to participate in managing the affairs of a manufacturer in respect of whom a conformity assessment certificate is issued; and
(c) whether the persons who have, or are likely to have, effective control over the applicant are fit and proper persons to have effective control over a manufacturer in respect of whom a conformity assessment certificate is issued.

(4) Without limiting the matters to which the Secretary may have regard in considering whether the applicant or person is a fit and proper person for the purposes of paragraph (3)(a), (b) or (c), the Secretary must have regard to:

(a) any suspension or revocation of a conformity assessment certificate issued to:

(i) the applicant or person; or
(ii) another person who controls the applicant or person (whether directly, or indirectly through one or more interposed entities); or
(iii) another person whom the applicant or person controlled (whether directly, or indirectly through one or more interposed entities) at the time of the suspension or revocation; or
(b) any conviction, for an offence against a law of the Commonwealth or a law of a State or Territory, against:
   (i) the applicant or person; or
   (ii) another person who controls the applicant or person (whether directly, or indirectly through one or more interposed entities); or
   (iii) another person whom the applicant or person controlled (whether directly, or indirectly through one or more interposed entities) at the time the offence was committed or the time of the conviction; or
(c) any failure to comply with a condition of a conformity assessment certificate by:
   (i) the applicant or person; or
   (ii) another person who controls the applicant or person (whether directly, or indirectly through one or more interposed entities); or
   (iii) another person whom the applicant or person controlled (whether directly, or indirectly through one or more interposed entities) at the time of the failure.

41ED Time for making decisions on applications

(1) If the application relates to the issuing of a conformity assessment certificate in relation to which a period has been prescribed under paragraph 63(2)(dc), a decision on the application must be made within that period, unless the application lapses under section 41EG.

(2) A failure to make a decision on the application within that period does not render the Commonwealth, the Secretary or a delegate of the Secretary liable to a person in respect of loss, damage or injury of any kind caused by, or arising out of, the failure.

41EE Procedure following making a decision whether to issue certificate

(1) After making a decision whether to issue a conformity assessment certificate, the Secretary must:
   (a) notify the applicant in writing of his or her decision within 20 working days; and

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(b) if the decision is not to issue the certificate—state in the notice the reasons for the decision; and
(c) if the decision is to issue the certificate and all assessment fees that are due and payable for the application have been paid:
   (i) issue the certificate to the manufacturer in relation to whom the application was made; and
   (ii) give the applicant a copy of the certificate (if the applicant is not the manufacturer).

(2) A conformity assessment certificate must specify whether it covers:
   (a) all medical devices manufactured by the manufacturer; or
   (b) only specified medical devices manufactured by the manufacturer.

**41EF Duration of certificate**

(1) The conformity assessment certificate commences on the day specified for the purpose in the certificate.

(2) A conformity assessment certificate has effect at all times:
   (a) unless the certificate is suspended under Division 3; or
   (b) until the end of the period (if any) specified in the certificate; or
   (c) until the certificate is revoked under Division 4.

**41EG Lapsing of applications**

An application for a conformity assessment certificate lapses if:
   (a) the applicant does not deliver to the office to which the application was made such information (in a form approved in writing by the Secretary) as will allow the certificate to be issued; or
   (b) the applicant does not comply with a requirement by the Secretary to deliver to the office to which the application was made a reasonable number of samples of the kind of medical device to which the application relates; or
   (c) the applicant fails to comply with a notice under section 41JA to give to the Secretary information relating to a
kind of medical device to which the application relates within a further 10 working days from the day specified in the notice; or

(d) information given to the Secretary by, or on behalf of, the applicant in connection with the application, including information given for the purpose of a notice under section 41JA, is false or misleading in a material particular; or

(e) the applicant fails to allow an authorised person to carry out any inspection as required under subsection 41EB(4); or

(f) the applicant fails to pay an assessment fee for the application within the period, specified in the regulations, after being notified of the decision to issue a conformity assessment certificate under section 41EE.

41EH Treating applications as having been refused

(1) The applicant for an application for a conformity assessment certificate may give the Secretary written notice that the applicant wishes to treat the application as having been refused if:

(a) a period is prescribed under paragraph 63(2)(dc) for making a decision on the application; and

(b) at the end of the period, the applicant has not been notified of a decision whether to issue the certificate.

(2) The notice may be given at any time before the applicant is notified of the decision.

(3) If a notice has been given, this Act (except for subsection 60(5)) has effect as if:

(a) the Secretary had decided not to issue the certificate; and

(b) the Minister had made a decision under subsection 60(3) confirming the decision of the Secretary; and

(c) the Minister’s decision had been made on the day on which notice was given to the Secretary.

41EI False statements

A person is guilty of an offence if:

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Section 41EI

(a) the person makes a statement (whether orally, in a document or in any other way); and

(b) the person knows that the statement is false or misleading in a material particular; and

(c) the statement is in or in connection with an application for a conformity assessment certificate.

Maximum penalty: 60 penalty units.
Division 2—Conditions

Note: Breaching conditions of the conformity assessment certificate may lead to suspension or revocation of the certificate (see Divisions 3 and 4), and may be an offence (see subsection 41MN(2)).

41EJ Automatic conditions on conformity assessment certificates

Entry and inspection powers

(1) A conformity assessment certificate is subject to the conditions that the manufacturer in respect of whom the certificate is issued will:
   (a) allow an authorised person:
       (i) to enter, at any reasonable time, premises (including premises outside Australia) at which the person or any other person deals with medical devices of a kind covered by the certificate; and
       (ii) while on those premises, to inspect those premises and medical devices of that kind at those premises and to take samples of devices of that kind; and
       (iii) to carry out tests, or require tests to be carried out, on the premises on medical devices of a kind covered by the certificate; and
   (b) if requested to do so by an authorised person:
       (i) produce to the person such documents relating to devices of that kind, or to the manufacturer’s quality management system, as the person requires; and
       (ii) allow the person to copy the documents.

Review

(2) A conformity assessment certificate is subject to the condition that the manufacturer in respect of whom the certificate is issued will cooperate in any review by the Secretary of the certificate to determine whether the conformity assessment procedures relating to the following matters have been applied to the kinds of medical devices covered by the certificate:
   (a) the application of quality management systems for the manufacture of medical devices;
   (b) the certification of compliance with the essential principles;
(c) any other requirement of the conformity assessment procedures specified in the regulations made for the purposes of subsection 41EC(2).

**Notification of substantial changes**

(3) A conformity assessment certificate is subject to the condition that the person in respect of whom the certificate is issued will notify the Secretary, in writing, of any plan for substantial changes to:
   (a) quality management systems; or
   (b) the product range covered by those systems; or
   (c) the product design of kinds of medical devices; in respect of which the certificate is issued.

**Fees**

(4) A conformity assessment certificate is subject to the condition that the applicant for the certificate will pay a fee, prescribed in the regulations, for a review under subsection (2), when the fee becomes due and payable.

(5) The regulations may prescribe different levels of fees for different kinds of manufacturers and medical devices.

**Conditions do not limit other conditions**

(6) A condition imposed under this section is in addition to any conditions imposed under this Division.

**41EK Conditions imposed when conformity assessment certificates are issued**

If the Secretary issues a conformity assessment certificate in respect of a manufacturer, the Secretary may, in writing, impose conditions on the certificate in respect of:

(a) one or more kinds of medical devices covered by the certificate; or

(b) the manufacturer’s quality management system.
41EL  Conditions imposed after issuing a conformity assessment certificate

(1) The Secretary may, by written notice given to a manufacturer in respect of whom a conformity assessment certificate has been issued:
   (a) impose new conditions on the certificate in respect of:
       (i) one or more kinds of medical devices covered by the certificate; or
       (ii) the manufacturer’s quality management system; or
   (b) vary or remove existing conditions.

The power may be exercised at the request of the applicant for the certificate or on the Secretary’s own initiative.

(2) The imposition or variation of a condition under this section takes effect:
   (a) if the notice states that the action is necessary to prevent imminent risk of death, serious illness or serious injury—on the day on which the notice is given to the person; or
   (b) in any other case—on the day specified for the purpose in the notice, being a day not earlier than 20 working days after the notice is given to the person.
Division 3—Suspension of conformity assessment certificates

Note: Suspension of a conformity assessment certificate leads to suspension from the Register of the kinds of medical devices to which the certificate applied (see subsection 41GF(1)). Applications to include such devices in the Register are not effective (see paragraph 41FC(2)(c)).

41EM Suspension of conformity assessment certificates

(1) The Secretary may, by written notice given to the person in relation to whom a conformity assessment certificate is issued, suspend the certificate if the Secretary is satisfied that it is likely that there are grounds for revoking the certificate under section 41ET.

(2) The suspension may be limited to some medical devices of that kind, as specified in the notice.

(3) The notice must specify the period of the suspension. The period must not exceed 6 months.

Note: The period of the suspension may be extended under section 41EO.

41EN Notice of proposed suspension

(1) However, before suspending a conformity assessment certificate, the Secretary must:

(a) inform the person in writing that the Secretary proposes the suspension and set out the reasons for it; and

(b) give the person a reasonable opportunity to make submissions to the Secretary in relation to the proposed suspension.

(2) The Secretary is not to make a decision relating to the proposed suspension until the Secretary has had regard to any submissions the person makes under paragraph (1)(b).

(3) This section does not apply if the notice under section 41EM states that the suspension is necessary to prevent imminent risk of death, serious illness or serious injury.
Section 41EO

41EO Duration of suspension

(1) The suspension takes effect:
   (a) if the notice under section 41EM states that the suspension is necessary to prevent imminent risk of death, serious illness or serious injury—on the day on which the notice is given to the person; or
   (b) in any other case—on the day specified for the purpose in the notice, being a day not earlier than 20 working days after the notice is given to the person.

(2) The suspension has effect until:
   (a) the Secretary revokes it under section 41EP; or
   (b) the expiry of:
      (i) the period specified in the notice under section 41EM; or
      (ii) if the period is extended under subsection (3) of this section, the period as so extended.

Note: Unless a suspension of a conformity assessment certificate has been revoked, the certificate is automatically revoked: see section 41ER.

(3) If a person in relation to whom a kind of medical device is included in the Register shows that he or she has taken steps to address the grounds for revoking the certificate under section 41ET, the Secretary may, by written notice given to the person, extend the period specified in the notice under section 41EM by a further specified period not exceeding 6 months.

41EP Revocation of suspension

(1) The Secretary must revoke the suspension if the Secretary is satisfied that:
   (a) the ground on which the conformity assessment certificate was suspended no longer applies; and
   (b) there are no other grounds for suspending the certificate.

(2) The Secretary’s power to revoke the suspension may be exercised:
   (a) if:
(i) the manufacturer in relation to whom the conformity assessment certificate was issued; or
(ii) the person who applied for the certificate (if the applicant was not the manufacturer); applies in writing to the Secretary; or
(b) on the Secretary’s own initiative.

(3) After revoking the suspension, the Secretary must, within 20 working days after the revocation, give written notice of the revocation to the person in relation to whom the conformity assessment certificate was issued.

(4) If the Secretary decides, after an application is made under paragraph (2)(a), not to revoke the suspension, the Secretary must:
   (a) notify the applicant in writing of his or her decision within 20 working days after the decision is made; and
   (b) state in the notice the reasons for the decision.

41EQ Powers of revocation of conformity assessment certificates unaffected

(1) This Division does not affect the Secretary’s powers to revoke a conformity assessment certificate under Division 4.

(2) To the extent that a suspension under this Division relates to a conformity assessment certificate to which such a revocation relates, the suspension ceases to have effect.
Chapter 4  Medical devices
Part 4-4  Conformity assessment certificates
Division 4  Revocation of conformity assessment certificates

Section 41ER

Division 4—Revocation of conformity assessment certificates

Note: Revocation of a conformity assessment certificate leads to cancellation of the entry from the Register of the kinds of medical devices to which the certificate applied (see paragraph 41GK(b)). Applications to include such devices in the Register are not effective (see paragraph 41FC(2)(c)).

41ER  Automatic revocation of conformity assessment certificates

The Secretary must, by written notice given to the person in relation to whom a conformity assessment certificate is issued, revoke the certificate if:

(a) the certificate has been suspended under section 41EM; and
(b) the period applying to the suspension under subsection 41EM(3) or 41EO(3) (as the case requires) expires before the suspension is revoked under section 41EP.

41ES  Immediate revocation of conformity assessment certificates

The Secretary may, by written notice given to the manufacturer in relation to whom a conformity assessment certificate is issued, revoke the certificate if the manufacturer requests in writing the revocation of the certificate.

41ET  Revocation of conformity assessment certificates after notice of proposed revocation

(1) The Secretary may, by written notice given to the person in relation to whom a conformity assessment certificate is issued, revoke the certificate if:

(a) the conformity assessment procedures have not been applied to medical devices of a kind to which the certificate applies; or
(b) the manufacturer in relation to whom the certificate is issued refuses or fails to comply with a condition to which the certificate is subject; or
(c) the Secretary gives to the person a notice under section 41JA that requires the person to give to the Secretary information or documents relating to:
   (i) a kind of medical device; or
   (ii) a quality management system to which the certificate applies;

   and the person fails to comply with that notice within a further 10 working days from the day specified in that notice; or

(d) the manufacturer in respect of whom the certificate is issued no longer manufactures any of the kinds of medical devices to which the certificate applies; or

(e) the manufacturer is not a fit and proper person to be a manufacturer in respect of whom a conformity assessment certificate is issued; or

(f) a person who is participating in managing the manufacturer’s affairs is not a fit and proper person to participate in managing the affairs of a manufacturer in respect of whom a conformity assessment certificate is issued; or

(g) a person who has effective control over the manufacturer is not a fit and proper person to have effective control over a manufacturer in respect of whom a conformity assessment certificate is issued.

(1A) Without limiting the matters to which the Secretary may have regard in considering whether the holder or another person is not a fit and proper person for the purposes of paragraph (1)(e), (f) or (g), the Secretary must have regard to the matters set out in paragraphs 41EC(4)(a), (b) and (c).

(2) However, before revoking the certificate, the Secretary must:

   (a) inform the person in writing that the Secretary proposes the revocation and set out the reasons for it; and

   (b) give the person a reasonable opportunity to make submissions to the Secretary in relation to the proposed revocation.

(3) The Secretary is not to make a decision relating to the proposed revocation until the Secretary has had regard to any submissions the person makes under paragraph (2)(b).
Chapter 4  Medical devices  
Part 4-4  Conformity assessment certificates  
Division 4  Revocation of conformity assessment certificates

Section 41EU

41EU  Limiting revocation of conformity assessment certificates to some medical devices of a particular kind

(1) If the Secretary is satisfied that the ground for revoking a conformity assessment certificate applies only to:
   (a) some of the kinds of medical devices to which the certificate applies; or
   (b) some medical devices of the kinds to which the certificate applies;
   the Secretary must limit the revocation to the medical devices to which that ground or any other ground for revocation applies.

(2) If the revocation of the certificate is so limited, the Secretary must vary the certificate so that it no longer applies to the medical devices referred to in subsection (1).

41EV  Publication of revocation etc. of conformity assessment certificates

The Secretary must cause to be published in the Gazette, as soon as practicable after revoking a conformity assessment certificate, or varying a conformity assessment certificate under subsection 41EU(2), a notice setting out particulars of the revocation or variation.

41EW  Date of effect of revocation etc. of conformity assessment certificates

If the Secretary revokes a conformity assessment certificate, or varies a conformity assessment certificate under subsection 41EU(2), the revocation or variation has effect:

(a) if the revocation is under section 41ER or 41ES, or the variation relates to a ground of revocation in section 41ER or 41ES—on the day on which the notice of revocation or variation is given to the person in relation to whom the certificate was issued; or

(b) in any other case—on such later day as is specified in the notice.
Part 4-5—Including medical devices in the Register

41F What this Part is about

Kinds of medical devices can be included in the Register if they comply with the essential principles, and conformity assessment procedures have been applied to the kinds of devices (and certain other requirements are complied with).

Inclusions in the Register are subject to certain automatic conditions and the Secretary may impose further conditions.

Division 1—Including medical devices in the Register

41FA What this Division is about

Kinds of medical devices are usually included in the Register automatically once a proper application is made, together with the required certification. However, applications may be selected for audit, which involves checking some or all aspects of the application and certification.

Note 1: In some cases, an application relating to a kind of medical device will not be effective unless that kind of device is covered by a conformity assessment certificate under Part 4-4: see paragraph 41FC(2)(c).

Note 2: Dealing in medical devices of a kind not included in the Register may be an offence: see Division 3 of Part 4-11.

41FB How this Division works

This diagram shows how this Division applies to an application for a kind of medical device to be included in the Register.
Subdivision A—Applications

41FC Applications

(1) An application for a kind of medical device to be included in the Register must:
   (a) be made in accordance with a form approved, in writing, by the Secretary or in such other manner as is approved, in writing, by the Secretary; and
   (b) be delivered to an office of the Department specified by the Secretary.

(2) An application is not effective if:
   (a) the application is not made in accordance with subsection (1); or
   (b) the prescribed application fee has not been paid; or
   (c) regulations made for the purposes of section 41EA require the manufacturer of the kind of device to have a conformity assessment certificate relating to the kind of medical device.
before an application under this section can be made, and no such certificate is in force; or
(d) the application contains information that is false or misleading in a material particular.

Note: A person might also be guilty of an offence if the person makes a statement in an application that is false or misleading in a material particular: see section 41FE.

(3) An approval of a form may require or permit an application or information to be given in accordance with specified software requirements:
(a) on a specified kind of data processing device; or
(b) by way of a specified kind of electronic transmission.

41FD Matters to be certified

The applicant must certify that:
(a) devices of the kind in question are medical devices; and
(b) devices of that kind are intended for a specified purpose, as ascertained under subsection 41BD(2); and
(c) the kind of device is correctly classified according to the medical device classifications; and
(d) devices of that kind comply with the essential principles; and
(e) the applicant:
   (i) has available sufficient information to substantiate that compliance with the essential principles; or
   (ii) has procedures in place, including a written agreement with the manufacturer of the kind of devices setting out the matters required by the regulations, to ensure that such information can be obtained from the manufacturer within the period specified in the regulations; and
(f) an appropriate conformity assessment procedure has been applied to devices of that kind; and
(g) the applicant:
   (i) has available sufficient information to substantiate the application of those conformity assessment procedures; or
   (ii) has procedures in place, including a written agreement with the manufacturer of the kind of devices setting out
the matters required by the regulations, to ensure that such information can be obtained from the manufacturer within the period specified in the regulations; and

(h) devices of that kind comply with every requirement (if any) relating to advertising applicable under Part 5-1 or under the regulations; and

(i) devices of that kind do not contain substances that are prohibited imports for the purposes of the *Customs Act 1901*; and

(j) the information included in or with the application is complete and correct.

Note: See section 41BH on compliance with the essential principles and section 41BI on applying the conformity assessment procedures.

### 41FE False statements

A person is guilty of an offence if:

(a) the person makes a statement (whether orally, in a document or in any other way); and

(b) the person knows that the statement is false or misleading in a material particular; and

(c) the statement is in or in connection with:

(i) an application for including a kind of medical device in the Register under this Chapter; or

(ii) a certification or purported certification under section 41FD.

Maximum penalty: Imprisonment for 12 months or 1,000 penalty units, or both.

### Subdivision B—Including kinds of medical devices in the Register

#### 41FF Obligation to include kinds of medical devices in the Register

(1) If:

(a) an application is made in accordance with section 41FC for a kind of medical device to be included in the Register in relation to a person; and

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Section 41FG

(b) the requirements of section 41FD have been complied with; the Secretary must include the kind of device in the Register in relation to the person, unless the application has been selected under section 41FH for audit.

(2) As soon as practicable after the kind of device has been included in the Register, the Secretary must give to the applicant a certificate of the inclusion of the kind of device in the Register.

(3) The certificate must specify the day on which the inclusion of the kind of device in the Register commences.

41FG Notification of unsuccessful applications

The Secretary must notify the applicant in writing, within 20 working days after receiving an application under subsection 41FC(1), if an application for a kind of medical device to be included in the Register is unsuccessful.

Subdivision C—Auditing of applications

41FH Selecting applications for auditing

(1) The Secretary:
   (a) must select for auditing any application for a kind of medical device to be included in the Register that is an application of the kind prescribed by the regulations; and
   (b) may select for auditing any other application for a kind of medical device to be included in the Register.

Note: An application audit assessment fee is payable in respect of any application that the Secretary must select for auditing: see Part 4-10.

(2) If an application is selected for auditing:
   (a) the Secretary must, within 20 working days after the application is made, give the applicant a written notice that:
      (i) informs the applicant of the selection; and
      (ii) requests the applicant to provide any further information necessary for the auditing of the application; and
   (b) the application must be dealt with under this Subdivision and not under Subdivision B.
Chapter 4  Medical devices
Part 4-5  Including medical devices in the Register
Division 1  Including medical devices in the Register

Section 41FI

41FI  Auditing of applications

(1) In auditing the application, the Secretary may consider all or some aspects of one or both of the following matters:
   (a) whether the application is in accordance with Subdivision A;
   (b) whether matters as to which the applicant has certified under section 41FD are correct.

(2) The Secretary must decide to include the kind of device to which the application relates in the Register, in relation to the person to whom the application relates, if the Secretary is satisfied as to all such aspects considered in the audit.

(3) The Secretary must decide not to include the kind of device to which the application relates in the Register if the Secretary is not so satisfied.

41FJ  Procedure following audits

After auditing the application, the Secretary must:
   (a) notify the applicant in writing of his or her decision within 20 working days after the decision is made; and
   (b) if the decision is not to include the kind of device to which the application relates in the Register—state in the notice the reasons for the decision; and
   (c) if the decision is to include the kind of device in the Register and all assessment fees for the application that are due and payable have been paid:
      (i) include the kind of device in the Register, in relation to the person in relation to whom the application was made; and
      (ii) give the applicant a certificate of the inclusion of the kind of device in the Register.

41FK  Lapsing of applications

An application that has been selected for auditing lapses if:
   (a) the applicant does not deliver to the office to which the application was made such information (in a form approved

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in writing by the Secretary) as will allow the audit of the application; or

(b) the applicant does not comply with a requirement by the Secretary to deliver to the office to which the application was made a reasonable number of samples of the kind of medical device to which the application relates; or

(c) the applicant fails to comply with a notice under section 41JA to give information relating to devices of that kind to the Secretary within a further 10 working days from the day specified in the notice; or

(d) information given to the Secretary by, or on behalf of, the applicant in connection with the application, including information given for the purpose of a requirement under section 41JA, is false or misleading in a material particular; or

(e) the applicant fails to pay an assessment fee for the application within the period, specified in the regulations, after being notified of the decision to include the kind of medical device in the Register under section 41FJ.

Subdivision D—Miscellaneous

41FL Device number

If a kind of medical device is included in the Register, the Secretary is to assign a unique device number to it.

41FM Duration of inclusion in the Register

(1) The inclusion of a kind of medical device in the Register commences on the day specified for the purpose in the certificate under section 41FF.

(2) The inclusion of a kind of medical device in the Register has effect at all times:

(a) unless the kind of device is suspended from the Register under Division 1 of Part 4-6; or

(b) until entry of the kind of device is cancelled from the Register under Division 2 of Part 4-6.
41FN Conditions applying automatically

Entry and inspection powers

(1) The inclusion of a kind of medical device in the Register is subject to the conditions that the person in relation to whom the kind of device is included in the Register:

(a) allow an authorised person:

(i) to enter, at any reasonable time, any premises (including premises outside Australia) at which that person or any other person deals with medical devices of that kind; and

(ii) while on those premises, to inspect those premises and medical devices of that kind at those premises and to take samples of medical devices of that kind; and

(b) if requested to do so by an authorised person, produce to the person such documents relating to devices of that kind as the person requires and allow the person to copy the documents.

Delivery of samples

(2) The inclusion of a kind of medical device in the Register is subject to a condition that the person in relation to whom the kind of device is included in the Register will deliver a reasonable number of samples of the kind of device if the Secretary so requests:

(a) within the period specified in the request; and

(b) in accordance with any other requirements specified in the request.

The period specified in the request must include at least 10 working days.
Availability etc. of information

(3) The inclusion of a kind of medical device in the Register is subject to conditions that:

(a) at all times while the inclusion in the Register has effect, the person in relation to whom the kind of device is included in the Register:
   (i) has available sufficient information to substantiate compliance with the essential principles; or
   (ii) has procedures in place, including a written agreement with the manufacturer of the kind of devices setting out the matters required by the regulations, to ensure that such information can be obtained from the manufacturer within the period specified in the regulations; and

(b) at all times while the inclusion in the Register has effect, the person in relation to whom the kind of device is included in the Register:
   (i) has available sufficient information to substantiate that the conformity assessment procedures have been applied to the kind of medical device; or
   (ii) has available information relating to changes to the kind of medical device, the product range, and quality management system, of the manufacturer of the device; or
   (iii) has procedures in place, including a written agreement with the manufacturer of the kind of devices setting out the matters required by the regulations, to ensure that such information can be obtained from the manufacturer within the period specified in the regulations; and

(c) at any time while the inclusion in the Register has effect, the person in relation to whom the kind of device is included in the Register will, if asked to do so by the Secretary, give the information to the Secretary; and

(d) the person in relation to whom the kind of device is included in the Register will give information of a kind mentioned in subsection 41MP(2) to the Secretary within the period specified in the regulations; and

(e) the person in relation to whom the kind of device is included in the Register will give the manufacturer of the kind of medical device information relevant to:
(i) the manufacturer’s obligations under the conformity assessment procedures; and
(ii) whether medical devices of that kind comply with the essential principles.

(4) The regulations may prescribe the amount, standard or kind of information or evidence required for the purposes of paragraphs (3)(c), (d) and (e).

Advertising material

(5) The inclusion of a kind of medical device in the Register is subject to a condition that advertising material relating to medical devices of that kind is consistent with the intended purpose as certified under section 41FD.

Conditions do not limit other conditions

(6) A condition imposed under this section is in addition to any conditions imposed under this Division.

41FO Conditions imposed when kinds of medical devices are included in the Register

(1) If the Secretary includes a kind of medical device in the Register in relation to a person, the Secretary may, in writing, impose conditions on the inclusion of the kind of device in the Register.

(2) Conditions referred to in subsection (1) may relate to:
(a) manufacture of devices of that kind; or
(b) custody, intended purpose, supply, disposal or destruction of devices of that kind; or
(c) keeping of records relating to devices of that kind, including records relating to the tracking and location of devices of that kind after their supply; or
(d) matters dealt with in, or matters additional to matters dealt with in, the essential principles; or
(e) such other matters relating to devices of that kind as the Secretary thinks appropriate.
41FP Conditions imposed after kinds of medical devices are included in the Register

(1) The Secretary may, by written notice given to the person in relation to whom a kind of medical device is included in the Register:

(a) impose new conditions on including the kind of device in the Register under this Chapter; or

(b) vary or remove existing conditions.

The power may be exercised at the person’s request or on the Secretary’s own initiative.

(2) The imposition or variation of a condition under this section takes effect:

(a) if the notice states that the action is necessary to prevent imminent risk of death, serious illness or serious injury—on the day on which the notice is given to the person; or

(b) in any other case—on the day specified for the purpose in the notice, being a day not earlier than 20 working days after the notice is given to the person.
Part 4-6—Suspension and cancellation from the Register

Division 1—Suspension from the Register

Subdivision A—General power of suspension

41G What this Part is about

Inclusions in the Register may be suspended in certain circumstances, such as when a conformity assessment certificate is suspended. A kind of medical device that is suspended is taken not to be included in the Register for most purposes.

Inclusions in the Register may also be cancelled in certain circumstances.

41GA Suspension of kinds of medical devices from the Register

(1) The Secretary may, by written notice given to the person in relation to whom a kind of medical device is included in the Register, suspend the kind of device from the Register if:

(a) the Secretary is satisfied that:

(i) there is a potential risk of death, serious illness or serious injury if the kind of device continues to be included in the Register; and

(ii) it is likely that the person will, within the period of the suspension, be able to take the action necessary to ensure that the kind of device would not cause a potential risk of death, serious illness or serious injury if it were to continue to be included in the Register; or

(b) the Secretary is satisfied that it is likely that there are grounds for cancelling the entry of the kind of device from the Register under Division 2 (other than under paragraph 41GL(a) or (d) or section 41GM).
Section 41GB

(2) The suspension may be limited to some medical devices of that kind, as specified in the notice.

(3) The notice must specify the period of the suspension. The period must not exceed 6 months.

Note: The period of the suspension may be extended under section 41GC.

(4) The Secretary must cause to be published in the Gazette, as soon as practicable after the suspension, a notice setting out particulars of the suspension.

41GB Notice of proposed suspension must be given in certain cases

(1) However, before suspending a kind of medical device from the Register because it is likely that there are grounds for cancelling the entry of the kind of device from the Register under section 41GN, the Secretary must:
   (a) inform the person by written notice that the Secretary proposes the suspension and set out the reasons for it; and
   (b) give the person a reasonable opportunity to make submissions to the Secretary in relation to the proposed suspension.

(2) The Secretary is not to make a decision relating to the proposed suspension until the Secretary has had regard to any submissions the person makes under paragraph (1)(b).

41GC Duration of suspension

(1) The suspension takes effect:
   (a) if the notice under subsection 41GA(1) states that the suspension is necessary to prevent a potential risk of death, serious illness or serious injury—on the day on which the notice is given to the person; or
   (b) in any other case—on the day specified for the purpose in the notice, being a day not earlier than 20 working days after the notice is given to the person.

(2) The suspension has effect until:
   (a) the Secretary revokes it under section 41GD; or
   (b) the end of:
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Division 1 Suspension from the Register

Section 41GD

(i) the period specified in the notice under subsection 41GA(3); or
(ii) if the period is extended under subsection (3) of this section, the period as so extended.

Note: Unless a suspension of a kind of medical device has been revoked, the entry of the kind of medical device is automatically cancelled from the Register: see section 41GK.

(3) If a person in relation to whom a kind of medical device is included in the Register shows that he or she has taken steps to remove the grounds for cancelling the entry of the kind of device from the Register under section 41GN, the Secretary may, by written notice given to the person, extend the period specified in the notice under subsection 41GA(1) by a further specified period not exceeding 6 months.

(4) The Secretary must cause to be published in the Gazette, as soon as practicable after the extension, a notice setting out particulars of the extension.

41GD Revocation of suspension

(1) The Secretary must revoke the suspension if the Secretary is satisfied that:
   (a) the ground on which the kind of medical device concerned was suspended from the Register no longer applies; and
   (b) there are no other grounds for suspending the kind of device from the Register.

(2) The Secretary’s power to revoke the suspension may be exercised:
   (a) if the person in relation to whom the kind of medical device concerned is included in the Register applies in writing to the Secretary; or
   (b) on the Secretary’s own initiative.

(3) After revoking the suspension, the Secretary must:
   (a) within 20 working days after the revocation, give written notice of the revocation to the person in relation to whom the kind of medical device concerned is included in the Register; and

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(b) as soon as practicable after the revocation, cause to be published in the Gazette a notice setting out particulars of the revocation.

(4) If the Secretary decides, after an application is made under paragraph (2)(a), not to revoke the suspension, the Secretary must:
(a) notify the applicant in writing of his or her decision within 20 working days after the decision is made; and
(b) state in the notice the reasons for the decision.

41GE Treating applications for revocation as having been refused

(1) The applicant for the suspension to be revoked may give the Secretary written notice that the applicant wishes to treat the application as having been refused if:
(a) a period is prescribed under paragraph 63(2)(dd) for the Secretary to make a decision on the application; and
(b) at the end of the period, the Secretary has not made a decision.

(2) The notice may be given at any time before the Secretary makes a decision on the application.

(3) If a notice has been given, this Act (except for subsection 60(5)) has effect as if:
(a) the Secretary had decided not to revoke the suspension; and
(b) the Minister had made a decision under subsection 60(3) confirming the decision of the Secretary; and
(c) the Minister’s decision had been made on the day on which notice was given to the Secretary.

Subdivision B—Suspension as a result of suspension under Part 4-4

41GF Suspension of kinds of medical devices from the Register

(1) The Secretary may, by written notice given to the person in relation to whom a kind of medical device is included in the Register, suspend the kind of device from the Register if the conformity
assessment certificate applying to that kind of device is suspended under Division 3 of Part 4-4.

(2) If the suspension under Division 3 of Part 4-4 is limited to some medical devices of that kind, the suspension under this section is taken to be limited to the same extent.

(3) The Secretary must cause to be published in the Gazette, as soon as practicable after the suspension, a notice setting out particulars of the suspension.

41GG Duration of suspension

(1) The suspension takes effect on the day on which the notice is given to the person.

(2) The suspension has effect until the Secretary revokes it under section 41GH.

41GH Revocation of suspension

(1) The Secretary must revoke the suspension if:
   (a) the suspension under Division 3 of Part 4-4 ceases to have effect; and
   (b) the Secretary is satisfied that there are no other grounds for suspending the kind of device from the Register.

(2) After revoking the suspension, the Secretary must:
   (a) within 20 working days after the revocation, give written notice of the revocation to the person in relation to whom the kind of medical device concerned is included in the Register; and
   (b) as soon as practicable after the revocation, cause to be published in the Gazette a notice setting out particulars of the revocation.
Subdivision C—Effect of suspension

41GI  Effect of suspension

If all or some medical devices of a particular kind are suspended, they are taken, for the purposes of this Act (other than Division 2 of Part 4-5, this Division and Part 4-8), not to be included in the Register while the suspension has effect.

Note: Dealing in medical devices that are not included in the Register may be an offence: see Division 3 of Part 4-11.

41GJ  Powers of cancellation from Register unaffected

(1) This Subdivision does not affect the Secretary’s powers to cancel the entry of kinds of medical devices from the Register under Division 2.

(2) To the extent that a suspension under this Division relates to medical devices to which such a cancellation relates, the suspension ceases to have effect.
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Part 4-6  Suspension and cancellation from the Register
Division 2  Cancellation of entries from the Register

Section 41GK

Division 2—Cancellation of entries from the Register

41GK  Automatic cancellation of entries of kinds of medical devices from the Register

The Secretary must, by written notice given to the person in relation to whom a kind of medical device is included in the Register, cancel the entry of the kind of device from the Register if:

(a) the kind of device has been suspended from the Register under section 41GA, and the period applying to the suspension under subsection 41GA(3) or 41GC(3) (as the case requires) expires before the suspension is revoked under section 41GD; or

(b) a conformity assessment certificate applying to that kind of device is revoked under Division 4 of Part 4-4.

41GL  Immediate cancellation of entries of kinds of medical devices from the Register

The Secretary may, by written notice given to the person in relation to whom a kind of medical device is included in the Register, cancel the entry of the kind of device from the Register if:

(a) the Secretary is satisfied that there would be an imminent risk of death, serious illness or serious injury if the kind of device continues to be included in the Register; or

(b) devices of that kind are no longer therapeutic goods; or

(c) devices of that kind are no longer medical devices; or

(d) the person requests in writing the cancellation of the entry of the kind of device from the Register; or

(e) the Secretary is satisfied that a statement made in or in connection with:

(i) the application for including the kind of device in the Register; or

(ii) the certification or purported certification under section 41FD relating to the application; was false or misleading in a material particular; or
the annual charge payable under subsection 4(1B) of the
Therapeutic Goods (Charges) Act 1989 in respect of the
inclusion of the kind of device in the Register is not paid
within 20 working days after it becomes payable; or
(g) both of the following apply:
   (i) under the regulations, an authority constituted by or
       under the regulations gives a direction to, or makes a
       requirement of, the person in relation to an
       advertisement of the kind of device to ensure that
       advertising complies with the Therapeutic Goods
       Advertising Code;
   (ii) the person does not comply with the direction or
        requirement; or
(h) there is a serious breach, involving the kind of device, of the
   requirements relating to advertising applicable under Part 5-1
   or under the regulations, and the Secretary is satisfied that:
   (i) the breach is significant; and
   (ii) as a result of the breach, the presentation of devices of
       that kind is misleading to a significant extent.

41GM Cancellation of entries of kinds of medical devices from the
Register after section 41JA notice

(1) The Secretary may, by written notice given to the person in relation
to whom a kind of medical device is included in the Register,
cancel the entry of the kind of device from the Register if:
   (a) the Secretary gives to the person a notice under section 41JA
       requiring the person to give to the Secretary information or
       documents relating to the kind of device; and
   (b) the notice under section 41JA is given for the purposes of
       ascertaining whether the kind of device should have been
       included in the Register; and
   (c) the person fails to comply with the notice under section 41JA
       within a further 10 working days from the day specified in
       that notice.

(2) The Secretary may, by written notice given to the person in relation
to whom a kind of medical device is included in the Register,
cancel the entry of the kind of device from the Register if:
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Section 41GN

(a) the Secretary gives to the person a notice under section 41JA requiring the person to give to the Secretary information or documents relating to whether medical devices of that kind are being:

(i) supplied in Australia; or

(ii) imported into Australia; or

(iii) exported from Australia; and

(b) either:

(i) the information or documents given are to the effect that medical devices of that kind are not being supplied in Australia, imported into Australia or exported from Australia; or

(ii) the person fails to comply with the notice under section 41JA within a further 10 working days from the day specified in that notice.

41GN  Cancellation of entries of kinds of medical devices from the Register after notice of proposed cancellation

(1) The Secretary may, by written notice given to the person in relation to whom a kind of medical device is included in the Register, cancel the entry of the kind of device from the Register if:

(a) medical devices that were devices of that kind when the kind of device was included in the Register have changed so those medical devices are no longer devices of that kind; or

(b) the person in relation to whom the kind of medical device is included in the Register refuses or fails to comply with a condition to which that inclusion is subject; or

(c) the Secretary gives to the person a notice under section 41JA:

(i) that requires the person to give to the Secretary information or documents relating to the kind of device; and

(ii) in respect of which section 41GM does not apply; and the person fails to comply with that notice within a further 10 working days from the day specified in that notice; or

(d) the person contravenes subsection 41MP(1) in relation to the kind of device; or

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Section 41GO

(e) the Secretary is satisfied that the safety or performance of the kind of device is unacceptable; or

(f) the Secretary is satisfied that any certification, or part of a certification, under section 41FD in relation to the application for inclusion of the kind of device in the Register is incorrect, or is no longer correct, in a material particular.

Note: The matters that must be certified under section 41FD include compliance with the essential principles and the application of conformity assessment procedures, being able to substantiate the compliance and application, and compliance with advertising requirements.

(2) However, before cancelling the entry of the kind of device from the Register, the Secretary must:

(a) inform the person in writing that the Secretary proposes the cancellation and set out the reasons for it; and

(b) give the person a reasonable opportunity to make submissions to the Secretary in relation to the proposed cancellation.

(3) The Secretary is not to make a decision relating to the proposed cancellation until the Secretary has had regard to any submissions the person makes under paragraph (2)(b).

41GO Limiting cancellation of entries from Register to some medical devices of a particular kind

(1) If the Secretary is satisfied that the ground for cancelling the entry of a kind of medical device from the Register applies only to some medical devices of that kind, the Secretary must limit the cancellation to the medical devices to which that ground or any other ground for cancellation applies.

(2) If the cancellation of the entry of a kind of medical device from the Register is limited to some medical devices of that kind, the Secretary:

(a) must vary the entry in the Register accordingly; and

(b) must not delete the entry from the Register because of the cancellation.
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Part 4-6  Suspension and cancellation from the Register  
Division 2  Cancellation of entries from the Register

Section 41GP

41GP  Publication of cancellation of entry from Register

The Secretary must cause to be published in the Gazette, as soon as practicable after cancelling an entry from the Register of a kind of medical device, or of some devices of a particular kind, a notice setting out particulars of the cancellation.

41GQ  Date of effect of cancellation of entries from Register

If the Secretary cancels an entry of a kind of medical device, or some devices of a particular kind, from the Register, the cancellation has effect:

(a) if the cancellation is under section 41GK or 41GL—on the day on which the notice of cancellation is given to the person in relation to whom the kind of device was included in the Register; or

(b) in any other case—on such later day as is specified in the notice, being a day not earlier than 20 working days after the notice is given to the person.
Part 4-7—Exempting medical devices from inclusion in the Register

41H  What this Part is about

There are 3 kinds of exemptions from the prohibitions in Division 3 of Part 4-11 on dealing in medical devices that are not included in the Register:

(a) medical devices exempted under the regulations;

(b) approval for medical devices to be used for special treatment of individuals or for experimental purposes;

(c) authorisation of particular medical practitioners to supply specified medical devices.

41HA  Devices exempted from inclusion in the Register

(1) The regulations may exempt from the operation of Division 3 of Part 4-11:

(a) all medical devices, except those medical devices of the kinds prescribed for the purposes of this paragraph; or

(b) specified kinds of medical devices.

Note: Division 3 of Part 4-11 contains offences relating to dealing in medical devices that are not included in the Register.

(2) An exemption may be subject to conditions that are prescribed in the regulations.

Note: Breach of the conditions may be an offence: see subsection 41MN(3).

(3) An exemption under paragraph (1)(a) has effect only for classes of persons prescribed in the regulations for the purposes of this subsection.
Section 41HB

(4) If the regulations revoke an exemption, the revocation takes effect on the day specified. The day must not be earlier than 20 working days after the day on which the regulations are made.

41HB Exemptions for special and experimental uses

(1) The Secretary may grant a written approval to a person for:
   (a) the importation into Australia; or
   (b) the exportation from Australia; or
   (c) the supply in Australia;
   of a specified medical device or kind of medical device (other than medical devices included in the Register or exempt devices):
   (d) for use in the treatment of another person; or
   (e) for use solely for experimental purposes in humans.

(2) The approval may be given subject to conditions specified in the approval, including a condition relating to charging for medical devices of the kinds in question.

Note: Breach of the conditions may be an offence: see subsection 41MN(3).

(3) In addition, the regulations may prescribe conditions that apply to a person’s approval to use specified kinds of medical devices solely for experimental purposes in humans. The conditions may relate to one or more of the following:
   (a) the preconditions on another person’s use of devices of those kinds for those purposes;
   (b) the principles to be followed in another person’s use of devices of those kinds for those purposes;
   (c) the monitoring of another person’s use, and the results of that use, of devices of those kinds for those purposes;
   (d) the circumstances in which that other person must cease using devices of those kinds for those purposes.

(4) An application to use specified medical devices in the treatment of another person must be accompanied by any information about the devices that is required by the Secretary.

(5) An application to use specified kinds of medical devices solely for experimental purposes in humans must:
   (a) be made in writing; and
(b) be accompanied by any information about the kinds of devices that is required by the Secretary; and
(c) be accompanied by the prescribed fee.

(6) The Secretary must:
(a) consider any application under this section; and
(b) assess any information submitted with the application; and
(c) notify the applicant, within 20 working days of making the decision:
(i) of the decision; and
(ii) in the case of a decision not to grant the approval—of the reasons for the decision.

(7) The use by a person for experimental purposes in humans of specified kinds of medical devices that are the subject of an approval granted to someone else under paragraph (1)(e) is subject to the conditions (if any) specified in the regulations relating to one or more of the following:
(a) the preconditions on the use of devices of those kinds for those purposes;
(b) the principles to be followed in the use of devices of those kinds for those purposes;
(c) the monitoring of the use, and the results of the use, of devices of those kinds for those purposes;
(d) the circumstances in which the person must cease the use of devices of those kinds for those purposes.

Note: Breach of the conditions may be an offence: see subsection 41MN(3).

41HC Exemptions for medical practitioners

(1) The Secretary may authorise, in writing, a specified medical practitioner to supply specified kinds of medical devices for use in the treatment of humans to a specified class of recipients.

(2) An authority may be given subject to conditions specified in the authority.

(3) The Secretary may impose conditions (or further conditions) on a person’s authority by giving the person written notice of the conditions.
(4) An authority may only be given:

(a) to a medical practitioner included in a class of medical practitioners prescribed by the regulations for the purposes of this paragraph; and

(b) to a medical practitioner who has the approval of an ethics committee to supply the specified kinds of medical devices or the specified class of such devices; and

(c) in relation to a class or classes of recipients prescribed by the regulations for the purposes of this paragraph. However, the regulations may prescribe exceptional circumstances in which paragraph (b) does not apply.

(5) The regulations may prescribe circumstances in which medical devices may be supplied under an authority.

(6) The giving of an authority in respect of kinds of medical devices does not render the Commonwealth, the Secretary or a delegate of the Secretary liable to a person for loss, damage or injury of any kind suffered by the person as a result of, or arising out of, the use of devices of those kinds by that person or another person.

(7) In this section:

*medical practitioner* means a person who is registered, in a State or internal Territory, as a medical practitioner.
Part 4-8—Obtaining information

41J What this Part is about

The Secretary may seek information or documents relating to:

- the application of conformity assessment procedures;
- compliance with the essential principles;
- compliance with other requirements;
- distribution of, and other matters relating to, medical devices covered by exemptions under Part 4-7.

Note: There are additional obligations relating to notifying defects in medical devices: see sections 41MP and 41MQ.

Division 1—Information relating to compliance with requirements and other matters

41JA Secretary may require information

(1) The Secretary may, by written notice given to a person:

(a) who is an applicant for a conformity assessment certificate that would relate to a kind of medical device; or
(b) who holds a conformity assessment certificate that relates to a kind of medical device; or
(c) who is an applicant for the inclusion of a kind of medical device in the Register; or
(d) in relation to whom a kind of medical device is, or was at any time during the notice period under subsection (2), included in the Register;

require the person to give to the Secretary information or documents, relating to devices of that kind, that are relevant to one or more of the following:

(e) whether the devices comply with the essential principles;
(f) whether the conformity assessment procedures have been applied to the devices;

(g) whether the devices comply with conditions (if any) imposed on a conformity assessment certificate issued in respect of the device or the inclusion of the device in the Register;

(h) whether the devices comply with every requirement (if any) relating to advertising applicable under Part 5-1 or under the regulations;

(i) if the kind of medical device is included in the Register in relation to the person—whether medical devices of that kind are being:
   (i) supplied in Australia; or
   (ii) imported into Australia; or
   (iii) exported from Australia;

(j) any other matter prescribed by the regulations for the purposes of this paragraph.

(2) For the purposes of paragraph (1)(d), the notice period is the period:

(a) of the length specified in the regulations; and

(b) ending on the day before the Secretary gives the notice under subsection (1).

41JB Complying with the Secretary’s requirements

(1) The person must give the information or documents to the Secretary:

   (a) within such reasonable time, being not less than 10 working days from the day on which the notice is given, as is specified in the notice; and

   (b) in such form as is specified in the notice.

(2) The form may require or permit information to be given in accordance with specified software requirements:

   (a) on a specified kind of data processing device; or

   (b) by way of a specified kind of electronic transmission.

(3) A person is guilty of an offence if the person:

   (a) is a person in relation to whom a notice is given under section 41JA; and

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(b) fails to comply with the notice.

Maximum penalty: 30 penalty units.

Note: Failure to comply with the notice might also lead to suspension or revocation of a conformity assessment certificate (see Divisions 3 and 4 of Part 4-4) or suspension or cancellation of the entry of a kind of medical device in the Register (see Part 4-6).

(4) A person is guilty of an offence if:
   (a) the person is a person in relation to whom a notice is given under section 41JA; and
   (b) the person gives information in purported compliance with the notice; and
   (c) the information is false or misleading in a material particular.

Maximum penalty: 60 penalty units.

41JC Self-incrimination

(1) A person is not excused from giving information or a document under section 41JB on the ground that to do so would tend to incriminate the person or expose the person to a penalty.

(2) However, in the case of an individual:
   (a) the information given; or
   (b) the giving of the document; or
   (c) any information, document or thing obtained as a direct or indirect consequence of giving the information or document; is not admissible in evidence in criminal proceedings against the individual, except proceedings under, or arising out of, subsection 41JB(4).
Division 2—Information relating to medical devices covered by exemptions

41JD Secretary may require information etc. about devices exempted under section 41HA from inclusion in the Register

(1) The Secretary may give the sponsor of kinds of medical devices exempted under subsection 41HA(1) from Division 3 of Part 4-11, a written notice requiring the sponsor to give to the Secretary specified information or documents relating to one or more of the following:
   (a) the supply of devices of those kinds;
   (b) the handling of devices of those kinds;
   (c) the monitoring of the supply of devices of those kinds;
   (d) the results of the supply of devices of those kinds;
   (e) any other matter prescribed by the regulations for the purposes of this paragraph in relation to devices of those kinds.

(2) If a medical device is exempt under subsection 41HA(1) because a medical practitioner has signed a statement in accordance with regulations made for the purposes of this section, the Secretary may give the medical practitioner a written notice requiring the medical practitioner to give to the Secretary specified information or documents relating to one or more of the following:
   (a) the condition of the person to whom the medical device is to be given or is given;
   (b) the supply of the device;
   (c) the handling of the device;
   (d) the monitoring of the supply of the device;
   (e) the results of the supply of the device;
   (f) any other matter prescribed by the regulations for the purposes of this paragraph in relation to medical devices of that kind.
Section 41JE

(3) A notice under this section must specify a reasonable period within which the person must comply. The period must be at least 10 working days starting on the day on which the notice is given.

41JE Secretary may require information relating to approvals under section 41HB

Approval under subsection 41HB(1)

(1) The Secretary may give to a person granted an approval under subsection 41HB(1) (special and experimental uses), in relation to specified kinds of medical devices, a written notice requiring the person to give to the Secretary specified information or documents relating to one or more of the following:
   (a) the supply of devices of those kinds;
   (b) the handling of devices of those kinds;
   (c) the monitoring of the supply of devices of those kinds;
   (d) the results of the supply of devices of those kinds;
   (e) any other matter prescribed by the regulations for the purposes of this paragraph in relation to devices of those kinds.

Approval under subsection 41HB(1)—use by another person

(2) The Secretary may give to a person using specified kinds of medical devices, that are the subject of an approval granted to someone else under paragraph 41HB(1)(e) (use solely for experimental purposes in humans), a written notice requiring the person to give to the Secretary specified information or documents relating to either of both of the following:
   (a) the use of devices of those kinds;
   (b) any other matter prescribed by the regulations for the purposes of this paragraph in relation to devices of those kinds.

Compliance period

(3) A notice under this section must specify a reasonable period within which the person to whom the notice is given must comply. The
Section 41JF

period must be at least 10 working days starting on the day on which the notice is given.

41JF Secretary may require information relating to authorities under section 41HC

(1) The Secretary may give to a person who is granted an authority under section 41HC (exemptions for medical practitioners), in relation to specified kinds of medical devices, a written notice requiring the person to give to the Secretary specified information or documents relating to one or more of the following:
(a) the supply of devices of those kinds;
(b) the handling of devices of those kinds;
(c) the monitoring of the supply of devices of those kinds;
(d) the results of the supply of devices of those kinds;
(e) any other matter prescribed by the regulations for the purposes of this paragraph in relation to devices of those kinds.

(2) The notice must specify a reasonable period within which the person to whom the notice is given must comply. The period must be at least 10 working days starting on the day on which the notice is given.

41JG Requirements in relation to information or documents sought under this Division

When information or documents must be given etc.

(1) A person to whom a notice is given under section 41JD, 41JE or 41JF must give to the Secretary, within the period specified in the notice:
(a) the information specified in the notice; and
(b) the documents specified in the notice in the form (if any) specified in the notice.

Way in which information given

(2) The notice may require information to be given in accordance with specified software requirements:
(a) on a specified kind of data processing device; or
(b) by way of a specified kind of electronic transmission.

**Offence**

(3) A person is guilty of an offence if the person fails to comply with subsection (1).

*Note:* The privilege against self incrimination is not a reasonable excuse for the purposes of this subsection. However, section 41J limits the use in prosecutions of information etc. obtained under sections 41JD, 41JE and 41JF.

(4) An offence against subsection (3) is punishable on conviction by a fine of not more than 30 penalty units.

**41JH False or misleading information**

A person to whom a notice is given under section 41JD, 41JE or 41JF is guilty of an offence if:

(a) the person gives information to the Secretary; and
(b) the person knows that the information:
   (i) is false or misleading; or
   (ii) omits any matter or thing without which the information is misleading; and
(c) the information is given in compliance or purported compliance with subsection 41JG(1).

Maximum penalty: 60 penalty units.

**41JI False or misleading documents**

(1) A person is guilty of an offence if:

(a) the person produces a document to the Secretary; and
(b) the person knows that the document is false or misleading; and
(c) the document is produced in compliance or purported compliance with subsection 41JG(1).

Maximum penalty: 60 penalty units.
Section 41JJ

(2) Subsection (1) does not apply to a person who produces a document if the document is accompanied by a written statement signed by the person or, in the case of a body corporate, by a competent officer of the body corporate:

(a) stating that the document is, to the knowledge of the first-mentioned person, false or misleading in a material particular; and

(b) setting out, or referring to, the material particular in which the document is, to the knowledge of the first-mentioned person, false or misleading.

Note: A defendant bears an evidential burden in relation to the matter in subsection (2) (see subsection 13.3(3) of the Criminal Code).

41JJ Self-incrimination

(1) A person is not excused from giving information or a document under section 41JG on the ground that to do so would tend to incriminate the person or expose the person to a penalty.

(2) However, in the case of an individual:

(a) the information given; or

(b) the giving of the document; or

(c) any information, document or thing obtained as a direct or indirect consequence of giving the information or document; is not admissible in evidence in criminal proceedings against the individual, except proceedings under, or arising out of, section 41JH or 41JI.
Part 4-9—Public notification and recovery of medical devices

41K What this Part is about

The Secretary can require action to recover medical devices, or to inform the public about medical devices, that do not comply with requirements or cannot lawfully be supplied.

41KA Public notification and recovery of medical devices

(1) The Secretary may, in writing, impose requirements, relating to a kind of medical device, on a person if:

   (a) any of the circumstances referred to in the second column of an item in the following table occur in relation to the kind of device; and

   (b) the person is referred to in the third column of that item of the table.

<table>
<thead>
<tr>
<th>Item</th>
<th>Circumstance relating to a kind of medical device</th>
<th>Person subject to requirements</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>It is supplied while it is included in the Register, but medical devices of that kind do not comply with the essential principles</td>
<td>The person in relation to whom it is included in the Register</td>
</tr>
<tr>
<td>2.</td>
<td>It is supplied while it is included in the Register, but the conformity assessment procedures have not been applied to medical devices of that kind</td>
<td>The person in relation to whom it is included in the Register</td>
</tr>
</tbody>
</table>
### Section 41KA

#### Circumstances in which requirements may be imposed

<table>
<thead>
<tr>
<th>Item</th>
<th>Circumstance relating to a kind of medical device</th>
<th>Person subject to requirements</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.</td>
<td>It is supplied while:</td>
<td>The person supplying the kind of medical device</td>
</tr>
<tr>
<td></td>
<td>(a) medical devices of that kind are exempt devices; or</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(b) there is an approval under section 41HB relating to devices of that kind; or</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(c) there is an authority under section 41HC relating to devices of that kind; but medical devices of that kind do not comply with the essential principles</td>
<td></td>
</tr>
<tr>
<td>4.</td>
<td>It is supplied while:</td>
<td>The person supplying the kind of medical device</td>
</tr>
<tr>
<td></td>
<td>(a) medical devices of that kind are exempt devices; or</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(b) there is an approval under section 41HB relating to devices of that kind; or</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(c) there is an authority under section 41HC relating to devices of that kind; but the conformity assessment procedures have not been applied to medical devices of that kind</td>
<td></td>
</tr>
<tr>
<td>5.</td>
<td>It is supplied while:</td>
<td>The person supplying the kind of medical device</td>
</tr>
<tr>
<td></td>
<td>(a) it is not included in the Register; and</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(b) it is not an exempt device; and</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(c) there is not an approval under section 41HB relating to devices of that kind; and</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(d) there is not an authority under section 41HC relating to devices of that kind.</td>
<td></td>
</tr>
<tr>
<td>6.</td>
<td>It has been suspended from the Register</td>
<td>The person in relation to whom it was included in the Register</td>
</tr>
<tr>
<td>7.</td>
<td>Its entry has been cancelled from the Register</td>
<td>The person in relation to whom it was included in the Register</td>
</tr>
</tbody>
</table>

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(2) The requirements may be one or both of the following:
(a) to take specified steps, in the specified manner and within such reasonable period as is specified, to recover medical devices of that kind that have been distributed;
(b) to inform the public or a specified class of persons, in the specified manner and within such reasonable period as is specified, to the effect that the circumstances referred to in paragraph (1)(a) have occurred in relation to medical devices of that kind.

(3) If the circumstances referred to in paragraph (1)(a) apply only to some medical devices of that kind, the Secretary may limit the imposition of the requirements to the medical devices of that kind to which those circumstances apply.

(4) A requirement to recover medical devices under this section does not apply to a medical device that cannot be recovered because it has been administered to, or applied in the treatment of, a person.

41KB Publication of requirements

The Secretary must cause to be published in the Gazette, as soon as practicable after imposing a requirement under section 41KA, a notice setting out particulars of the requirement.

41KC Non-compliance with requirements

A person is guilty of an offence if:
(a) the person does an act, or omits to do an act; and
(b) the act or omission constitutes a contravention of a requirement imposed on the person under section 41KA.

Maximum penalty: 60 penalty units.

41KD Powers of suspension and cancellation unaffected

Imposition of a requirement under section 41KA does not affect the Secretary’s powers to:
(a) suspend the entry of a kind of medical device, or some medical devices of a particular kind, from the Register under Part 4-6; or
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(b) cancel the entry of a kind of medical device, or some medical devices of a particular kind, in the Register under Part 4-6.
Part 4-10—Assessment fees

41L. What this Part is about

Conformity assessment fees must be paid for consideration of applications for conformity assessment certificates. Application audit assessment fees must be paid for auditing applications that are required to be selected for auditing under paragraph 41FH(1)(a).

41LA Assessment fees

(1) A conformity assessment fee specified in or determined in accordance with the regulations is payable by a person in respect of consideration of an application for a conformity assessment certificate under Part 4-4.

(2) The regulations may prescribe different levels of conformity assessment fees in relation to any one or more of the following:
   (a) different kinds of manufacturers;
   (b) different kinds of medical devices;
   (c) different parts of the conformity assessment procedures that are considered in relation to an application for a conformity assessment certificate under Part 4-4.

(3) An application audit assessment fee specified in or determined in accordance with the regulations is payable by a person in respect of the auditing of an application for inclusion of a kind of medical device in the Register under Part 4-5, if paragraph 41FH(1)(a) required the Secretary to select the application for audit.

(4) The regulations may prescribe different levels of application audit assessment fees in relation to any one or more of the following:
   (a) different kinds of manufacturers;
   (b) different kinds of medical devices;
   (c) different levels of assessment of kinds of medical devices.
Section 41LB

(5) The application audit assessment fee payable because of subsection (3) is payable only in respect of considering the matters set out in subsection 41FI(1).

41LB  When assessment fee due for payment

Subject to sections 41LC and 41LE, an assessment fee payable by an applicant is due and payable on the day, and in the manner, specified in the regulations.

41LC  Payment of assessment fee by instalments

(1) The regulations may provide for the payment of an assessment fee to be made by such instalments and at such times as are ascertained in accordance with the regulations, and the assessment fee is due and payable accordingly.

(2) Regulations made for the purposes of subsection (1) may provide that a person is not allowed to pay an assessment fee by instalments if any part of an instalment of:
   (a) that or any other assessment fee payable by the person; or
   (b) any evaluation fee under section 24 payable by the person;
   was unpaid immediately after the time when it became due for payment.

(3) Subsection (2) does not limit the generality of subsection (1).

41LD  Recovery of assessment fee

An assessment fee may be recovered by the Commonwealth as a debt due to the Commonwealth.

41LE  Reduction of conformity assessment fee where decision not made within prescribed period

(1) Nothing in section 41LA, 41LB or 41LC requires the applicant to pay more than $3/4 of the conformity assessment fee before the making of the decision if:
   (a) the application is for the issuing of a conformity assessment certificate under Part 4-4; and
Section 41LE

(b) consideration of the application will involve an examination of the design of medical devices; and
(c) a period is prescribed under paragraph 63(2)(dc) for making a decision on the application.

(2) If the decision is not made within that period, the conformity assessment fee is $\frac{3}{4}$ of the fee that, apart from this subsection, would have been the conformity assessment fee.

(3) If:
   (a) the decision is made within that period; and
   (b) part of the conformity assessment fee under section 41LA is, because of subsection (1) of this section, unpaid when the decision is made;
   
   that part becomes due and payable on the making of the decision.

(4) For the purposes of this section, a decision is taken to be made on the application when the applicant is notified under subsection 41EE(1) of the Secretary’s decision on the application.
Chapter 4  Medical devices
Part 4-11  Offences relating to medical devices
Division 1  Non-compliance with essential principles

Section 41M

Part 4-11—Offences relating to medical devices

41M  What this Part is about

This Part contains offences that are aimed at ensuring that:

- the essential principles are complied with (see Division 1);
- the conformity assessment procedures have been applied to kinds of medical devices (see Division 2);
- administrative processes put in place by Parts 4-4 to 4-9 are followed (see Divisions 3 and 4).

Note: There are also some offences in the earlier Parts of this Chapter. They generally relate to matters ancillary to administrative processes in those Parts (e.g. false or misleading statements in applications).

Division 1—Non-compliance with essential principles

41MA  Non-compliance with essential principles

(1) A person is guilty of an offence if:

- the person imports a medical device into Australia; and
- the medical device does not comply with the essential principles relating to matters other than the labelling of the device; and
- the Secretary has not consented to the importation.

Maximum penalty: Imprisonment for 12 months or 1,000 penalty units, or both.

(2) A person is guilty of an offence if:

- the person supplies a medical device for use in Australia; and
- the medical device does not comply with the essential principles; and
- the Secretary has not consented to the supply.
Chapter 4

Offences relating to medical devices

Part 4-11

Non-compliance with essential principles

Division 1

Section 41MB

Maximum penalty: Imprisonment for 12 months or 1,000 penalty units, or both.

(3) A person is guilty of an offence if:
   (a) the person exports a medical device from Australia; and
   (b) the medical device does not comply with the essential principles; and
   (c) the Secretary has not consented to the exportation.

Maximum penalty: Imprisonment for 12 months or 1,000 penalty units, or both.

(4) Paragraph (3)(b) does not apply to the extent that the essential principles in question relate to labelling medical devices for supply in Australia.

Note: A defendant bears an evidential burden in relation to the matters in this subsection (see subsection 13.3(3) of the Criminal Code).

(5) The Secretary must not give consent relating to an exportation unless satisfied that there are exceptional circumstances that justify giving the consent.

41MB Exceptions

(1) Section 41MA does not apply if:
   (a) the medical device complies with one or more medical device standards that apply to it; and
   (b) the medical device fails to comply with the essential principles only in respect of a part or parts of the essential principles to which that medical device standard, or one or more of those medical device standards, relate.

Note: Medical device standards are determined under Division 2 of Part 4-2.

(2) For the purposes of this section, a medical device standard relates to a part or parts of the essential principles only if the standard specifies that part or parts.

Note: A defendant bears an evidential burden in relation to the matters in this section (see subsection 13.3(3) of the Criminal Code).
Section 41MC

41MC  Consent may be subject to conditions etc.

(1) The consent of the Secretary under section 41MA may be given:
   (a) unconditionally or subject to conditions; or
   (b) in respect of particular medical devices or kinds of medical devices.

(2) A person is guilty of an offence if:
   (a) the person does an act, or omits to do an act; and
   (b) the act or omission constitutes a breach of a condition of such a consent.

Maximum penalty:  120 penalty units.

41MD  Treating medical devices as prohibited imports or exports

If:
   (a) the importation or exportation of a medical device is prohibited under subsection 41MA(1) or (3); and
   (b) the Secretary notifies the Chief Executive Officer of Customs in writing that the Secretary wishes the Customs Act 1901 to apply to that importation or exportation;

   the Customs Act 1901 has effect as if the device included in that importation or exportation were goods described as forfeited to the Crown under section 229 of that Act because they were:
   (c) prohibited imports within the meaning of that Act; or
   (d) prohibited exports within the meaning of that Act; as the case requires.
Division 2—Failure to apply conformity assessment procedures

41ME  Failure to apply conformity assessment procedures—manufacturers

(1) A person is guilty of an offence if:
   (a) the person supplies in Australia a medical device that the person has manufactured; and
   (b) the conformity assessment procedures have not been applied to the device.

   Maximum penalty:  Imprisonment for 12 months or 1,000 penalty units, or both.

(2) A person is guilty of an offence if:
   (a) the person exports from Australia a medical device that the person has manufactured; and
   (b) the conformity assessment procedures have not been applied to the device.

   Maximum penalty:  Imprisonment for 12 months or 1,000 penalty units, or both.

41MF  Failure to apply conformity assessment procedures—sponsors

(1) A person is guilty of an offence if:
   (a) the person supplies a medical device in Australia; and
   (b) the conformity assessment procedures have not been applied to the device.

   Maximum penalty:  Imprisonment for 12 months or 1,000 penalty units, or both.

(2) A person is guilty of an offence if:
   (a) the person exports a medical device from Australia; and
   (b) the conformity assessment procedures have not been applied to the device.
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Maximum penalty:  Imprisonment for 12 months or 1,000 penalty units, or both.

(3) This section does not apply if the defendant was not the sponsor of the device at the time of the supply or export, as the case may be.

Note:  A defendant bears an evidential burden in relation to the matters in subsection (3) (see subsection 13.3(3) of the Criminal Code).

41MG Exceptions

(1) Sections 41ME and 41MF do not apply to the extent that:
   (a) the quality management systems applied to the medical device comply with one or more conformity assessment standards that apply to them; and
   (b) the conformity assessment procedures have not been applied to the device only in respect of a part or parts of the conformity assessment procedures to which one or more of those conformity assessment standards relate.

Note:  Conformity assessment standards are determined under Division 2 of Part 4-3.

(2) For the purposes of this section, a conformity assessment standard relates to a part or parts of the conformity assessment procedures only if the standard specifies that part or parts.

Note:  A defendant bears an evidential burden in relation to the matters in this section (see subsection 13.3(3) of the Criminal Code).

41MH False statements in declarations

A person is guilty of an offence if:
   (a) the person makes a statement in or in connection with a declaration, relating to the application of conformity assessment procedures to a medical device that the person has manufactured; and
   (b) the statement is false or misleading in a material particular.

Maximum penalty:  Imprisonment for 12 months or 1,000 penalty units, or both.
Division 3—Medical devices not included in the Register and related matters

41MI  Importation, exportation, supply or manufacture of medical devices not included in the Register

(1) A person is guilty of an offence if:
   (a) the person:
      (i) imports a medical device into Australia; or
      (ii) exports a medical device from Australia; or
      (iii) supplies a medical device in Australia; or
      (iv) manufactures a medical device in Australia; and
   (b) none of the following subparagraphs applies in relation to the device:
      (i) the device is of a kind included in the Register in relation to the person;
      (ii) the device is an exempt device;
      (iii) the device is the subject of an approval under section 41HB or an authority under section 41HC.

   Maximum penalty: Imprisonment for 12 months or 1,000 penalty units, or both.

(2) Strict liability applies to paragraph (1)(b).
   Note: For strict liability, see section 6.1 of the Criminal Code.

(3) Subsection (1) does not apply if the defendant proves that the defendant was not the sponsor of the device at the time of the importation, export, supply, or manufacture, as the case may be.
   Note: A defendant bears a legal burden in relation to the matters in subsection (3) (see section 13.4 of the Criminal Code).

41MJ  Treating medical devices as prohibited imports or exports

If:
   (a) the importation or exportation of a medical device is prohibited under subsection 41MI(1); and
Section 41MK

(b) the Secretary notifies the Chief Executive Officer of Customs in writing that the Secretary wishes the *Customs Act 1901* to apply to that importation or exportation; the *Customs Act 1901* has effect as if the device included in that importation or exportation were goods described as forfeited to the Crown under section 229 of that Act because they were:

(c) prohibited imports within the meaning of that Act; or

(d) prohibited exports within the meaning of that Act; as the case requires.

41MK Wholesale supply of medical devices not included in the Register

A person is guilty of an offence if:

(a) the person supplies a medical device in Australia; and

(b) none of the following subparagraphs applies in relation to the device:

(i) the device is of a kind included in the Register;

(ii) the device is an exempt device;

(iii) the device is the subject of an approval under section 41HB or an authority under section 41HC; and

(c) the person to whom the device is supplied is not the ultimate consumer of the device.

Maximum penalty: 120 penalty units.

41ML Misrepresentations etc. about medical devices

(1) A person is guilty of an offence if:

(a) the person makes a representation of a kind referred to in subsection (2); and

(b) the representation is false or misleading.

Maximum penalty: 60 penalty units.

(2) Subsection (1) applies to these representations:

(a) representations that medical devices are of a kind included in the Register;

(b) representations that medical devices are exempt devices;
Section 41MM

(c) representations that medical devices are the subject of an approval under section 41HB or an authority under section 41HC.

(3) A person is guilty of an offence if:
   (a) a kind of medical device is included in the Register in respect of a person; and
   (b) the person, by any means, advertises the goods as being for a purpose other than that accepted in relation to that inclusion.

Maximum penalty: 60 penalty units.

41MM  Claims about arranging supplies of medical devices not included in the Register

A person is guilty of an offence if:
   (a) the person claims, by any means, that the person or another person can arrange the supply of medical devices; and
   (b) the devices are not:
       (i) medical devices of a kind included in the Register; or
       (ii) exempt devices.

Maximum penalty: 60 penalty units.

41MN  Breaches of conditions

(1) A person is guilty of an offence if:
   (a) a kind of medical device is included in the Register in relation to the person; and
   (b) the person does an act, or omits to do an act; and
   (c) the act or omission constitutes a breach of a condition of the inclusion of the kind of device in the Register.

Maximum penalty: 60 penalty units.

(2) A person is guilty of an offence if:
   (a) a conformity assessment certificate is issued in respect of a person; and
   (b) the person does an act, or omits to do an act; and
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(c) the act or omission constitutes a breach of a condition of the conformity assessment certificate.

Maximum penalty: 60 penalty units.

(3) A person is guilty of an offence if:

(a) the person does an act, or omits to do an act; and

(b) the act or omission constitutes a breach of:

(i) a condition of an exemption applicable under regulations made for the purposes of section 41HA; or

(ii) a condition of an approval under section 41HB; or

(iii) a condition applicable under regulations made for the purposes of subsection 41HB(7).

Maximum penalty: 60 penalty units.
Division 4—Other offences

41MO Misuse of medical devices exempted for special or experimental uses

(1) A person is guilty of an offence if he or she:
   (a) has been granted an authority under section 41HC relating to a specified kind of medical device; and
   (b) supplies a medical device of that kind:
       (i) other than in accordance with the authority; or
       (ii) other than in accordance with any conditions to which the authority is subject; or
       (iii) other than in accordance with any regulations made for the purpose of subsection 41HC(5).

   Maximum penalty: 60 penalty units.

(2) A person is guilty of an offence if:
   (a) he or she has been granted an approval under section 41HB relating to a specified medical device or specified kind of medical device; and
   (b) he or she uses a medical device of that kind:
       (i) in the treatment of another person; or
       (ii) solely for experimental purposes in humans;

   in a way that is not in accordance with the approval.

   Maximum penalty: 60 penalty units.

41MP Notification of adverse events etc.

(1) A person is guilty of an offence if:
   (a) the person is a person in relation to whom a kind of medical device is included in the Register; and
   (b) the person knows that particular information is information of a kind mentioned in subsection (2); and
   (c) the person fails to give that information to the Secretary within the period specified in the regulations (whether or not
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the person has already given to the Secretary other
information relating to the same matter).

Maximum penalty: 400 penalty units.

(2) The information with which subsection (1) is concerned is
information of the following kinds:

(a) information relating to:
   (i) any malfunction or deterioration in the characteristics or
       performance of a kind of device; or
   (ii) any inadequacy in the design, production, labelling,
       instructions for use or advertising materials of a kind of
       device; or
   (iii) any use in accordance with, or contrary to, the use
       intended by the manufacturer of the kind of device;
       that might lead, or might have led, to the death of a patient or
       a user of the device, or to a serious deterioration in his or her
       state of health;

(b) information relating to any technical or medical reason for a
malfunction or deterioration of a kind referred to in
subparagraph (a)(i) that has led the manufacturer to take
steps to recover devices of that kind that have been
distributed;

(c) information that indicates that a device of that kind does not
comply with the essential principles;

(d) information that indicates that a certificate (other than one
issued under this Act) used for the purpose of an application
under subsection 41FC(1) to signify:
   (i) compliance with the essential principles; or
   (ii) the application of relevant conformity assessment
       procedures to a particular device;
       has been restricted, suspended, revoked or is no longer in
effect.

41MQ  Notification of adverse events etc. where application
withdrawn or lapses

(1) If an application for inclusion of a kind of medical device in the
Register is withdrawn or lapses, the Secretary may give the
applicant written notice requiring the applicant:

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(a) to inform the Secretary in writing whether the applicant is aware of any information of a kind mentioned in subsection 41MP(2) relating to the kind of device; and
(b) if the applicant is aware of such information, to give the information to the Secretary in writing.

(2) Notice under subsection (1) may only be given within 10 working days after an application is withdrawn or lapses.

(3) A person is guilty of an offence if the person fails to comply with the requirements of a notice under subsection (1) within 20 working days after the notice is given to the person.

Maximum penalty: 400 penalty units.

(4) A person is guilty of an offence if:
   (a) the person gives information in purported compliance with a notice under this section; and
   (b) the information is false or misleading in a material particular.

Maximum penalty: 400 penalty units.
Chapter 5—Advertising, counterfeit therapeutic goods and product tampering

Part 5-1—Advertising and generic information

Division 1—Preliminary

42AA This Part not to apply to advertisements directed at health professionals etc.

(1) This Part does not apply to advertisements directed exclusively to:

(a) medical practitioners, psychologists, dentists, veterinary surgeons, pharmacists, physiotherapists, dietitians, scientists working in medical laboratories or nurses; or

(b) persons who are:

(i) engaged in the business of wholesaling therapeutic goods; or

(ii) purchasing officers in hospitals; or

(c) herbalists, homoeopathic practitioners, chiropractors, naturopaths, nutritionists, practitioners of traditional Chinese medicine, podiatrists or osteopaths registered under a law of a State or Territory.

(2) This Part does not apply to advertisements directed exclusively to persons who are members of an Australian branch (however described) of one of the bodies prescribed for the purposes of this subsection.

(3) For the purposes of subsection (2), a person is taken to be a member of an Australian branch of one of those bodies if, and only if, the person has the qualifications and training that are necessary or appropriate for membership of the relevant body.

(4) This Part does not apply to advice or information given directly to a patient by a person referred to in paragraph (1)(a) or (c) or subsection (2) in the course of treatment of that patient.
42AB  This Part not to apply to advertisements for goods not for human use

This Part does not apply to advertisements in respect of goods that are not for use in humans.

42AC  This Part not to apply to advertisements for exported goods

(1) Subject to subsection (2), this Part does not apply to advertisements solely for therapeutic goods that have been exported or are intended exclusively for export.

(2) Section 42DC applies to advertisements of that kind.

42B  Definitions

In this Part, unless the contrary intention appears:

approval number means the distinguishing number allocated to an approved advertisement by the Secretary under regulation 5J of the Therapeutic Goods Regulations.

approved advertisement means an advertisement:

(a) approved under regulation 5G, or taken to be approved by the Secretary under subregulation 5H(2), or approved by the Minister on review under regulation 5M, of the Therapeutic Goods Regulations; and

(b) the approval of which has not been withdrawn.

broadcaster, in relation to an advertisement for therapeutic goods, means a person (other than a person who is required to enter those goods on the Register) who undertakes, as a business activity in its own right:

(a) the broadcasting of the advertisement in broadcast media; or

(b) the placement of the advertisement for such broadcasting.

broadcast media, in relation to an advertisement or generic information, means any means (other than a means declared in the regulations to be an exempted means) by which the information is disseminated electronically in a visible or audible form or a combination of such forms.
generic information, in relation to therapeutic goods, includes any statement, pictorial representation or design, however made, about the composition, properties or other characteristics of therapeutic goods, but does not include:

(a) an advertisement about the goods; or  
(b) generic information included in an advertisement about the goods; or  
(c) bona fide news.

mainstream media means any magazine or newspaper for consumers containing a range of news, public interest items, advertorials, advertisements or competitions.

prohibited representation means a representation referred to in subsection 42DJ(1).

publisher, in relation to an advertisement for therapeutic goods, means a person (other than a person who is required to enter those goods on the Register) who undertakes, as a business activity in its own right:

(a) the publishing of the advertisement in specified media other than broadcast media; or  
(b) the placement of the advertisement for such publication.

publishing, in relation to an advertisement, includes inserting material within the pages of an item of mainstream media.

required representation means a representation referred to in subsection 42DJ(2).

restricted representation means a representation referred to in subsection 42DD(1).

specified media, in relation to an advertisement or generic information, means:

(a) mainstream media; or  
(b) broadcast media; or  
(c) cinematograph films; or  
(d) displays about goods, including posters:  
   (i) in shopping malls (except inside an individual shop); and
(ii) in or on public transport; and
(iii) on billboards.

*visual broadcast media* means broadcast media that is intended to be viewed by its audience.
**Section 42BA**

**Division 2—Therapeutic goods advertisements for which an approval is required**

**42BA  Application of Division**

This Division applies only to advertisements to which Part 2 of the Therapeutic Goods Regulations applies.

**42C  Offences relating to publication of advertisements**

(1) A person is guilty of an offence if:

(a) the person:

(i) publishes or broadcasts; or

(ii) causes to be published or broadcast;

in specified media, an advertisement that is required by the Therapeutic Goods Regulations to be an approved advertisement; and

(b) the advertisement is not an approved advertisement.

Penalty: 60 penalty units.

(2) A person is guilty of an offence if:

(a) the person:

(i) publishes or broadcasts; or

(ii) causes to be published or broadcast;

an advertisement in specified media; and

(b) the advertisement is not an approved advertisement in that it differs, in any respect, from the advertisement that was approved.

Penalty: 60 penalty units.

(3) It is a defence to a prosecution under subsection (2) if:

(a) the person prosecuted is a publisher or broadcaster who received the advertisement to which the prosecution relates for publication or broadcasting in specified media in the ordinary course of business; or
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Section 42C

(b) the particular advertisement to which the prosecution relates differs only in respect of a matter mentioned in paragraph 5C(2)(b), (e) or (f) of the Therapeutic Goods Regulations.

Note: A defendant bears an evidential burden in relation to the matters in subsection (3) (see subsection 13.3 of the Criminal Code).

(4) A person is guilty of an offence if:
   (a) the person:
      (i) publishes or broadcasts; or
      (ii) causes to be published or broadcast;
   in specified media referred to in paragraph (a), (c) or (d) of the definition of specified media, or in visual broadcast media, a particular advertisement; and
   (b) the advertisement:
      (i) does not display its approval number; or
      (ii) displays a number purporting to be its approval number but that is not its approval number; or
      (iii) displays an approval number that has expired.

Penalty: 30 penalty units.

(5) It is a defence to a prosecution under subsection (4) if the person prosecuted:
   (a) is a publisher who received the advertisement to which the prosecution relates for publication in specified media referred to in paragraph (a), (c) or (d) of the definition of specified media; or
   (b) is a broadcaster who received the advertisement to which the prosecution relates for broadcasting in visual broadcast media;
   in the ordinary course of business.

Note: A defendant bears an evidential burden in relation to the matters in subsection (5) (see subsection 13.3 of the Criminal Code).

(6) A person is guilty of an offence if:
   (a) the person:
      (i) publishes or broadcasts; or
      (ii) causes to be published or broadcast;
   in specified media, an approved advertisement; and
(b) the person’s action is in contravention of a condition to which the approval of the advertisement is subject.

Penalty: 60 penalty units.

(7) It is a defence to a prosecution under subsection (6) if the person prosecuted is a publisher or broadcaster who received the advertisement to which the prosecution relates for publication or broadcasting in specified media in the ordinary course of business.

Note: A defendant bears an evidential burden in relation to the matters in subsection (7) (see subsection 13.3 of the Criminal Code).

(8) An offence against this section is an offence of strict liability.
Division 3—General provisions about advertising therapeutic goods

42DA  Application of Division

This Division applies to advertisements about therapeutic goods other than advertisements for which an approval is required under Part 2 of the Therapeutic Goods Regulations.

42DB  Definitions

In this Division:

applicant means an applicant for approval of the use of a restricted representation in an advertisement about therapeutic goods.

approval holder, in relation to a restricted representation, means the person to whom notice of approval of the use of the restricted representation was given.

42DC  Certain representations not to be published or broadcast

If a representation in an advertisement about therapeutic goods is false or misleading, the Secretary may, by notice given to the person apparently responsible for publishing or broadcasting the advertisement, prevent that person from publishing or broadcasting, or causing to be published or broadcast, an advertisement containing that representation (whether express or implied) about those goods.

42DD  Restricted representations

(1) For the purposes of this Part, a representation in an advertisement about therapeutic goods that refers to a serious form of a disease, condition, ailment or defect specified in a part of the Therapeutic Goods Advertising Code that is prescribed by the regulations for the purposes of this subsection is a restricted representation about therapeutic goods.
Section 42DE

(2) A person must not use a restricted representation in an advertisement about therapeutic goods unless the Secretary:
   (a) has approved its use under subsection 42DF(1); or
   (b) has permitted its use under subsection 42DK(1).

42DE Applications for approval of use of restricted representation

An application for approval of the use of a restricted representation must be:
   (a) made to the Secretary in writing, in a form approved by the Secretary; and
   (b) signed by or on behalf of the applicant.

42DF Approval of use of restricted representation

(1) If an application for approval of the use of a restricted representation is made, the Secretary must approve the use of the restricted representation if the Secretary is satisfied that:
   (a) the representation is accurate and balanced; and
   (b) the representation is not misleading or likely to be misleading.

(2) Otherwise, the Secretary must refuse to approve the use of the restricted representation.

(3) An approval may be subject to conditions imposed by the Secretary.

(4) In deciding whether to approve or refuse to approve the use of a restricted representation, the Secretary must take into consideration:
   (a) any recommendation of the Therapeutic Goods Advertising Code Council; and
   (b) any advice of the Complementary Medicines Evaluation Committee or the Medicines Evaluation Committee; and
   (c) the public interest criteria mentioned in a part of the Therapeutic Goods Advertising Code that is prescribed by the regulations made for the purposes of this paragraph.
42DG  Notice of approval or refusal

(1) The Secretary must give written notice to the applicant of the approval of, or of the refusal to approve, the use of a restricted representation.

(2) If written notice is not given to the applicant within the period of 60 days after the day on which the application was made (or within such longer period as the Secretary specifies by written notice to the applicant before the end of that period), the Secretary is taken to have approved the use of the restricted representation at the end of the period.

(3) If an approval is subject to conditions, the conditions must be set out in the notice.

(4) A notice of refusal to approve the use of a restricted representation must:
   (a) give the Secretary’s reasons for the refusal; and
   (b) inform the applicant of the applicant’s right to have the Secretary’s decision reviewed by the Minister under section 60.

42DH  Variation of conditions of approval

(1) The Secretary, by written notice to an approval holder, may vary any condition of approval of the use of a restricted representation.

(2) The notice must:
   (a) give the Secretary’s reasons for the variation; and
   (b) inform the approval holder of the approval holder’s right to have the Secretary’s decision reviewed by the Minister under section 60.

42DI  Withdrawal of approval

(1) The Secretary, by written notice, may withdraw the approval of the use of a restricted representation if:
   (a) the Secretary is satisfied that:
      (i) information given by the applicant in the application was false or incorrect and the Secretary, or the Minister
on review of a decision of the Secretary under section 42DF or 42DH, relied on the information in deciding to approve the use of the representation; or
(ii) the restricted representation has become a prohibited representation; or
(iii) there has been a breach of a condition of approval; or
(b) both:
(i) additional information about the safety of the therapeutic goods becomes available; and
(ii) the Secretary is satisfied that, if that information had been available at the time of the approval, the Secretary would not have approved the use of the restricted representation.

(2) The notice must:
(a) give the Secretary’s reasons for the withdrawal; and
(b) inform the approval holder of the approval holder’s right to have the Secretary’s decision reviewed by the Minister under section 60.

**42DJ Prohibited and required representations**

(1) For the purposes of this Part, representations of a kind specified in regulations made for the purposes of this subsection are prohibited representations about therapeutic goods of a kind specified in those regulations.

(2) For the purposes of this Part, representations of a kind specified in regulations made for the purposes of this subsection are required representations about the therapeutic goods of a kind specified in those regulations.

**42DK Use of restricted or prohibited representations**

(1) The Secretary may, by notice in writing published in the Gazette or on the Department’s web site on the Internet, permit, in relation to therapeutic goods, the use of a restricted representation (including its use on the label of the goods or in information included in the package in which the goods are contained).
(2) The Secretary may, by notice in writing published in the Gazette or on the Department’s web site on the Internet, permit a prohibited representation to be included on the label of therapeutic goods, or in information included in the package in which therapeutic goods are contained, if the representation is necessary for the appropriate use of the goods.

42DL Advertising offences

(1) A person must not publish or broadcast an advertisement about therapeutic goods:
   (a) that contains a prohibited representation (whether in express terms or by necessary implication) about those goods; or
   (b) that does not contain a required representation about those goods; or
   (c) that contains a restricted representation, about those goods, the use of which has not been approved under subsection 42DF(1) or permitted under subsection 42DK(1); or
   (d) that is in contravention:
      (i) of a notice referred to in section 42DC that was served on the person; or
      (ii) of a notice referred to in section 42DK of which the person was aware when the advertisement was published; or
   (e) that contains:
      (i) a reference to the Act other than in a statement of the registration number, listing number or device number of the goods; or
      (ii) a statement suggesting or implying the goods have been recommended or approved by or on behalf of a government or government authority (including a foreign government or foreign government authority), other than a statement of their availability as a pharmaceutical benefit or a statement authorised or required by a government or government authority (including a foreign government or foreign government authority); or
(f) that refers to goods, or substances or preparations containing goods, included in Schedule 3, 4 or 8 to the Poisons Standard; or

(g) that are not entered in the Register; or

(h) if the goods are therapeutic goods, or come within a class of therapeutic goods, that:
   (i) are exempt goods or exempt devices prescribed in the regulations for the purposes of this provision; or
   (ii) have been approved under subsection 19(1) or section 41HB of this Act for importation into, exportation from, or supply within, Australia.

Penalty: 60 penalty units.

(2) For the purposes of an offence against subsection (1), strict liability applies to the following physical elements:

(a) that the use of a restricted representation, as referred to in paragraph (1)(c), has not been approved under subsection 42DF(1) or permitted under subsection 42DK(1);

(b) that the notice referred to in paragraph (1)(d):
   (i) in a case to which subparagraph (1)(d)(i) applies—is a notice referred to in section 42DC; and
   (ii) in a case to which subparagraph (1)(d)(ii) applies—is a notice referred to in section 42DK;

(c) that goods, substances or preparations referred to in paragraph (1)(f) are included in Schedule 3, 4 or 8 to the Poisons Standard;

(d) that the therapeutic goods, or class of therapeutic goods, referred to in paragraph (1)(h):
   (i) are exempt goods or exempt devices prescribed in the regulations made for the purposes of subparagraph (1)(h)(i); or
   (ii) have been approved under subsection 19(1) or section 41HB of the Act for importation into, exportation from or supply within, Australia.

(3) It is a defence to a prosecution under subsection (1) if:

(a) in relation to an advertisement mentioned in paragraph (1)(a) or (f)—the advertisement is made by, or on behalf of, the Commonwealth; and
Section 42DM

42DM Compliance with Code

(1) A person is guilty of an offence if:
   (a) the person publishes or broadcasts an advertisement about therapeutic goods; and
   (b) the advertisement does not comply with the Therapeutic Goods Advertising Code.

Penalty: 60 penalty units.

(2) An offence against this section is an offence of strict liability.
Division 4—Generic information about ingredients or components of therapeutic goods

42DN Application of Division

This Division applies to generic information about goods that:
(a) may be used as an ingredient or component in the manufacture of therapeutic goods; and
(b) although not presented for supply as therapeutic goods, come within the meaning of therapeutic goods because they are represented to be:
   (i) for therapeutic use; or
   (ii) for use as an ingredient or component in the manufacture of other therapeutic goods.

42DO Compliance with the Code

Generic information to which this Division applies must comply with principles of the Therapeutic Goods Advertising Code specified in regulations made for the purposes of this section as if those principles applied to generic information in the same way as they apply to advertisements.

42DP Offences—publication of generic information

(1) A person is guilty of an offence if:
   (a) the person publishes or broadcasts generic information about therapeutic goods; and
   (b) the publication or broadcasting of that generic information does not comply with principles contained in the part of the Therapeutic Goods Advertising Code that are specified in Regulations.

Penalty: 60 penalty units.

(2) An offence against this section is an offence of strict liability.
Part 5-2—Counterfeit therapeutic goods

42E Offence of dealing with counterfeit therapeutic goods

(1) A person is guilty of an offence if:
   (a) the person intentionally:
       (i) manufactures goods in Australia; or
       (ii) supplies goods in Australia; or
       (iii) imports goods into Australia; or
       (iv) exports goods from Australia; and
   (b) the goods are therapeutic goods; and
   (c) the goods are counterfeit and the person knows that fact or is reckless as to whether that fact exists.

(2) Goods are counterfeit if any of the following contain a false representation of a matter listed in subsection (3):
   (a) the label or presentation of the goods;
   (b) any document or record relating to the goods or their manufacture;
   (c) any advertisement for the goods.

(3) The matters are as follows:
   (a) the identity or name of the goods;
   (b) the formulation, composition or design specification of the goods or of any ingredient or component of them;
   (c) the presence or absence of any ingredient or component of the goods;
   (d) the strength or size of the goods (other than the size of any pack in which the goods are contained);
   (e) the strength or size of any ingredient or component of the goods;
   (f) the sponsor, source, manufacturer or place of manufacture of the goods.

(4) An offence against this section is punishable on conviction by imprisonment for not more than 5 years, a fine not more than 2,000 penalty units or both.
Section 42F

Note: Subsection 4B(3) of the Crimes Act 1914 lets a court fine a body corporate up to 5 times the maximum amount the court can fine an individual.

(5) To avoid doubt, a term that is defined in subsection 3(1) in relation to therapeutic goods and used in this section in relation to goods has in this section the meaning given by subsection 3(1).

42F Customs treatment of counterfeit therapeutic goods

Imported counterfeit therapeutic goods

(1) If the Secretary notifies the Chief Executive Officer of Customs in writing that the Secretary wishes the Customs Act 1901 to apply to an import of counterfeit therapeutic goods, that Act has effect as if the goods included in the import were goods described as forfeited to the Crown under section 229 of that Act because they were prohibited imports within the meaning of that Act.

Exported counterfeit therapeutic goods

(2) If the Secretary notifies the Chief Executive Officer of Customs in writing that the Secretary wishes the Customs Act 1901 to apply to an export of counterfeit therapeutic goods, that Act has effect as if the goods included in the export were goods described as forfeited to the Crown under section 229 of that Act because they were prohibited exports within the meaning of that Act.
Part 5-3—Product tampering

42T Notifying of actual or potential tampering

(1) A person is guilty of an offence if:
   (a) the person supplies, manufactures or is a sponsor of, or proposes to supply, manufacture or become a sponsor of, therapeutic goods; and
   (b) either:
      (i) the person knows that some or all of those therapeutic goods, or any other therapeutic goods, are or have been subject to actual or potential tampering; or
      (ii) some or all of those therapeutic goods, or any other therapeutic goods, are or have been subject to actual or potential tampering, and the person is reckless as to that fact; and
   (c) the person fails, within 24 hours after becoming aware of, or becoming aware of a substantial risk of, the actual or potential tampering, to notify the Secretary or the National Manager of the Therapeutic Goods Administration.

Maximum penalty: 400 penalty units.

(2) A person is guilty of an offence if:

   (a) the person supplies, manufactures or is a sponsor of, or proposes to supply, manufacture or become a sponsor of, therapeutic goods; and

   (b) the person receives information or a demand; and

   (c) either:
      (i) the person knows that the information or demand relates (either expressly or by implication) to actual or potential tampering with some or all of those therapeutic goods, or any other therapeutic goods; or
      (ii) the information or demand relates (either expressly or by implication) to actual or potential tampering with some or all of those therapeutic goods, or any other
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Section 42U

therapeutic goods, and the person is negligent as to that fact; and
(d) the person fails to notify the Secretary or the National Manager of the Therapeutic Goods Administration of the information or demand within 24 hours after receiving it.

Maximum penalty: 240 penalty units.

(3) For the purposes of subparagraph (2)(c)(ii), the person is only taken to be negligent as to the fact that the information or demand is of the kind referred to in that subparagraph if:
(a) the person’s acts or omissions involve such a great falling short of the standard of care that a reasonable person would exercise in the circumstances; and
(b) there is such a high risk that the information or demand is of that kind;
that the acts or omissions merit criminal punishment.

(4) For the purposes of this section, it does not matter whether, at the time of receipt of the information or demand:
(a) the person has possession or control of the therapeutic goods to which the information or demand relates; or
(b) the therapeutic goods are in existence.

42U Meaning of actual or potential tampering etc.

Actual or potential tampering, in relation to therapeutic goods, means:
(a) tampering with the therapeutic goods; or
(b) causing the therapeutic goods to be tampered with; or
(c) proposing to tamper with the therapeutic goods; or
(d) proposing to cause the therapeutic goods to be tampered with.

42V Recovery of therapeutic goods because of actual or potential tampering

(1) The Secretary may, in writing, impose requirements under this section on a person if:
Section 42V

(a) the person supplies or has supplied therapeutic goods of a particular kind, or a particular batch of therapeutic goods of that kind; and

(b) the Secretary is satisfied that therapeutic goods of that kind, or included in that batch, are, have been or could possibly be, subject to actual or potential tampering.

(2) The requirements may be one or more of the following:

(a) to take specified steps, in the specified manner and within such reasonable period as is specified, to recover therapeutic goods of that kind, or included in that batch, that the person has supplied;

(b) to inform the public or a specified class of persons, in the specified manner and within such reasonable period as is specified, that therapeutic goods of that kind, or included in that batch, are, or have been, subject to actual or potential tampering;

(c) to inform the public or a specified class of persons, in the specified manner and within such reasonable period as is specified, that therapeutic goods of that kind, or included in that batch, could possibly be subject to actual or potential tampering.

(3) Requirements referred to in paragraph (2)(a) do not apply to therapeutic goods that cannot be recovered because they have been administered to, or applied in the treatment of, a person or animal.

(4) The Secretary must cause to be published in the Gazette, as soon as practicable after imposing such requirements, a notice setting out particulars of the requirements.

(5) The Secretary may impose requirements under this section whether or not the Secretary has been notified under section 42T.

(6) A person who intentionally refuses or fails to comply with a requirement under subsection (1) is guilty of an offence.

   Maximum penalty: 240 penalty units.

(7) This section does not prevent the Secretary from taking action under section 30 or Division 1 or 2 of Part 4-6.
Section 42W

42W Supply etc. of therapeutic goods that are subject to recovery requirements

(1) A person is guilty of an offence if:
   (a) the person supplies therapeutic goods in Australia; and
   (b) either:
      (i) the person knows that the therapeutic goods are of a kind, or are included in a batch, in respect of which requirements have been imposed under section 42V, on that person or another person, to recover therapeutic goods; or
      (ii) the therapeutic goods are of such a kind, or are included in such a batch, and the person is reckless as to that fact; and
   (c) the Secretary has not consented in writing to the supply.

Maximum penalty: 240 penalty units.

(2) A person is guilty of an offence if:
   (a) the person exports therapeutic goods from Australia; and
   (b) either:
      (i) the person knows that the therapeutic goods are of a kind, or are included in a batch, in respect of which requirements have been imposed under section 42V, on that person or another person, to recover therapeutic goods; or
      (ii) the therapeutic goods are of such a kind, or are included in such a batch, and the person is reckless as to that fact; and
   (c) the Secretary has not consented in writing to the exportation.

Maximum penalty: 240 penalty units.

(3) The Secretary must not give consent relating to an exportation unless satisfied that there are exceptional circumstances that justify giving the consent.
42X Saving of other laws

This Part is not intended to exclude or limit the operation of any other law of the Commonwealth or any law of a State or Territory.
Chapter 6—Administration

Part 6-1—Payment of charges

43 By whom charges payable

(1) An annual registration charge, annual listing charge or annual charge for inclusion in the Register is payable by the person in relation to whom the therapeutic goods concerned are registered, listed or included in the Register.

(2) An annual licensing charge is payable by the holder of the licence to which the charge relates.

44 Time for payment of charges

(1) An annual registration charge, annual listing charge or annual charge for inclusion in the Register for a financial year that relates to therapeutic goods other than grouped therapeutic goods becomes payable:

   (a) if the charge is imposed in respect of therapeutic goods the registration or listing of which commenced before 1 July 1990:

      (i) in the case of charge for the financial year commencing on 1 July 1990—on that day; or

      (ii) in the case of a later financial year—on 1 July in that financial year or, if the Secretary has, by notice in writing given before 1 July 1991 to the person in relation to whom the therapeutic goods concerned are registered or listed, specified another day as being the day on which the charge becomes payable, on the specified day or on an anniversary of that day, as the case requires; or

   (b) in any other case:

      (i) if the year is the financial year (in this paragraph called the first year) during which the registration, listing or inclusion in the Register of the therapeutic goods concerned commenced—on that commencement; or
(ii) if the year is a later financial year—on the anniversary of that commencement or, if the Secretary has, by notice in writing given before the end of the first year to the person in relation to whom the therapeutic goods concerned are registered, listed or included in the Register, specified another day as being the day on which charge becomes payable, on the specified day or on an anniversary of that day, as the case requires.

(1A) An annual registration charge or annual listing charge for a financial year that relates to grouped therapeutic goods becomes payable by a person on the day specified in relation to those grouped therapeutic goods in a written notice given by the Secretary to the person.

(2) An annual licensing charge for a financial year becomes payable:
   (a) if the licence commenced before 1 July 1990—on 1 July 1990 and on each anniversary of that day; or
   (b) in any other case—on the day on which the licence commenced and on each anniversary of that day.

(3) The Secretary may, by agreement with the person by whom an annual registration charge, an annual listing charge, an annual charge for inclusion in the Register or an annual licensing charge is payable, vary the day on which the charge becomes payable in a financial year.

45 Therapeutic Goods Administration Account

(1) There is continued in existence the Therapeutic Goods Administration Account.

Note: The Account was established by subsection 5(3) of the Financial Management Legislation Amendment Act 1999.

(2) The Account is a Special Account for the purposes of the Financial Management and Accountability Act 1997.

(3) There must be credited to the Account amounts equal to:
   (a) amounts received by the Commonwealth by way of annual registration charge, annual listing charge, annual charge for inclusion in the Register and annual licensing charge; and
Section 45

(b) interest received by the Commonwealth from the investment of an amount standing to the credit of the Account; and

c) money received by the Commonwealth in relation to property paid for after a debit from the Account; and

d) money received by the Commonwealth for services provided or to be provided, by or on behalf of the Commonwealth, using amounts standing to the credit of the Account (including amounts received by way of fees payable under the regulations); and

e) donations for the furtherance of a purpose of the Account that are received by the Commonwealth; and

(f) receipts relating to the recovery of debts (other than debts in respect of statutory fines and penalties) by the Commonwealth that are associated with expenditure of an amount standing to the credit of the Account.

Note: An Appropriation Act provides for amounts to be credited to a Special Account if any of the purposes of the Account is a purpose that is covered by an item in the Appropriation Act.

(4) The purposes of the Account are to make payments:

(a) to further the objects of this Act (as set out in section 4); and

(b) to enable the Commonwealth to participate in the international harmonisation of regulatory controls on therapeutic goods and other related activities.
Part 6-2—Entry, searches and warrants

45A Definitions

In this Part, unless the contrary intention appears:

**evidential material** means:
(a) any thing with respect to which an offence against this Act has been committed or is suspected, on reasonable grounds, to have been committed; or
(b) any thing as to which there are reasonable grounds for suspecting that it will afford evidence as to the commission of any such offence; or
(c) any thing as to which there are reasonable grounds for suspecting that it is intended to be used for the purpose of committing any such offence.

**occupier**, in relation to premises, includes a person present at the premises who is in apparent control of the premises.

**seize** includes secure against interference.

**thing** includes a substance, and a thing in electronic or magnetic form.

46 Searches to monitor compliance with Act

(1) Subject to subsections (2) and (3), an authorised person may, for the purpose of finding out whether this Act or the regulations have been complied with:
(a) enter any premises; and
(b) exercise the powers set out in subsection 48(1).

(2) The authorised person must not enter the premises unless:
(a) the occupier of the premises has consented to the entry; or
(b) the entry is made under a warrant issued under section 49.

(3) An authorised person is not entitled to exercise any powers under subsection (1) in relation to premises if:
Section 46A

(a) the occupier of the premises has required the authorised person to produce his or her identity card for inspection by the occupier; and
(b) the authorised person fails to comply with the requirement.

46A Searches of certain premises to monitor compliance with Act

(1) An authorised person may, subject to subsections (2) and (3), and to the extent that it is reasonably necessary for the purpose of finding out whether this Act or the regulations have been complied with, enter premises to which this section applies and do any of the following:
   (a) search the premises and any thing on the premises;
   (b) inspect, examine, take measurements of, or conduct tests (including by the taking of samples) concerning, any thing on the premises that relates to therapeutic goods;
   (c) take photographs (including video recordings) or make sketches of the premises or any thing on the premises;
   (d) inspect any book, record or document on the premises.

(2) An authorised person must not, under subsection (1), enter premises that are a residence unless:
   (a) the occupier of the premises has consented to the entry; or
   (b) the premises are used for commercial purposes in relation to therapeutic goods, in addition to residential purposes.

(3) An authorised person is not entitled to exercise any powers under subsection (1) in relation to premises if:
   (a) the occupier of the premises has required the authorised person to produce his or her identity card for inspection by the occupier; and
   (b) the authorised person fails to comply with the requirement.

(4) This section applies to:
   (a) premises of a person:
      (ia) who is required to comply with a condition of an exemption of therapeutic goods under section 18A; or
      (i) who has been granted an approval or authority under section 19; or
      (ii) who has been granted an approval under section 19A; or
(ia) who has been granted an approval or authority under section 41HB or 41HC; or

(iii) in relation to whom therapeutic goods are registered, listed or included in the Register;

being premises connected with the importation, export, manufacture or supply of therapeutic goods, or the keeping of records relating to the importation, export, manufacture or supply of therapeutic goods; and

(b) premises to which the person in relation to whom therapeutic goods are registered, listed or included in the Register, or the sponsor of the goods, must allow access as a condition of the registration or listing of the therapeutic goods; and

(c) premises in relation to which a licence has been granted under Part 3-3 for, or a conformity assessment certificate issued under Part 4-4, in relation to the manufacture of therapeutic goods, or premises at which records are kept in relation to such manufacture.

46B  Searches and seizures on public health grounds

(1) Subject to subsection (2), if an authorised person has reasonable grounds for suspecting that:

(a) there may be on any premises a particular thing in respect of which this Act or the regulations have not been complied with; and

(b) it is necessary in the interests of public health to exercise powers under this section in order to avoid an imminent risk of death, serious illness or serious injury;

the authorised person may, to the extent that it is reasonably necessary for the purpose of avoiding an imminent risk of death, serious illness or serious injury, enter the premises and do any of the following:

(c) search the premises for the thing;

(d) if the authorised person finds the thing on the premises—seize it.

(2) An authorised person is not entitled to exercise any powers under subsection (1) in relation to premises if:
Section 47

(a) the occupier of the premises has required the authorised person to produce his or her identity card for inspection by the occupier; and
(b) the authorised person fails to comply with the requirement.

47 Searches and seizures related to offences

(1) Subject to subsections (2) and (3), if an authorised person has reasonable grounds for suspecting that there may be evidential material on any premises, the authorised person may:
(a) enter the premises; and
(b) exercise the powers set out in subsection (4) and subsection 48(1); and
(c) if the authorised person finds the thing on the premises—seize it.

(2) The authorised person must not enter the premises unless:
(a) the occupier of the premises has consented to the entry; or
(b) the entry is made under a warrant issued under section 50.

(3) An authorised person is not entitled to exercise any powers under subsection (1) in relation to premises if:
(a) the occupier of the premises has required the authorised person to produce his or her identity card for inspection by the occupier; and
(b) the authorised person fails to comply with the requirement.

(4) If:
(a) in the course of searching, in accordance with a warrant, for a particular thing, an authorised person finds another thing that the authorised person believes on reasonable grounds to be evidential material; and
(b) the authorised person believes, on reasonable grounds, that it is necessary to seize that other thing in order to prevent its concealment, loss or destruction, or its use in committing, continuing or repeating an offence against this Act; the warrant is taken to authorise the authorised person to seize that other thing.
48 General powers of authorised persons in relation to premises

(1) The powers an authorised person may exercise under paragraphs 46(1)(b) and 47(1)(b) are as follows:
   (a) to search the premises and any thing on the premises;
   (b) to inspect, examine, take measurements of, or conduct tests (including by the taking of samples) concerning, any thing on the premises that relates to therapeutic goods;
   (c) to take photographs (including video recordings) or make sketches of the premises or any thing on the premises;
   (d) if the authorised person was only authorised to enter the premises because the occupier of the premises consented to the entry—to require the occupier to:
      (i) answer any questions put by the authorised person; and
      (ii) produce any book, record or document requested by the authorised person;
   (e) if the authorised person was authorised to enter the premises by a warrant under section 49 or 50—to require any person in or on the premises to:
      (i) answer any questions put by the authorised person; and
      (ii) produce any book, record or document requested by the authorised person;
   (f) to inspect any book, record or document on the premises;
   (g) to take extracts from or make copies of any such book, record or document;
   (h) to take onto the premises such equipment and materials as the authorised person requires for the purpose of exercising powers in relation to the premises.

(3) A person must not refuse or fail to comply with a requirement under paragraph (1)(e).

Maximum penalty: 30 penalty units.

(3A) Subsection (3) does not apply if the person has a reasonable excuse.

Note: The defendant bears an evidential burden in relation to the matter in subsection (3A). See subsection 13.3(3) of the Criminal Code.
Section 48A

(4) It is a reasonable excuse for a person to refuse or fail to answer a question or produce a document if answering the question, or producing the document, would tend to incriminate the person.

48A Details of warrant to be given to occupier etc.

(1) If a warrant in relation to premises is being executed and the occupier of the premises or another person who apparently represents the occupier is present at the premises, the authorised person must make available to that person a copy of the warrant.

(2) The authorised person must identify himself or herself to that person.

(3) The copy of the warrant referred to in subsection (1) need not include the signature of the magistrate who issued the warrant.

48B Announcement before entry

(1) An authorised person must, before entering the premises under a warrant:

(a) announce that he or she is authorised to enter the premises; and

(b) give any person at the premises an opportunity to allow entry to the premises.

(2) An authorised person is not required to comply with subsection (1) if he or she believes on reasonable grounds that immediate entry to the premises is required to ensure:

(a) the safety of a person; or

(b) that the effective execution of the warrant is not frustrated.

48C Use of electronic equipment at premises

(1) The authorised person may operate electronic equipment at the premises to see whether evidential material is accessible by doing so if he or she believes on reasonable grounds that the operation of the equipment can be carried out without damage to the equipment.

(2) If the authorised person, after operating the equipment, finds that evidential material is accessible by doing so, he or she may:
Section 48C

(a) seize the equipment and any disk, tape or other associated device; or

(b) if the material can, by using facilities at the premises, be put in documentary form—operate the facilities to put the material in that form and seize the documents so produced; or

(c) if the material can be transferred to a disk, tape or other storage device that:
   i) is brought to the premises; or
   ii) is at the premises and the use of which for the purpose has been agreed to in writing by the occupier of the premises;

operate the equipment or other facilities to copy the material to the storage device and take the storage device from the premises.

(3) An authorised person may seize equipment under paragraph (2)(a) only if:

   (a) it is not practicable to put the material in documentary form as mentioned in paragraph (2)(b) or to copy the material as mentioned in paragraph (2)(c); or

   (b) possession by the occupier of the equipment could constitute an offence.

(4) If the authorised person believes on reasonable grounds that:

   (a) evidential material may be accessible by operating electronic equipment at the premises; and
   (b) expert assistance is required to operate the equipment; and
   (c) if he or she does not take action under this subsection, the material may be destroyed, altered or otherwise interfered with;

he or she may do whatever is necessary to secure the equipment, whether by locking it up, placing a guard or otherwise.

(5) The authorised person must give notice to the occupier of the premises of his or her intention to secure equipment and of the fact that the equipment may be secured for up to 24 hours.

(6) The equipment may be secured:

   (a) for a period not exceeding 24 hours; or
   (b) until the equipment has been operated by the expert; whichever happens first.
(7) If the authorised person believes on reasonable grounds that the expert assistance will not be available within 24 hours, he or she may apply to the magistrate for an extension of that period.

(8) The authorised person must give notice to the occupier of the premises of his or her intention to apply for an extension, and the occupier is entitled to be heard in relation to the application.

48D  Compensation for damage to electronic equipment

(1) If:
   (a) damage is caused to equipment as a result of it being operated as mentioned in section 48C; and
   (b) the damage was caused as a result of:
      (i) insufficient care being exercised in selecting the person who was to operate the equipment; or
      (ii) insufficient care being exercised by the person operating the equipment;

compensation for the damage is payable to the owner of the equipment.

(2) Compensation is payable out of money appropriated by the Parliament for the purpose.

(3) In determining the amount of compensation payable, regard is to be had to whether the occupier of the premises and his or her employees and agents, if they were available at the time, had provided any warning or guidance as to the operation of the equipment that was appropriate in the circumstances.

48E  Copies of seized things to be provided

(1) Subject to subsection (2), if an authorised person seizes, under a warrant relating to premises:
   (a) a document, film, computer file or other thing that can be readily copied; or
   (b) a storage device the information in which can be readily copied;

the authorised person must, if requested to do so by the occupier of the premises or another person who apparently represents the
occupier and who is present when the warrant is executed, give a copy of the thing or the information to that person as soon as practicable after the seizure.

(2) Subsection (1) does not apply if:
   (a) the thing that has been seized was seized under paragraph 48C(2)(b) or (c); or
   (b) possession by the occupier of the document, film, computer file, thing or information could constitute an offence.

48F Occupier entitled to be present during search

(1) If a warrant in relation to premises is being executed and the occupier of the premises or another person who apparently represents the occupier is present at the premises, the person is entitled to observe the search being conducted.

(2) The right to observe the search being conducted ceases if the person impedes the search.

(3) This section does not prevent 2 or more areas of the premises being searched at the same time.

48G Receipts for things seized under warrant

(1) If a thing is seized under this Part, the authorised person must provide a receipt for the thing.

(2) If 2 or more things are seized or moved, they may be covered in the one receipt.

48H Retention of seized things

(1) Subject to any contrary order of a court, if an authorised person seizes a thing under this Part, an authorised person must return it if:
   (a) the reason for its seizure no longer exists or it is decided that it is not to be used in evidence; or
   (b) the period of 90 days after its seizure ends;
whichever first occurs, unless the thing is forfeited or forfeitible to the Commonwealth.
Section 48J

(2) At the end of the 90 days specified in subsection (1), an authorised person must take reasonable steps to return the thing to the person from whom it was seized, unless:
   (a) proceedings in respect of which the thing may afford evidence were instituted before the end of the 90 days and have not been completed (including an appeal to a court in relation to those proceedings); or
   (b) an authorised person may retain the thing because of an order under section 48J; or
   (c) an authorised person is otherwise authorised (by a law, or an order of a court, of the Commonwealth or of a State or Territory) to retain, destroy or dispose of the thing.

(3) The thing may be returned under subsection (2) either unconditionally or on such terms and conditions as the Secretary sees fit.

48J Magistrate may permit a thing to be retained

(1) An authorised person may apply to a magistrate for an order that he or she may retain the thing for a further period if:
   (a) before the end of 90 days after the seizure; or
   (b) before the end of a period previously specified in an order of a magistrate under this section; proceedings in respect of which the thing may afford evidence have not commenced.

(2) If the magistrate is satisfied that it is necessary for an authorised person to continue to retain the thing:
   (a) for the purposes of an investigation as to whether an offence against this Act has been committed; or
   (b) to enable evidence of an offence against this Act to be secured for the purposes of a prosecution;
the magistrate may order that an authorised person may retain the thing for a period (not being a period exceeding 3 years) specified in the order.

(3) Before making the application, the authorised person must:
   (a) take reasonable steps to discover who has an interest in the retention of the thing; and
(b) if it is practicable to do so, notify each person whom the authorised person believes to have such an interest of the proposed application.

49 Monitoring warrants

(1) An authorised person may apply to a magistrate for a warrant under this section in relation to premises.

(2) Subject to subsection (3), the magistrate may issue the warrant if the magistrate is satisfied, by information on oath, that it is reasonably necessary that one or more authorised persons should have access to the premises for the purposes of finding out whether this Act or the regulations have been complied with.

(3) The magistrate must not issue the warrant unless the authorised person or some other person has given to the magistrate, either orally or by affidavit, such further information (if any) as the magistrate requires concerning the grounds on which the issue of the warrant is being sought.

(4) The warrant must:

(a) authorise one or more authorised persons (whether or not named in the warrant), with such assistance and by such force as is necessary and reasonable:
   (i) to enter the premises; and
   (ii) to exercise the powers set out in subsection 48(1) in relation to the premises; and

(b) state whether the entry is authorised to be made at any time of the day or night or during specified hours of the day or night; and

(c) specify the day (not more than 6 months after the issue of the warrant) on which the warrant ceases to have effect; and

(d) state the purpose for which the warrant is issued.

50 Offence related warrants

(1) An authorised person may apply to a magistrate for a warrant under this section in relation to premises.
Section 51

(2) Subject to subsection (3), the magistrate may issue the warrant if the magistrate is satisfied, by information on oath, that there are reasonable grounds for suspecting that there is, or there may be within the next 72 hours, in or on the premises evidential material.

(3) The magistrate must not issue the warrant unless the authorised person or some other person has given to the magistrate, either orally or by affidavit, such further information (if any) as the magistrate requires concerning the grounds on which the issue of the warrant is being sought.

(4) The warrant must:

(a) name one or more authorised persons; and

(b) authorise the persons so named, with such assistance and by such force as is necessary and reasonable:

(i) to enter the premises; and

(ii) to exercise the powers set out in subsections 47(4) and 48(1); and

(iii) to seize the evidential material; and

(c) state whether the entry is authorised to be made at any time of the day or night or during specified hours of the day or night; and

(d) specify the day (not more than one week after the issue of the warrant) on which the warrant ceases to have effect; and

(e) state the purpose for which the warrant is issued.

51 Offence related warrants by telephone

(1) If, in an urgent case, an authorised person considers it necessary to do so, the person may apply to a magistrate by telephone for a warrant under section 50 in relation to premises.

(2) Before applying for the warrant, the person must prepare an information of the kind mentioned in subsection 50(2) in relation to the premises that sets out the grounds on which the warrant is sought.

(3) If it is necessary to do so, the person may apply for the warrant before the information is sworn.

(4) If the magistrate is satisfied:
(a) after having considered the terms of the information; and
(b) after having received such further information (if any) as the
magistrate requires concerning the grounds on which the
issue of the warrant is being sought;
that there are reasonable grounds for issuing the warrant, the
magistrate may complete and sign the same warrant that the
magistrate would issue under section 50 if the application had been
made under that section.

(5) If the magistrate completes and signs the warrant:
(a) the magistrate must:
   (i) tell the authorised person what the terms of the warrant
       are; and
   (ii) tell the authorised person the day on which and the time
        at which the warrant was signed; and
   (iii) tell the authorised person the day (not more than one
        week after the magistrate completes and signs the
        warrant) on which the warrant ceases to have effect; and
   (iv) record on the warrant the reasons for granting the
        warrant; and
(b) the authorised person must:
   (i) complete a form of warrant in the same terms as the
       warrant completed and signed by the magistrate; and
   (ii) write on the form the name of the magistrate and the
day on which and the time at which the warrant was
signed.

(6) The authorised person must also, not later than the day after the
day of expiry or execution of the warrant, whichever is the earlier,
send to the magistrate:
(a) the form of warrant completed by the person; and
(b) the information referred to in subsection (2), which must
have been duly sworn.

(7) When the magistrate receives those documents, the magistrate
must:
(a) attach them to the warrant that the magistrate completed and
signed; and
Section 51A

(b) deal with them in the way in which the magistrate would have dealt with the information if the application had been made under section 50.

(8) A form of warrant duly completed under subsection (5) is authority for any entry, search, seizure or other exercise of a power that the warrant signed by the magistrate authorises.

(9) If:
    (a) it is material, in any proceedings, for a court to be satisfied that an exercise of a power was authorised by this section; and
    (b) the warrant signed by the magistrate authorising the exercise of the power is not produced in evidence;

the court must assume, unless the contrary is proved, that the exercise of the power was not authorised by such a warrant.

(10) A reference in this Part to a warrant under section 50 includes a reference to a warrant signed by a magistrate under this section.

51A Searches at request of manufacturer

(1) A person may request the Secretary to arrange for an authorised person to inspect premises, and specified processes being carried out on those premises, for the purposes of paragraph 2 of Article 3 of the Mutual Recognition Convention.

(2) An authorised person may make an inspection in accordance with arrangements under subsection (1).

51B Offences relating to warrants

(1) A person must not make, in an application for a warrant, a statement that the person knows to be false or misleading in a material particular.

    Maximum penalty: Imprisonment for 2 years.

(2) A person must not:
    (a) state in a document that purports to be a form of warrant under section 51 the name of a magistrate unless that magistrate issued the warrant; or
Section 52

(b) state on a form of warrant under that section a matter that, to the person’s knowledge, departs in a material particular from the form authorised by the magistrate; or

(c) purport to execute, or present to another person, a document that purports to be a form of warrant under that section that the first-mentioned person knows:
   (i) has not been approved by a magistrate under that section; or
   (ii) to depart in a material particular from the terms authorised by a magistrate under that section; or

(d) give to a magistrate a form of warrant under that section that is not the form of warrant that the person purported to execute.

Maximum penalty:  Imprisonment for 2 years.

52 Identity cards

(1) The Secretary is to ensure that each authorised person is issued with an identity card that incorporates a recent photograph of the person.

(3) Where a person ceases to be an authorised person, the person must, as soon as practicable after so ceasing, return the person’s identity card to the Secretary.

Maximum penalty:  1 penalty unit.

(4) An offence under subsection (3) is an offence of strict liability.

Note: For strict liability, see section 6.1 of the Criminal Code.
Part 6-3—National Drugs and Poisons Schedule Committee

52A Definitions

In this Part, unless the contrary intention appears:

Committee means the National Drugs and Poisons Schedule Committee referred to in section 52B.

current Poisons Standard means:
(a) if no document has been prepared under paragraph 52D(2)(b)—the first Poisons Standard; or
(b) otherwise—the document last prepared under that paragraph.

first Poisons Standard means the latest edition at the commencement of this Part of the document known as the Standard for the Uniform Scheduling of Drugs and Poisons published by the Australian Health Ministers’ Advisory Council.

scheduling, in relation to a substance, means determining the schedule or schedules to the current Poisons Standard in which the name or a description of the substance is to be included.

substance means any medicine or poison.

52B Establishment and constitution of Committee

(1) There is established a committee to be known as the National Drugs and Poisons Schedule Committee.

(2) Subject to subsection (3), the Committee is to be constituted, and to hold meetings and to make decisions, in accordance with the regulations.

(3) The Commonwealth, each State, the Northern Territory and the Australian Capital Territory are each entitled to nominate a representative on the Committee.
52C Functions of Committee

The functions of the Committee are:

(a) to make decisions in relation to the classification and scheduling of substances; and

(b) to provide technical advice to governments in relation to:
   (i) the legislative restrictions (including restrictions as to accessibility and availability) to be imposed in respect of particular substances; and
   (ii) the policies to be adopted with respect to the labelling, packaging and advertising of substances; and

(c) to maintain the current Poisons Standard; and

(d) to facilitate the harmonisation between Australia and New Zealand of the legislative provisions relating to the classification and scheduling of substances; and

(e) to undertake public consultation with respect to matters relating to the classification and scheduling of substances that are of public interest or significance; and

(f) to consider any matters referred to it by:
   (i) the Minister or Secretary; or
   (ii) the subcommittee of the Australian Health Ministers’ Advisory Council known as the National Co-ordinating Committee on Therapeutic Goods;

and report to the Minister, Secretary or subcommittee the results of its consideration; and

(g) any other functions that are prescribed by the regulations.

52D Poisons Standard

(1) On the commencement of this Part, the first Poisons Standard is taken to have been prepared and made available by the Committee.

(2) Subject to this Act and the regulations, the Committee has power:
   (a) to amend the current Poisons Standard; or
   (b) to prepare a document (a new Poisons Standard) that includes schedules containing the names or descriptions of substances, in substitution for the current Poisons Standard.
(3) As soon as practicable after a new Poisons Standard is prepared, the Committee must cause a notice to be published in the Gazette stating:
   (a) that a new Poisons Standard has been prepared; and
   (b) the date on which the new Poisons Standard comes into effect; and
   (c) a place from which copies of the new Poisons Standard can be purchased.

(4) As soon as practicable after an amendment is made to the current Poisons Standard, the Committee must cause a notice to be published in the Gazette stating:
   (a) that an amendment has been made to the Poisons Standard; and
   (b) the date on which the amendment comes into effect; and
   (c) a place from which copies of the amendment can be purchased.

(5) In this section:

   *amend*, in relation to the current Poisons Standard, means:
   (a) alter any provision (including a reference to a substance) in the current Poisons Standard; or
   (b) omit any provision (including a reference to a substance) from the current Poisons Standard; or
   (c) insert any provision (including a reference to a substance) in the current Poisons Standard.

52E Matters to be taken into account in exercising powers

(1) In exercising its powers under subsection 52D(2), the Committee must take the following matters into account (where relevant):
   (a) the toxicity and safety of a substance;
   (b) the risks and benefits associated with the use of a substance;
   (c) the potential hazards associated with the use of a substance;
   (d) the extent and patterns of use of a substance;
   (e) the dosage and formulation of a substance;
Section 52E

(f) the need for access to a substance, taking into account its toxicity compared with other substances available for a similar purpose;
(g) the potential for abuse of a substance;
(h) the purposes for which a substance is to be used;
(i) any other matters that the Committee considers necessary to protect public health, including the risks (whether imminent or long-term) of death, illness or injury resulting from its use; and may take into account the labelling, packaging and presentation of a substance.

(2) In taking into account the matters referred to in subsection (1), the Committee must comply with any guidelines of the Australian Health Ministers’ Advisory Council or the subcommittee of the Council known as the National Co-ordinating Committee on Therapeutic Goods, notified to the Committee for the purposes of this section.
Section 52F

Part 6-4—Complementary medicines

52F Definitions

In this Part, unless the contrary intention appears:

*active ingredient* means the therapeutically active component in a medicine’s final formulation that is responsible for its physiological or pharmacological action.

*complementary medicines* means therapeutic goods consisting wholly or principally of one or more designated active ingredients, each of which has a clearly established identity and:

(a) a traditional use; or

(b) any other use prescribed in the regulations.

*designated active ingredient* means an active ingredient, or a kind of active ingredient, mentioned in Schedule 14 to the Therapeutic Goods Regulations.

*traditional use*, in relation to a designated active ingredient, means use of the designated active ingredient that:

(a) is well documented, or otherwise established, according to the accumulated experience of many traditional health care practitioners over an extended period of time; and

(b) accords with well-established procedures of preparation, application and dosage.

Note: An example of traditional use is use in Chinese traditional medicine.

52G Establishment and constitution of Committee

(1) There is established a committee to be known as the Complementary Medicines Evaluation Committee.

(2) The Committee is to have the functions prescribed in the regulations in relation to complementary medicines.

(3) The Committee is to be constituted, and to hold meetings and to make recommendations, in accordance with the regulations.
Chapter 7—Miscellaneous

53 Retention of material on withdrawal of application

Where a person withdraws an application for:
(a) registration; or
(b) listing; or
(c) a conformity assessment certificate; or
(d) inclusion of a kind of medical device in the Register; or
(e) a licence;
the Department may retain the application and any material submitted in connection with the application.

54 Indictable offences and forfeiture

(1) An offence against section 22A, 29A, 29B, 41FE, 41MP or 41MQ is an indictable offence.

(3) Where a court convicts a person of an offence against this Act in relation to any therapeutic goods, the court may order that the goods be forfeited to the Commonwealth and, where such an order is made, the goods become the property of the Commonwealth.

(4) Where goods are so forfeited, the Secretary may cause notice of the forfeiture to be published in the Gazette.

(5) Goods forfeited under an order referred to in subsection (3) are to be disposed of in such manner as the Secretary directs.

54AA Offences for contravening conditions or requirements imposed under the regulations

(1) If:
(a) a person holds a licence or a permission to import or export therapeutic goods; and
(b) the person engages in conduct; and
Section 54AB

(c) the conduct breaches a condition or a requirement to which the licence or permission is subject under the regulations; the person is guilty of an offence punishable on conviction by a fine of no more than the number of penalty units specified in whichever of n or (3) applies whichever of subsection (2) or (3) applies.

(1A) In subsection (1):

engage in conduct means:

(a) do an act; or
(b) omit to perform an act.

(2) If:

(a) the condition or requirement relates to the possession, custody, transport, use or disposal of the goods; or
(b) the regulation providing for the condition or requirement states that the purpose of the condition or requirement is to protect the safety of the public;

the number of penalty units for the contravention is 240 penalty units.

(3) If subsection (2) does not apply, the number of penalty units for the contravention is 50 penalty units.

54AB Damage etc. to documents

(1) A person is guilty of an offence if:

(a) the person damages, destroys, alters, conceals or falsifies a document; and
(b) the document is created, retained or issued for the purposes of this Act, or for purposes that include the purposes of this Act.

Maximum penalty: Imprisonment for 5 years or 2,000 penalty units, or both.

(2) Strict liability applies to paragraph (1)(b).

Note: For strict liability, see section 6.1 of the Criminal Code.
54A Time for bringing prosecutions

A prosecution for an offence against this Act may be commenced at any time within 3 years after the commission of the offence.

55 Conduct by directors, servants and agents

(1) Where, in proceedings for an offence against this Act, it is necessary to establish the state of mind of a body corporate in relation to particular conduct, it is sufficient to show:

(a) that the conduct was engaged in by a director, servant or agent of the body corporate within the scope of his or her actual or apparent authority; and

(b) that the director, servant or agent had the state of mind.

(2) Any conduct engaged in on behalf of a body corporate by a director, servant or agent of the body corporate within the scope of his or her actual or apparent authority is to be taken, for the purposes of a prosecution for an offence against this Act, to have been engaged in also by the body corporate unless the body corporate establishes that the body corporate took reasonable precautions and exercised due diligence to avoid the conduct.

(3) Where, in proceedings for an offence against this Act, it is necessary to establish the state of mind of a person other than a body corporate in relation to particular conduct, it is sufficient to show that:

(a) the conduct was engaged in by a servant or agent of the person within the scope of his or her actual or apparent authority; and

(b) the servant or agent had the state of mind.

(4) Any conduct engaged in on behalf of a person other than a body corporate (in this subsection called the employer) by a servant or agent of the employer within the scope of his or her actual or apparent authority is to be taken, for the purposes of a prosecution for an offence against this Act, to have been engaged in also by the employer unless the employer establishes that he or she took reasonable precautions and exercised due diligence to avoid the conduct.
(5) Where:
   (a) a person other than a body corporate is convicted of an
       offence; and
   (b) the person would not have been convicted of the offence if
       subsections (3) and (4) had not been enacted;
       the person is not liable to be punished by imprisonment for that
       offence.

(6) A reference in subsection (1) or (3) to the state of mind of a person
    includes a reference to:
    (a) the knowledge, intention, opinion, belief or purpose of the
        person; and
    (b) the person’s reasons for the intention, opinion, belief or
        purpose.

(7) A reference in this section to a director of a body corporate
    includes a reference to a constituent member of a body corporate
    incorporated for a public purpose by a law of the Commonwealth,
    of a State or of a Territory.

(8) A reference in this section to engaging in conduct includes a
    reference to failing or refusing to engage in conduct.

56  Judicial notice

All courts (except in proceedings under Chapter 4) are to take
judicial notice of the British Pharmacopoeia and of the British
Pharmacopoeia (Veterinary).

56A  Certificates to provide evidence of certain matters

(1) The Secretary or a person authorised in writing by him or her to
give certificates under this section may certify in writing that, at a
specified time, or at all times during a specified period:
   (a) there was no exemption in effect under section 18 in relation
       to particular therapeutic goods; or
   (aa) particular medical devices were not exempt devices;
   (b) there was no approval or authority under section 19 granted
       to a particular person in relation to particular therapeutic
       goods; or
(ba) there was no approval or authority in effect under section 41HB or 41HC granted to a particular person in relation to particular medical devices;
(c) there was no approval under section 19A granted to a particular person in relation to particular therapeutic goods; or
(d) particular therapeutic goods were or were not included in the Register as registered goods; or
(e) particular therapeutic goods were or were not included in the Register as listed goods; or
(ea) particular medical devices were or were not medical devices of a kind included in the Register; or
(eb) particular medical devices were suspended from the Register; or
(f) particular therapeutic goods were included in the Register subject to conditions including those specified in the certificate; or
(g) the registration, listing or inclusion in the Register of the particular therapeutic goods had been cancelled; or
(h) there was no declaration under section 7 which applied to particular therapeutic goods; or
(i) a person was or was not the holder of a licence in force under Part 3-3; or
(j) the licence is subject to conditions including those specified in the certificate; or
(k) there was no exemption in effect under subsection 34(1) that applied to particular therapeutic goods or a particular class of therapeutic goods; or
(l) there was no exemption in effect under subsection 34(2) that applied to a particular person in relation to one or more of the following:
   (i) the manufacture of particular therapeutic goods;
   (ii) a particular step in the manufacture of particular therapeutic goods;
   (iii) the manufacture of a particular class of therapeutic goods;
   (iv) a particular step in the manufacture of a particular class of therapeutic goods; or
Section 57

(m) a conformity assessment certificate has been issued relating to a particular kind of medical device; or
(n) a conformity assessment certificate was subject to conditions including those specified in the certificate under this section; or
(o) a conformity assessment certificate was suspended.

(2) A certificate under subsection (1) may relate to more than one of the matters referred to in paragraphs (1)(a) to (o).

(3) In proceedings for an offence against this Act, a certificate under subsection (1) is prima facie evidence of the matters specified in the certificate.

(4) In proceedings for an offence against section 14 or 41MA, a certificate by the Secretary to the effect that:
   (a) the Secretary did not consent to the importation, supply or exportation that is the subject of the proceedings; or
   (b) the Secretary consented to that importation, supply or exportation subject to conditions specified in the certificate; is prima facie evidence of the matters specified in the certificate.

(5) In proceedings for an offence against this Act, a document purporting to be a certificate given under this section is, unless the contrary is proved, taken to be such a certificate and to have been duly given.

57 Delegation

(1) Subject to subsections (2), (6) and (8) to (10), the Minister or the Secretary may, by signed instrument, delegate to:
   (a) an officer of the Department; or
   (b) an officer of an authority of the Commonwealth that has functions in relation to therapeutic goods; or
   (ba) an APS employee in an Agency (within the meaning of the Public Service Act 1999) that has functions in relation to therapeutic goods; or
(c) a person occupying or acting in an office, or holding an appointment, declared by the regulations to be an office or appointment the occupant or holder of which may be a delegate under this section;

all or any of his or her powers and functions under this Act.

(2) The powers of the Secretary under paragraph 19(1)(a) or 41HB(1)(d) may be delegated under subsection (1) only to a person referred to in paragraph (1)(a) or (c) who is registered, or eligible for registration, in a State or internal Territory, as a medical or dental practitioner or as a pharmacist.

(3) Subject to the regulations, the Secretary may, in such circumstances as are prescribed, by signed instrument, delegate all or any of his or her powers under paragraph 19(1)(a) or 41HB(1)(d) to a person who is registered, in a State or internal Territory, as a medical or dental practitioner.

(4) A delegate under subsection (3) is, in the exercise of a delegated power, subject to the directions of:

(a) the Secretary; or

(b) an officer of the Department authorised in writing by the Secretary; or

(c) a person referred to in paragraph (1)(c).

(5) Without limiting the generality of matters that may be dealt with by regulations made for the purposes of subsection (3), the regulations may make provision in relation to the following:

(a) the persons who may be delegates;

(b) the circumstances in which delegates may grant approvals for the purposes of paragraph 19(1)(a) or 41HB(1)(d);

(c) the conditions to which any approvals granted by delegates are to be subject;

(d) requiring information to be given by delegates to the Secretary.

(6) The powers of the Secretary under subsection 19(5) or 41HC(1) may be delegated only to a person referred to in paragraph (1)(a) or (c) who is registered, or eligible for registration, in a State or internal Territory as a medical or dental practitioner.
Section 58

(7) The regulations may prescribe the circumstances in which, and the requirements subject to which, delegates may grant authorities under subsection 19(5) or 41HC(1).

(8) The powers of the Secretary under section 19A may be delegated only to either or both of the following persons:
   (a) the National Manager of the Therapeutic Goods Administration;
   (b) the Director of the Drug Safety and Evaluation Branch of the Therapeutic Goods Administration.

(9) The Minister must not delegate his or her powers or functions under section 6AA or 23AA.

(10) The power of the Minister under paragraph 18A(2)(a) may be delegated only to the Secretary.

58 Export certifications

(1) The Secretary may issue export certification for goods for therapeutic use in humans, including certifications for the purposes of the World Health Organisation Certification Scheme on the Quality of Pharmaceutical Products Moving in International Commerce.

(2) A State or Territory must not issue export certifications for goods for therapeutic use in humans.

(3) Such fee as is prescribed is payable in respect of:
   (a) an application for a certification under this section; and
   (b) where an inspection of manufacturing premises is necessary for the purposes of the issue of a certification under this section—the inspection of those premises.

59 Fees

(1) No fees are payable under this Act in respect of an event occurring before 1 July 1990.

(2) Fees prescribed under this Act must not be such as to amount to taxation.
(3) No licence or inspection fees are to apply to non-profit hospital supply units.

60 Review of decisions

(1) In this section and section 60A:

*decision* has the same meaning as in the *Administrative Appeals Tribunal Act 1975*.

*initial decision* means a decision of the Secretary or of a delegate of the Secretary:

(a) under the definition of *therapeutic devices* in subsection 3(1) or under subsection 7(1) or 41BD(3); or

(ab) under section 9C or 9D; or

(b) refusing to grant a consent under section 14; or

(c) under Part 3-2 (registration and listing of therapeutic goods); or

(d) under Part 3-3 (manufacturing of therapeutic goods); or

(e) under Part 4-4 (conformity assessment certificates); or

(f) under Part 4-5 (including medical devices in the Register), other than:

(i) a decision under section 41FH (selecting applications for auditing); or

(ii) a decision about which aspects of the matters referred to in paragraphs 41FI(1)(a) and (b) to consider in auditing an application under Subdivision C of Division 1 of Part 4-5; or

(g) under Part 4-6 (suspension and cancellation from the Register); or

(h) under Part 4-7 (exempting medical devices from inclusion in the Register); or

(i) under Part 4-8 (obtaining information); or

(j) under Part 4-9 (public notification and recovery of medical devices); or

(k) refusing to grant, or imposing conditions on a grant of, a consent for the purposes of section 41MA (non-compliance with essential principles); or

(l) under section 42DF, 42DH or 42DI.
Section 60

A *reviewable decision* means a decision of the Minister under subsection (3).

(1A) For the avoidance of doubt, the following are not initial decisions for the purposes of this section or section 60A:

(a) a proposal to suspend a conformity assessment certificate under section 41EM;
(b) a proposal to revoke a conformity assessment certificate under section 41ET;
(c) a proposal to suspend a kind of medical device from the Register under section 41GA;
(d) a proposal to cancel the entry of a kind of medical device on the Register under section 41GN.

(2) A person whose interests are affected by an initial decision may, by notice in writing given to the Minister:

(a) in the case of a decision particulars of which are required to be notified in the Gazette—within 90 days after those particulars are so notified; or
(b) in any other case—within 90 days after the decision first comes to the person’s notice;

request the Minister to reconsider the decision.

(3) Subject to paragraph 60A(2)(b), the Minister must, as soon as practicable after receiving a request under subsection (2), reconsider the initial decision and, as a result of that reconsideration, may:

(a) confirm the initial decision; or
(b) revoke the initial decision, or revoke that decision and make a decision in substitution for the initial decision.

(4) Where a person who has made a request under subsection (2) does not receive notice of the decision of the Minister on reconsideration, or (if applicable) notice that the matter has been remitted under paragraph 60A(2)(b), within 60 days of the making of the request, the Minister is to be taken to have confirmed the original decision.

(5) After reconsideration of an initial decision, the Minister must give the applicant a notice in writing stating the result of the reconsideration and that the applicant may, except where
subsection 28(4) of the *Administrative Appeals Tribunal Act 1975* applies, apply for a statement setting out the reasons for the decision on reconsideration and may, subject to that Act, make an application to the Administrative Appeals Tribunal for review of that decision.

(6) Where written notice of the making of an initial decision is given to a person whose interests are affected by the decision, the notice is to include a statement to the effect that a person whose interests are affected by the decision may:

(a) seek a reconsideration of the decision under this section; and
(b) subject to the *Administrative Appeals Tribunal Act 1975*, if the person is dissatisfied with the decision upon reconsideration, make an application to the Administrative Appeals Tribunal for review of that decision.

(7) Any failure to comply with the requirements of subsection (5) or (6) in relation to a decision does not affect the validity of the decision.

(8) An application may be made to the Administrative Appeals Tribunal for review of a reviewable decision.

### 60A New information on review—discretion to remit

(1) This section applies only if the Secretary or an authorised delegate makes a decision under section 25 or 41EC in relation to therapeutic goods.

(2) If a person (the *appellant*) whose interests are affected by the decision requests the Minister to reconsider the decision, and lodges new information in support of that request, the Minister must either:

(a) take that information into account when he or she reconsiders the decision; or
(b) remit the matter to an authorised delegate for a fresh decision.

(3) If the appellant applies to the Administrative Appeals Tribunal for review of the decision on reconsideration, and lodges new information in support of that application, the Tribunal may, if the Tribunal thinks fit, remit the matter to an authorised delegate for a fresh decision.
Section 60A

(4) The Tribunal must not remit the matter under subsection (3) if all of the new information is information that the Minister took into account under paragraph (2)(a) in making the decision on reconsideration.

(5) If:

(a) the appellant lodges new information in support of an application to the Administrative Appeals Tribunal for review of the decision on reconsideration; and
(b) the Tribunal does not remit the matter under subsection (3);

the Tribunal, in reviewing the decision on reconsideration:

(c) may consider new information (if any) that the Minister took into account under paragraph (2)(a) in making the decision on reconsideration; and
(d) must not consider any other new information, except new information that indicates that the quality, safety or efficacy of the therapeutic goods is unacceptable.

(6) If:

(a) the matter relates to a decision under section 25; and

(b) the Minister or the Tribunal remits the matter; and

the authorised delegate must make a decision under section 25, taking into account the new information, as if a fresh application for registration had been made.

(6A) If:

(a) the matter relates to a decision under section 41EC; and
(b) the Minister or the Tribunal remits the matter; and
(c) the appellant has paid, as a further conformity assessment fee, the conformity assessment fee that the appellant would have to pay under section 41LA on making a new application for a conformity assessment certificate;

the authorised delegate must make a decision under section 41EC, taking into account the new information, as if a fresh application for a conformity assessment certificate had been made.
Section 61

(7) To remove any doubt, the authorised delegate’s fresh decision is to be treated, for the purposes of subsequent applications of section 60 and this section, as a decision under Part 3-2 or 4-4.

(8) In this section:

authorised delegate means a delegate of the Secretary exercising a power to decide whether to register therapeutic goods.

new information means information that:

(a) was in existence at the time the decision referred to in subsection (1) was made; and
(b) was not made available to the Secretary or authorised delegate for the purpose of making the decision; and
(c) is relevant to that decision;

and includes any opinions that are wholly or substantially based on such information (whether or not the opinions were formed before or after the decision was made).

61 Release of information

(1) In this section:

therapeutic goods information means information in relation to therapeutic goods that came into the possession of the Department in connection with the performance of the Department’s functions (including functions relating to the EC Mutual Recognition Agreement or the EFTA Mutual Recognition Agreement).

(2) The Secretary may:

(a) release to the Director-General of the World Health Organisation therapeutic goods information relating to:

(i) notifications concerning therapeutic goods the consumption or supply of which in Australia has been prohibited or severely restricted, or relating to the reasons for that action; or

(ii) the licensing status of Australian manufacturers of therapeutic goods and their compliance with the manufacturing principles; or

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(iii) the content of reports to the Department concerning adverse effects of therapeutic goods; or
(iv) the issue of, imposition of conditions on, or revocation of, conformity assessment certificates;
for use in the development of policies relating to the regulation of therapeutic goods or for the provision of information to regulatory authorities of member countries of the World Health Organisation; or
(b) release, in confidence, therapeutic goods information to the Director-General of the World Health Organisation, being information concerning proceedings of committees established under the regulations.

(3) The Secretary may release to the head of an authority of the Commonwealth, a State or a Territory that has functions relating to therapeutic goods, therapeutic goods information relating to:
(a) reported problems and complaints concerning therapeutic goods, the Department’s investigation of those problems and complaints and any action that the Department has taken or proposes to take in relation to those problems and complaints; or
(b) reports of inspections conducted under this Act or the regulations; or
(c) decisions to revoke or suspend, or not to issue, licences for the manufacturing of therapeutic goods; or
(d) conditions of licences; or
(e) reports of the testing of samples of therapeutic goods; or
(f) the issue of, imposition of conditions on, or revocation of, conformity assessment certificates;
for use in the performance of those functions.

(3A) The Secretary may release information obtained in response to a notice under section 31A, 31B, 41JD, 41JE or 41JF to:
(a) the head of an authority, or an authority, of the Commonwealth, a State or a Territory that has functions relating to therapeutic goods; and
(b) the body in a State or Territory responsible for the registration of medical practitioners in that State or Territory; and
(c) the body in a State or Territory responsible for the registration of pharmacists in that State or Territory.

(4) The Secretary may release to the head of a national regulatory authority of another country, being an authority that has national responsibility relating to therapeutic goods, therapeutic goods information relating to:

(a) recommendations of advisory committees on therapeutic goods supplied in or proposed for supply in Australia, and any conditions that are or will be applicable to that supply; or

(b) decisions on the registration or listing, or the cancellation of the registration or listing, of therapeutic goods; or

(ba) decisions on the inclusion of kinds of medical devices in the Register, or the cancellation of the inclusion of kinds of medical devices in the Register; or

(c) the withdrawal from supply in Australia of therapeutic goods and the reasons for that action; or

(d) the licensing status of Australian manufacturers of therapeutic goods and their compliance with the manufacturing principles; or

(e) proceedings of committees established under the regulations; or

(f) the issue of, imposition of conditions on, or revocation of, conformity assessment certificates;

for use in the performance of those functions or for furthering international co-operation in the regulation of therapeutic goods.

(4A) The Secretary may release to:

(a) the head of an authority of the Commonwealth, a State or a Territory that has functions relating to therapeutic goods, health or law enforcement; or

(b) the head of a national regulatory authority of another country that has national responsibility relating to therapeutic goods, health or law enforcement;

therapeutic goods information relating to one or more of the following:

(c) notifications received under section 42T;

(d) action taken by the Secretary under Part 5-3;

(e) contraventions, or possible contraventions, of Part 5-3;
Section 61

(f) any cases, or possible cases, of actual or potential tampering with therapeutic goods.

(5) The Secretary may release to the head of a national regulatory authority of another country, or the head of an international organisation, being another country or an organisation with which the Commonwealth has co-operative arrangements relating to the assessment or regulation of therapeutic goods, therapeutic goods information the release of which is consistent with those arrangements.

(6) The Secretary may release to a person, on application by that person, therapeutic goods information of a kind identified in the regulations relating to:
   (a) therapeutic goods included in the Register; or
   (b) therapeutic goods in relation to which an application for registration, listing or inclusion in the Register has been made.

(6A) Regulations made for the purposes of subsection (6) may:
   (a) relate to therapeutic goods generally or to a class of such goods; and
   (b) authorise the release of therapeutic goods information to persons generally or to a class of persons; and
   (c) authorise the release of information on application or otherwise.

(7) The Secretary may release therapeutic goods information:
   (a) the release of which is necessary to ensure the safe use of particular therapeutic goods; or
   (b) relating to the reasons for the withdrawal of therapeutic goods from supply in Australia.

(8) Subject to section 25A, therapeutic goods information provided to the Department in relation to a matter may:
   (a) be used by the Department in the consideration of another matter within its functions relating to therapeutic goods; and
   (b) be provided to a committee appointed to advise the Minister or the Secretary on matters relating to therapeutic goods, including a committee of the National Health and Medical Research Council.

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Section 62

(8A) Regulations prescribing fees in respect of applications for information under the regulations:
(a) may include provision for the payment of deposits on account of such fees; and
(b) may provide for fees that take into account the time spent by officers of the Department in:
(i) searching for or retrieving information; or
(ii) making, or doing anything related to the making of, a decision on an application; and
(c) may provide for fees that take into account the direct costs incurred by the Commonwealth in making available an officer to supervise the inspection by an applicant of any document containing information to which an application relates.

(8C) If, under the regulations, a person is liable to pay a fee in respect of an application for information, the Secretary must notify the person, in writing, accordingly, and must give to the person, together with that notification, a statement setting out the basis on which the amount of that fee is calculated.

(9) Civil proceedings do not lie against the Secretary or a delegate of the Secretary in respect of loss, damage or injury of any kind suffered by another person as a result of the release of information in good faith under this section or the regulations.

(10) Nothing in this or any other Act requires the Secretary to disclose to any person, court or tribunal information referred to in subsection 25(2E) or 26(2D) if the disclosure would constitute a breach of the Mutual Recognition Convention.

(11) This section (except subsection (10)) has effect subject to the Freedom of Information Act 1982.

62 Consequential amendments

The Acts specified in the Schedule are amended as set out in the Schedule.
Section 63

63 Regulations

(1) The Governor-General may make regulations, not inconsistent with this Act, prescribing matters:
   (a) required or permitted to be prescribed by this Act; or
   (b) necessary or convenient to be prescribed for carrying out or giving effect to this Act.

(2) The regulations may:
   (a) make provision in relation to:
      (i) the establishment of committees to advise the Minister or the Secretary on matters relating to therapeutic goods; and
      (ii) the functions and powers of those committees; and
      (iii) the payment of remuneration and allowances to members of those committees; and
   (b) prescribe requirements for the storage and transport of therapeutic goods; and
   (c) prescribe requirements for the advertising of therapeutic goods; and
   (d) provide for the procedures to be followed by the Department in the sampling and testing of therapeutic goods; and
   (da) provide for the periods within which evaluations under section 25 in relation to specified therapeutic goods or specified classes of such goods are to be completed; and
   (db) provide for the periods within which decisions under section 41EP to revoke suspensions of conformity assessment certificates are to be made, in cases where applications for revocation have been made under paragraph 41EP(2)(a); and
   (dc) provide for the periods within which decisions on applications for the issuing of conformity assessment certificates under Part 4-4 are to be made if considering the applications involves examining the design of medical devices; and
   (dd) provide for the periods within which decisions under section 41GD to revoke suspensions of entries on the Register are to be made, in cases where applications for revocation have been made under paragraph 41GD(2)(a); and
(e) prescribe requirements for informational material that is included with therapeutic goods; and

(f) make provision for the transfer of registration, listing or inclusion in the Register of therapeutic goods and of licences; and

(g) make provision for the testing of therapeutic goods, the inspection of manufacturing operations or the evaluation of data concerning therapeutic goods by the Department at the request of persons; and

(h) prescribe fees in respect of matters under this Act or the regulations; and

(j) prescribe penalties not exceeding 10 penalty units for offences against the regulations.

(3) The regulations may:

(a) prescribe different fees under this Act in relation to:
   (i) different classes of goods; or
   (ii) in the case of fees under Part 3-3—different steps in the manufacture of goods; or

(b) provide for the refund, reduction or waiving of fees or charges in cases identified in the regulations; or

(c) specify the type of information relating to therapeutic goods manufactured by licence holders that the Secretary may, under subsection 37(2), require to be supplied by the holders of licences at the time of payment of annual licensing charges in respect of the licences.

(3A) The regulations may provide for:

(a) the granting of a licence or permission to import or export therapeutic goods; and

(b) licences or permissions to import or export therapeutic goods to be subject to conditions or requirements; and

(c) the assignment of a licence or permission to import or export therapeutic goods; and

(d) the surrender of a licence or permission to import or export therapeutic goods; and

(e) the revocation of a licence or permission to import or export therapeutic goods.
Section 63

(4) The regulations may make provision for a matter by applying, adopting or incorporating, with or without modification, any matter contained in an instrument:
   (a) as that instrument is in force at the time when the regulations take effect; or
   (b) as that instrument is in force from time to time.

(5) For the purposes of section 2, regulations may be made before the commencement of this Act as if this Act were in force, but do not come into effect on a day earlier than the day on which this Act commences.
Chapter 8—Repeal and transitional provisions

64 Interpretation

In this Part, former Act means the Therapeutic Goods Act 1966.

65 Repeal

The former Act is repealed.

66 Transitional arrangements for goods required to be registered or listed

(1) This section applies to therapeutic goods in relation to a person if, immediately before the commencement of this Act, the person was supplying goods of that kind in Australia for use in humans.

(2) Where:
   (a) this section applies to therapeutic goods in relation to a person; and
   (b) the Secretary is not aware of the person having been convicted of an offence against a law of the Commonwealth, of a State or of an internal Territory in respect of goods of that kind during the period of 2 years ending on the commencement of this Act; and
   (c) if the goods are imported goods—the Secretary is not aware of the person having, during that period, imported goods of that kind into Australia otherwise than in accordance with regulations in force under the Customs Act 1901;
   subsections 20(1) and (2) do not apply to goods of that kind in relation to the person during the period of 3 months after that commencement.

(3) Where:
   (a) this section applies to therapeutic goods in relation to a person; and
Section 66

(b) the person makes an application for registration or listing of goods of that kind in accordance with section 23 and within 3 months after the commencement of this Act;

then:

(c) subsection 20(1) does not apply to goods of that kind in relation to the person during the period of 6 months after that commencement or before the end of such longer period as the Secretary specifies by notice published in the Gazette before the end of that first-mentioned period; and

(d) subsection 20(2) does not apply to goods of that kind in relation to the person during the period of 12 months after that commencement or before the end of such longer period as the Secretary specifies by notice published in the Gazette before the end of that first-mentioned period.

(3A) If, on an application under subsection (3), goods have been registered without having been evaluated, the Secretary may, if he or she thinks it appropriate, give the person in relation to whom the goods are registered written notice that the goods are to be evaluated to determine whether they should continue to be registered.

(4) A person who makes an application in accordance with subsection (3) is not required to pay:

(a) any application fee for the registration or listing of the goods to which the application relates; or

(b) in the case of an application for the registration of goods—any fee for the evaluation of the goods for registration;

but where the goods are later evaluated to determine whether the goods should continue to be registered, such fee as is prescribed is payable in respect of that evaluation.

(4A) In relation to an evaluation conducted for the purposes of this section:

(a) section 25 has effect as if:

(i) the person in respect of whom the goods are registered were an applicant for the registration of the goods; and

(ii) the reference in paragraph (1)(b) to an evaluation fee under section 24 were a reference to a fee payable under subsection (4) of this section; and
(b) sections 24A, 24B and 24C have effect as if any reference in those sections to section 24 were a reference to subsection (4) of this section; and
(c) sections 24D and 24E do not apply.

(4B) If, on an application under subsection (3), goods have been listed without consideration of the matters mentioned in paragraphs 26(1)(c) to (m), the Secretary may, if he or she thinks it appropriate, give the person in relation to whom the goods are listed written notice that the Secretary intends to determine whether the goods should continue to be listed.

(4C) If notice is given under subsection (4B), section 26 applies as if the person in relation to whom the goods are listed were an applicant for the listing of the goods.

(5) Section 21 does not apply, during the period of 15 months after the commencement of this Act or during such longer period as the Secretary specifies by notice published in the Gazette before the end of that first-mentioned period, to any goods.

(6) Where a person suffers any kind of loss, damage or injury caused by, or arising out of, the use by the person of therapeutic goods to which this section applies, no liability in respect of that loss, damage or injury attaches to the Commonwealth, the Secretary or any delegate of the Secretary.

67 Transitional provision for therapeutic goods for export only

Section 20 does not apply, during the period of 6 months after the commencement of this Act, to therapeutic goods manufactured in Australia solely for export from Australia.

68 Transitional arrangements for Part 3-3

(1) This section applies to a step in the manufacture of therapeutic goods in relation to a person in relation to premises in Australia if, before the commencement of this Act, the person was carrying out that step in relation to goods of that kind at those premises.
Section 69

(2) Where:
   (a) this section applies to a step in the manufacture of therapeutic goods in relation to a person in relation to premises; and
   (b) the Secretary is not aware of the person having been convicted of an offence against a law of the Commonwealth, of a State or of an internal Territory in respect of goods of that kind during the period of 2 years ending on the commencement of this Act;
subsection 35(1) does not apply the carrying out of that step by the person in relation to goods of that kind at those premises during the period of 4 months after that commencement.

(3) Where:
   (a) this section applies to a step in the manufacture of therapeutic goods in relation to a person in relation to premises; and
   (b) the person makes an application for a licence to carry out that step in relation to goods of that kind at those premises in accordance with section 37 and within 4 months after the commencement of this Act;
subsection 35(1) does not apply to the carrying out of that step by the person in relation to goods of that kind at those premises until the application is determined.

69 Continuation of standards and requirements

Any standards that were in force immediately before the commencement of this Act under Part 2 of the former Act, and any requirements that were in force at that time under section 15 of the former Act, continue in force as if they were standards made under Part 3-1 of this Act.
Schedule—Consequential Amendments

Section 62

Note:
The amendments made by this Schedule are incorporated in the compilations on ComLaw.

_Agricultural and Veterinary Chemicals Act 1988_
[repealed by Act No. 36, 1964, s. 24]

/Commonwealth Serum Laboratories Act 1961/

_Sea Installations Act 1987_

For access to the wording of the amendments made by this Schedule, _see_ Act No. 21, 1990.
Notes to the *Therapeutic Goods Act 1989*

Note 1

The *Therapeutic Goods Act 1989* as shown in this compilation comprises Act No. 21, 1990 amended as indicated in the Tables below.

All relevant information pertaining to application, saving or transitional provisions prior to 30 May 2000 is not included in this compilation. For subsequent information see Table A.

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<td>S. 2(6) [see Table A]</td>
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<td>Sch. 1 (items 38, 46, 55) [see Table A]</td>
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Notes to the Therapeutic Goods Act 1989

Act Notes

(a) The Therapeutic Goods Act 1989 was amended by Part 8 (sections 78–81) only of the Community Services and Health Legislation Amendment Act (No. 2) 1990, subsection 2(6) of which provides as follows:

(6) Part 8 commences immediately after the commencement of the Therapeutic Goods Act 1989.


(b) The Therapeutic Goods Act 1989 was amended by sections 82–88 only of the Health, Housing and Community Services Legislation Amendment Act 1992, subsection 2(1) of which provides as follows:

(1) Subject to this section, this Act commences on the day on which it receives the Royal Assent.

(c) The Therapeutic Goods Act 1989 was amended by the Customs, Excise and Bounty Legislation Amendment Act 1995, subsections 2(1) and (5) of which provide as follows:

(1) Subject to subsections (2), (3), (4), (5) and (6), this Act commences on the day on which it receives the Royal Assent.

(5) Schedules 2 and 3, items 1, 26 to 45, 49 to 53 and 67 of Schedule 4, Schedule 6, items 6 to 11 of Schedule 7 and Schedules 8 and 10 commence on 1 July 1995.

(d) The Therapeutic Goods Act 1989 was amended by Schedule 2 (item 1249) only of the Audit (Transitional and Miscellaneous) Amendment Act 1997, subsection 2(2) of which provides as follows:

(2) Schedules 1, 2 and 4 commence on the same day as the Financial Management and Accountability Act 1997.

(e) Subsection 2(3) of the Therapeutic Goods Legislation Amendment Act 1999 provides as follows:

(3) Schedule 2 is taken to have commenced immediately after the commencement of Part 2 of Schedule 1 to the Therapeutic Goods Amendment Act 1997.

Part 2 of Schedule 1 commenced on 1 January 1999 (see Gazette 1998, No. S609).

(f) The Therapeutic Goods Act 1989 was amended by Schedule 1 (items 936–938) only of the Public Employment (Consequential and Transitional) Amendment Act 1999, subsections 2(1) and (2) of which provide as follows:

(1) In this Act, commencing time means the time when the Public Service Act 1999 commences.

(2) Subject to this section, this Act commences at the commencing time.

(g) The Therapeutic Goods Act 1989 was amended by Schedule 3 (item 8) only of the Australia New Zealand Food Authority Amendment Act 2001, subsections 2(1)(a), (2) and (5) of which provide as follows:

(1) The following provisions commence on the day on which this Act receives the Royal Assent:

(a) sections 1, 2 and 3;

(2) Part 1 of Schedule 1 (other than item 120A), and Schedule 3, commence on the first day after the commencement of this section on which an amendment of the Australia New Zealand Joint Food Standards Agreement comes into force in accordance with Article 10 of that Agreement.

(5) As soon as practicable after the commencement of the following provisions:

(a) Part 1 of Schedule 1 (other than item 120A);

(b) Schedule 3;

the Minister must cause to be published in the Gazette a notice specifying the day on which those provisions commenced.
Notes to the  *Therapeutic Goods Act 1989*

**(h)** Subsection 2(1) (item 3) of the *Therapeutic Goods Amendment (Medical Devices) Act 2002* provides as follows:

> (1) Each provision of this Act specified in column 1 of the table commences, or is taken to have commenced, on the day or at the time specified in column 2 of the table.

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<th>Provision(s)</th>
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<tr>
<td>3. Schedule 2</td>
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**(i)** Subsection 2(1) (items 2(b), 4, 6 and 7) of the *Therapeutic Goods and Other Legislation Amendment Act 2002* provides as follows:

> (1) Each provision of this Act specified in column 1 of the table commences, or is taken to have commenced, on the day or at the time specified in column 2 of the table.

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<td>2. Schedule 1, items 6 and 7</td>
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<td>4. Schedule 3, item 4</td>
<td>Immediately after item 22 of Schedule 3 commences, subject to subsection (3)</td>
<td>4 October 2002</td>
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<td>6. Schedule 3, item 22</td>
<td>Immediately after Schedule 1 to the <em>Therapeutic Goods Amendment (Medical Devices) Act 2002</em> commences, subject to subsection (3)</td>
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<td>7. Schedule 4, item 1</td>
<td>Immediately after Schedule 1 to the <em>Therapeutic Goods Amendment (Medical Devices) Act 2002</em> commences</td>
<td>4 October 2002</td>
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**(j)** Subsection 2(1) (item 4) of the *Therapeutic Goods Amendment Act (No. 1) 2003* provides as follows:

> (1) Each provision of this Act specified in column 1 of the table commences, or is taken to have commenced, on the day or at the time specified in column 2 of the table.

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ad. = added or inserted  am. = amended  rep. = repealed  rs. = repealed and substituted

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Notes to the *Therapeutic Goods Act 1989*

**Note 2**

The following amendments commence on 4 October 2007:

**Schedule 2**

1. **Subsection 3(1) (definition of *gazetted therapeutic devices group*)**
   Repeal the definition.

2. **Subsection 3(1) (paragraph (b) of the definition of *grouped therapeutic goods*)**
   Repeal the paragraph.

3. **Subsection 3(1) (definition of *listable devices*)**
   Repeal the definition.

4. **Subsection 3(1) (definition of *medicine*)**
   Repeal the definition, substitute:
   
   *medicine* means therapeutic goods that are represented to achieve, or are likely to achieve, their principal intended action by pharmacological, chemical, immunological or metabolic means in or on the body of a human or animal.

5. **Subsection 3(1) (definition of *therapeutic device*)**
   Repeal the definition.

6. **Paragraph 7B(2)(b)**
   Omit “or therapeutic devices”.

7. **Chapter 3 (note following the heading)**
   Repeal the note.

8. **Section 10A**
   Omit “unless Part 3-2 applies to the device”.

282   *Therapeutic Goods Act 1989*
9 Section 15A
Repeal the section, substitute:

15A Application of this Part to medical devices
This Part does not apply to a medical device.

10 Subsection 16(2)
Omit “or therapeutic devices”.

11 Subsection 16(3)
Repeal the subsection.

12 Paragraphs 20(2)(a) and (b)
Repeal the paragraphs, substitute:

(a) the registration number or listing number of the goods is set out on the label of the goods in the prescribed manner; or

(b) in the case of an importation, that number is so set out, or is to be so set out before the goods are supplied in Australia.

13 Section 21
Omit “(other than listable devices)”.

14 Subparagraph 25(2)(a)(i)
Omit “if the goods are not therapeutic devices”, substitute “if the goods are medicines”.

15 Paragraph 25A(2)(a)
Repeal the paragraph, substitute:

(a) the information was given to the Secretary in relation to an application to register therapeutic goods (the new goods) consisting of, or containing, an active component; and

16 Section 25B
Repeal the section.

17 Subsection 26(1)
Omit “, subject to section 26AA,”.
Note 2

18 Paragraph 26(1)(g)
   Omit “not being therapeutic devices other than devices”, substitute “being medicines or other therapeutic goods”.

19 Subparagraph 26(2)(a)(i)
   Omit “if the goods are not therapeutic devices”, substitute “if the goods are medicines”.

20 Section 26AA
   Repeal the section.

21 Chapter 4 (note following the heading)
   Repeal the note.

22 Section 41BJ
   Repeal the section.

23 Subsection 60(1) (paragraph (a) of the definition of *initial decision*)
   Omit “under the definition of *therapeutic devices* in subsection 3(1) or”.

As at 4 March 2005 the amendments are not incorporated in this compilation.
Table A

Application, saving or transitional provisions

Therapeutic Goods Amendment Act (No. 2) 2000 (No. 56, 2000)

Schedule 1

5 Application

The amendment of section 30 of the Therapeutic Goods Act 1989 by this Schedule applies to directions given, and requirements made, of persons after the commencement of the amendment.

Therapeutic Goods Amendment Act (No. 3) 2000 (No. 120, 2000)

Schedule 1

4 Saving

(1) Regulations in force for the purposes of subsection 18(1) of the Therapeutic Goods Act 1989 immediately before the commencement of the amendment of that subsection made by this Schedule have effect, after that commencement, as if they had been made for the purposes of that subsection after that commencement.

(2) Subitem (1) does not prevent the amendment or repeal of the regulations.

6 Application

Conditions specified in regulations made for the purposes of subsection 19(1A) of the Therapeutic Goods Act 1989 apply to an approval for the purpose mentioned in paragraph 19(1)(b) of that Act granted before or after the commencement of subsection 19(1A).

8 Application

Conditions specified in regulations made for the purposes of subsection 19(4A) of the Therapeutic Goods Act 1989 apply to an approval for the purpose mentioned in paragraph 19(1)(b) of that Act granted before or after the commencement of subsection 19(4A).
Table A

10 Application

Subsection 19(5B) of the Therapeutic Goods Act 1989 applies to an authority given under subsection 19(5) of that Act before or after the commencement of subsection 19(5B).

17 Application

(1) Subsections 31B(1) and (2) of the Therapeutic Goods Act 1989 apply to an approval granted under subsection 19(1) of that Act before or after the commencement of section 31B of that Act.

(2) Subsection 31B(3) of the Therapeutic Goods Act 1989 applies to an authority given under subsection 19(5) of that Act before or after the commencement of section 31B of that Act.

Therapeutic Goods Amendment Act 2001 (No. 14, 2001)

Schedule 1

36 Application of amendments

(1) The amendments made by this Schedule do not apply, and the Therapeutic Goods Act 1989 as in force immediately before the commencement of this Schedule continues to apply, in relation to any application made under that Act, before that commencement, for listing of therapeutic goods.

(2) However, the amendment made by item 5 of this Schedule applies, after that commencement, to therapeutic goods that were listed before that commencement.

(3) The amendment made by item 31 of this Schedule applies, after that commencement, to medicines listed under section 26A before that commencement.
Table A

**Australia New Zealand Food Authority Amendment Act 2001** (No. 81, 2001)

2(6) In this section:


**Health and Aged Care Legislation Amendment (Application of Criminal Code) Act 2001** (No. 111, 2001)

4 Application of amendments

(1) Each amendment made by this Act applies to acts and omissions that take place after the amendment commences.

(2) For the purposes of this section, if an act or omission is alleged to have taken place between 2 dates, one before and one on or after the day on which a particular amendment commences, the act or omission is alleged to have taken place before the amendment commences.

**Therapeutic Goods Amendment (Medical Devices) Act 2002** (No. 24, 2002)

**Schedule 1**

38 Transitional for publication of list of goods on Register

The first list published under section 9E of the *Therapeutic Goods Act 1989* after the commencement of this Schedule must be published within 12 months after the last list published under section 33 of the *Therapeutic Goods Act 1989*.

46 Saving of existing registrations and listings

(1) Despite the repeal of section 17 of the *Therapeutic Goods Act 1989* by item 45, any therapeutic goods that, immediately before the
Table A

commencement of this Schedule, were included in the Register are taken to be included in the Register after that commencement.

(2) Despite the repeal of subsection 17(5) of the Therapeutic Goods Act 1989, any notices that were in force immediately before the commencement of this Schedule, are taken to have been made under section 9A after that commencement.

55 Saving of existing requests for inspection or variation of the Register

(1) A request for a copy of an entry in the Register under section 32 of the Therapeutic Goods Act 1989:
   (a) that was made before the commencement of this Schedule; and
   (b) that was not dealt with before that commencement; is taken, after that commencement, to be a request made under section 9C of the Therapeutic Goods Act 1989 as in force after that commencement.

(2) A request for a variation of an entry in the Register under section 32 of the Therapeutic Goods Act 1989:
   (a) that was made before the commencement of this Schedule; and
   (b) that was not dealt with before that commencement; is taken, after that commencement, to be a request made under section 9D of the Therapeutic Goods Act 1989 as in force after that commencement.

Therapeutic Goods and Other Legislation Amendment Act 2002 (No. 56, 2002)

Schedule 3

20 Application

(1) The amendment made by item 12 applies in relation to applications made on or after the commencement of that item.

(2) The amendment made by item 18 applies in relation to applications for licences made on or after the commencement of that item.
(3) Paragraph 40(4)(a) of the Therapeutic Goods Act 1989 as amended by this Schedule applies in relation to a request made under that paragraph on or after the commencement of this item, even if the licence to which the request relates was granted before that commencement.

21 Transitional

(1) An approval of a body as an approved conformity assessment body that was in force under the Therapeutic Goods Act 1989 immediately before the commencement time has effect after the commencement time as if it were an approval of the body by the Secretary in writing for the purposes of the definition of EC/EFTA attestation of conformity in subsection 3(1) of the Therapeutic Goods Act 1989 as in force after the commencement time.

(2) In subitem (1):

   commencement time means the time at which this item commences.

Therapeutic Goods Amendment Act (No. 1) 2003 (No. 39, 2003)

Schedule 1

41 Saving provisions

(1) The repeal of subsections 30(6) and (7) of the Therapeutic Goods Act 1989 does not affect the application, after the commencement of this item, of paragraph 30(6)(b) and subsection 30(7) of that Act in relation to a requirement imposed under paragraph 30(6)(a) of that Act before that commencement.

(2) The repeal of section 30A of the Therapeutic Goods Act 1989 does not affect the application, after the commencement of this item, of subsections 30A(3) and (4) of that Act in relation to a requirement imposed under subsection 30A(2) of that Act before that commencement.

(3) The repeal of section 30B of the Therapeutic Goods Act 1989 does not affect the application, after the commencement of this item, of subsections 30B(3), (4) and (5) of that Act in relation to a requirement imposed under subsection 30B(2) of that Act before that commencement.
Table A

55 Saving provision
The repeal and substitution of subsection 40(1) of the *Therapeutic Goods Act 1989* does not affect the application, after the commencement of this item, of any condition that:

(a) was imposed on a licence under that subsection; and
(b) was in force immediately before that commencement.

60 Application provision
Subsection 40(4) of the *Therapeutic Goods Act 1989* as amended by this Act, and subsection 40(5) of that Act, apply to any licence in force after the commencement of this item, whether or not the licence was granted after that commencement.

US Free Trade Agreement Implementation Act 2004 (No. 120, 2004)

Schedule 7

7 Application of amendments
(1) The amendments made by this Schedule apply to applications for registration or listing under section 23 of the *Therapeutic Goods Act 1989* made on or after the day on which this Schedule commences.

(2) The amendments made by item 6 apply to legal proceedings commenced on or after the day on which this Schedule commences.

Financial Framework Legislation Amendment Act 2005 (No. 8, 2005)

4 Saving of matters in Part 2 of Schedule 1
(1) If:

(a) a decision or action is taken or another thing is made, given or done; and
(b) the thing is taken, made, given or done under a provision of a Part 2 Act that had effect immediately before the commencement of this Act;
then the thing has the corresponding effect, for the purposes of the Part 2 Act as amended by this Act, as if it had been taken, made, given or done under the Part 2 Act as so amended.

(2) In this section:

Part 2 Act means an Act that is amended by an item in Part 2 of Schedule 1.

Schedule 1

496 Saving provision—Finance Minister’s determinations

If a determination under subsection 20(1) of the Financial Management and Accountability Act 1997 is in force immediately before the commencement of this item, the determination continues in force as if it were made under subsection 20(1) of that Act as amended by this Act.