Health Act 1993
A1993-13

Republication No 40
Effective: 2 July 2019

Republication date: 2 July 2019

Last amendment made by A2018-42
(republication for expiry of transitional provisions (pt 23))
About this republication

The republished law

This is a republication of the Health Act 1993 (including any amendment made under the Legislation Act 2001, part 11.3 (Editorial changes)) as in force on 2 July 2019. It also includes any commencement, amendment, repeal or expiry affecting this republished law to 2 July 2019. The legislation history and amendment history of the republished law are set out in endnotes 3 and 4.

Kinds of republications

The Parliamentary Counsel’s Office prepares 2 kinds of republications of ACT laws (see the ACT legislation register at www.legislation.act.gov.au):

• authorised republications to which the Legislation Act 2001 applies
• unauthorised republications.

The status of this republication appears on the bottom of each page.

Editorial changes

The Legislation Act 2001, part 11.3 authorises the Parliamentary Counsel to make editorial amendments and other changes of a formal nature when preparing a law for republication. Editorial changes do not change the effect of the law, but have effect as if they had been made by an Act commencing on the republication date (see Legislation Act 2001, s 115 and s 117). The changes are made if the Parliamentary Counsel considers they are desirable to bring the law into line, or more closely into line, with current legislative drafting practice.

This republication does not include amendments made under part 11.3 (see endnote 1).

Uncommenced provisions and amendments

If a provision of the republished law has not commenced, the symbol \[U\] appears immediately before the provision heading. Any uncommenced amendments that affect this republished law are accessible on the ACT legislation register (www.legislation.act.gov.au). For more information, see the home page for this law on the register.

Modifications

If a provision of the republished law is affected by a current modification, the symbol \[M\] appears immediately before the provision heading. The text of the modifying provision appears in the endnotes. For the legal status of modifications, see the Legislation Act 2001, section 95.

Penalties

At the republication date, the value of a penalty unit for an offence against this law is $160 for an individual and $810 for a corporation (see Legislation Act 2001, s 133).
# Health Act 1993

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Health Act 1993

An Act relating to the provision of health services
Part 1 Preliminary

Section 1

Part 1 Preliminary

1 Name of Act

This Act is the Health Act 1993.

2 Dictionary

The dictionary at the end of this Act is part of this Act.

Note 1 The dictionary at the end of this Act defines certain terms used in this Act, and includes references (signpost definitions) to other terms defined elsewhere in this Act.

For example, the signpost definition ‘health facility—see section 6.’ means that the term ‘health facility’ is defined in that section.

Note 2 A definition in the dictionary (including a signpost definition) applies to the entire Act unless the definition, or another provision of the Act, provides otherwise or the contrary intention otherwise appears (see Legislation Act, s 155 and s 156 (1)).

3 Notes

A note included in this Act is explanatory and is not part of this Act.

Note See Legislation Act, s 127 (1), (4) and (5) for the legal status of notes.
4 Offences against Act—application of Criminal Code etc

Other legislation applies in relation to offences against this Act.

Note 1 Criminal Code

The [Criminal Code](#), ch 2 applies to all offences against this Act (see Code, pt 2.1).

The chapter sets out the general principles of criminal responsibility (including burdens of proof and general defences), and defines terms used for offences to which the Code applies (e.g. conduct, intention, recklessness and strict liability).

Note 2 Penalty units

The [Legislation Act](#), s 133 deals with the meaning of offence penalties that are expressed in penalty units.
Part 2  Important concepts

Section 5

Part 2  Important concepts

5  What is a health service?
For this Act, a health service is a service provided to someone (the service user) for any of the following purposes:

(a) assessing, recording, maintaining or improving the physical, mental or emotional health, comfort or wellbeing of the service user;

(b) diagnosing, treating or preventing an illness, disability, disorder or condition of the service user.

6  What is a health facility?
In this Act:

health facility means the following facilities where health services are provided:

(a) a hospital, including a day hospital;

(b) a hospice;

(c) a nursing home;

(d) a health practitioner’s consulting room;

(e) another facility ordinarily used by the Territory to provide health services;

(f) any other facility prescribed by regulation for this section.

7  Who is a health service provider?
In this Act:

health service provider—

(a) means a health practitioner or other person who provides a health service; and
(b) for a health facility, means a health service provider who—

(i) provides a health service at the health facility; or

(ii) uses the equipment or other facilities of the health facility to provide a health service elsewhere.

**Examples of people who may be health service providers**

1. a chiropractor
2. a dentist
3. a dental technician
4. a dental prosthetist
5. a doctor
6. a nurse
7. an osteopath
8. an optometrist
9. a pharmacist
10. a physiotherapist
10 **Objectives**

In providing health services the Territory must have regard to the following objectives:

(a) to improve the efficiency, effectiveness and quality of health services;

(b) to guarantee equitable access to and participation in health services and to ensure that language and cultural differences are not barriers to such access or participation;

(c) to maintain a strong and viable public hospital system and a full range of community health services;

(d) to support worker and community participation in the development of policies for the delivery of health services;

(e) to ensure that the community is aware of the range of health services that is available and that patients have information that is sufficient to enable them to make informed choices;

(f) to foster disease prevention and primary health care;

(g) to cooperate with community groups in the provision of health services.
11 Medicare principles and commitments

(1) The following guidelines govern the delivery of public hospital services to eligible persons in the ACT:

Note The guidelines focus on the provision of public hospital services to eligible persons, but operate in an environment where eligible persons have the right to choose private health care in public and private hospitals supported by private health insurance.

(a) eligible persons must be given the choice to receive public hospital services free of charge as public patients;

Note 1 Hospital services include in-patient, outpatient, emergency services (including primary care where appropriate) and day patient services consistent with currently acceptable medical and health service standards.

Note 2 At the time of admission to a hospital, or as soon as practicable after that, an eligible person will be required to elect or confirm whether he or she wishes to be treated as a public or private patient.

(b) access to public hospital services is to be on the basis of clinical need;

Note 1 None of the following factors are to be a determinant of an eligible person’s priority for receiving hospital services:

(a) whether or not an eligible person has health insurance;
(b) an eligible person’s financial status or place of residence;
(c) whether or not an eligible person intends to elect, or elects, to be treated as a public or private patient.

Note 2 This guideline applies equally to waiting times for elective surgery.

(c) to the maximum practicable extent, the Territory will ensure the provision of public hospital services equitably to all eligible persons, regardless of their geographical location;

Note 1 This guideline does not require a local hospital to be equipped to provide eligible persons with every hospital service they may need.

Note 2 In rural and remote areas, the Territory should ensure provision of reasonable public access to a basic range of hospital services that are in accord with clinical practices.
Part 3  
Health care principles

Section 12

(d) the Commonwealth and the Territory must make available information on the public hospital services eligible persons can expect to receive as public patients;

Note 1 The joint Commonwealth/Territory development of a Public Patients Hospital Charter for the Territory will be a vehicle for the public dissemination of this information.

Note 2 The Charter will set out the public hospital services available to public patients.

(e) the Commonwealth and the Territory are committed to making improvements in the efficiency, effectiveness and quality of hospital service delivery.

Note This includes a commitment to quality improvement, outcome measurement, management efficiency and effort to integrate the delivery of hospital and other health and community services.

(2) A word or expression used in the Medicare Agreements Act 1992 (Cwlth) has the same meaning in subsection (1).

12 Legal effect

Nothing in this part is to be taken to create any legal rights not in existence before the enactment of this part or to affect any legal rights in existence before that enactment or that would, apart from this part, have come into existence after that enactment.
Part 4  Quality assurance

Division 4.1  Quality assurance—important concepts

20  Definitions—pt 4

In this part:

CEO—

(a) of a health facility—see section 22; and

(b) of a health professional organisation—see section 23.

health facility QAC, for a health facility, means a committee approved under section 25 as a quality assurance committee for the health facility.

health professional organisation—see section 21.

health professional organisation QAC, for a health professional organisation, means a committee approved under section 26 as a quality assurance committee for the health professional organisation.

health service report—see section 38.

ministerial report—see section 41.

special purpose QAC means a committee approved under section 27.

Note  Quality assurance committee is defined for the Act in s 24.

21  What is a health professional organisation?

In this part:

health professional organisation means an entity that—

(a) is an association, society, college, faculty or other body of professionals who provide a health service; and

(b) is prescribed by regulation for this section.
22 Who is the CEO of a health facility?

In this part:

CEO, of a health facility, means—

(a) for a health facility operated by the Territory—the director-general; or

(b) in any other case—the person with overall responsibility for the control of the health facility.

23 Who is the CEO of a health professional organisation?

In this part:

CEO, of a health professional organisation, means the person with overall responsibility for the control of the health professional organisation.

Division 4.2 Quality assurance—quality assurance committees

24 What is a quality assurance committee?

In this Act:

quality assurance committee means—

(a) a health facility QAC; or

(b) a health professional organisation QAC; or

(c) a special purpose QAC.
25 Approval of health facility QACs

(1) The Minister may approve a stated committee as a quality assurance committee for a stated health facility.

(2) An approval is a notifiable instrument.

Note A notifiable instrument must be notified under the Legislation Act.

26 Approval of health professional organisation QACs

(1) The Minister may approve a stated committee as a quality assurance committee for a stated health professional organisation.

(2) An approval is a notifiable instrument.

Note A notifiable instrument must be notified under the Legislation Act.

27 Approval of special purpose QACs

(1) The Minister may approve a stated committee as a quality assurance committee for a stated purpose.

(2) An approval is a notifiable instrument.

Note A notifiable instrument must be notified under the Legislation Act.

27A Quality Assurance Committees—term

The Minister may not approve a committee under section 25, section 26 or section 27 for a term longer than 3 years.

28 Quality assurance committees—criteria for approval

The Minister may approve a committee as a quality assurance committee under section 25, section 26 or section 27 only if satisfied that—

(a) the committee’s functions would be facilitated by the members, and other people mentioned in section 34, being protected from liability under section 34 (Quality assurance committees—protection of members etc from liability); and
(b) it is in the public interest for part 8 (Secrecy) to apply to information held by the committee members.

29  Quality assurance committees—revocation of approval

The Minister may revoke the approval of a committee as a quality assurance committee if—

(a) the Minister is not satisfied about 1 or both of the criteria mentioned in section 28 in relation to the committee; or

(b) the committee has failed to prepare a health service report as required under section 38; or

(c) the committee has failed to give a health service report as required under section 39; or

(d) the committee has failed to prepare, or give, a ministerial report as required under section 41 (Annual quality assurance committee report to Minister); or

(e) the committee has failed to prepare a report as required by a regulation made under section 42 (Other quality assurance reports); or

(f) for the last year, none of the members of the committee has held sensitive information in the exercise of a function under this Act.

Note 1  Sensitive information is defined in s 124.

Note 2  Power to make a statutory instrument includes power to amend or repeal the instrument. The power to amend or repeal the instrument is exercisable in the same way, and subject to the same conditions, as the power to make the instrument (see Legislation Act, s 46).
30 **Quality assurance committees—functions**

A quality assurance committee has the following functions:

(a) to facilitate the improvement of health services provided in the ACT;

(b) any other function given to the committee under this Act.

31 **Quality assurance committees—appointment of members**

(1) The CEO of a health facility must appoint the members of a health facility QAC for the health facility.

(2) The CEO of a health professional organisation must appoint the members of a health professional organisation QAC for the health professional organisation.

(3) The director-general must appoint the members of a special purpose QAC.

*Note 1* For the making of appointments (including acting appointments), see the *Legislation Act*, pt 19.3.

*Note 2* In particular, an appointment may be made by naming a person or nominating the occupant of a position (see *Legislation Act*, s 207).

*Note 3* A person may be reappointed to a position if the person is eligible to be appointed to the position (see *Legislation Act*, s 208 and dict, pt I, def appoint).

32 **Quality assurance committees—disclosure of interests**

(1) Section 190 (Disclosure of interests by committee members) applies to quality assurance committees.

(2) If a person acting under the direction of a quality assurance committee has a material interest in an issue being considered, or about to be considered, by the committee, the person must disclose the nature of the interest at a committee meeting as soon as practicable after the relevant facts come to the person’s knowledge.
(3) In this section:

material interest—see section 190 (4).

33 Quality assurance committees—procedure

In exercising its functions, a quality assurance committee—

(a) must comply with the rules of natural justice; and

(b) is not bound by the rules of evidence but may inform itself of anything in the way it considers appropriate; and

(c) may do whatever it considers necessary or convenient for the fair and prompt conduct of its functions.

34 Quality assurance committees—protection of members etc from liability

(1) In this section:

relevant person, for a quality assurance committee—

(a) means a person who is, or has been, a member of the committee; and

(b) includes anyone engaging in conduct under the direction of a person who is a member of the committee.

(2) A relevant person for a quality assurance committee is not personally liable for anything done or omitted to be done honestly and without recklessness—

(a) in the exercise of a function under this Act; or

(b) in the reasonable belief that the act or omission was in the exercise of a function under this Act.

Note A reference to an Act includes a reference to the statutory instruments made or in force under the Act, including any regulation (see Legislation Act, s 104).
(3) Any civil liability that would, apart from this section, attach to a relevant person for a quality assurance committee attaches instead to—

(a) if the committee is a health facility QAC for a health facility—the health facility; or

(b) if the committee is a health professional organisation QAC for a health professional organisation—the health professional organisation; or

(c) if the committee is a special purpose QAC—the Territory.

35 Quality assurance committees—obtaining information

(1) A quality assurance committee carrying out a function under this Act may ask anyone to give the committee information, including protected information, that is relevant to the committee carrying out the function.

Note The identity of a person who gives information to a committee under this section is protected (see pt 8).

(2) When asking anyone for information, the committee must tell the person that giving false or misleading information is an offence against the Criminal Code, section 338 (Giving false or misleading information).

(3) If someone gives information honestly and without recklessness to a quality assurance committee under subsection (1)—

(a) the giving of the information is not—

(i) a breach of confidence; or

(ii) a breach of professional etiquette or ethics; or

(iii) a breach of a rule of professional conduct; and

(b) the person does not incur civil or criminal liability only because of the giving of the information.
Division 4.3  
Assessment and evaluation of health services

36  
Assessment and evaluation of health services

(1) A health facility QAC for a health facility may assess and evaluate health services provided by health service providers for the health facility by carrying out a quality assurance activity with the health service providers.

(2) A health professional organisation QAC for a health professional organisation may assess and evaluate health services provided by health service providers who are members of a health professional organisation by carrying out a quality assurance activity with the health service providers.

(3) A special purpose QAC may, for a purpose for which it was approved, assess and evaluate health services provided by health service providers for any health facility by carrying out a quality assurance activity with the health service providers.

(4) In this section:

*quality assurance activity* means an activity approved as a quality assurance activity under section 37.
37 Approval of quality assurance activities

(1) The Minister may approve an activity as a quality assurance activity if satisfied that the activity is designed to evaluate, monitor or improve the quality of a health service.

(2) An approval is a notifiable instrument.

Note A notifiable instrument must be notified under the Legislation Act.

Examples of activities designed to evaluate, monitor or improve the quality of a health service
1. clinical audits
2. records audits
3. peer review
4. quality review
5. investigation into disease and death.

38 Preparing health service reports

(1) This section applies to a quality assurance committee if it completes an assessment and evaluation under section 36.

(2) The quality assurance committee must prepare a report (a health service report) about the assessment and evaluation.

Note The report must be prepared as soon as possible (see Legislation Act, s 151B).

(3) The health service report must include the following:
(a) details of the health services assessed and evaluated;
(b) the results of the assessment and evaluation;
(c) the committee’s conclusions;
(d) the committee’s recommendations (if any).
38A Extraordinary reports

(1) This section applies if—

(a) a quality assurance committee is assessing and evaluating health services under section 36; and

(b) the quality assurance committee becomes aware of something that is sufficiently serious to require urgent action to prevent or limit any adverse effect it might have on the health service.

(2) The quality assurance committee must report the thing to the director-general as soon as possible, even if the committee has not completed the assessment and evaluation.

(3) Subsection (2) applies even if the thing is not related to the quality assurance activity the committee is carrying out.

(4) A report under subsection (2) must be in writing and may include sensitive information.

Note Sensitive information—see s 124.

38B Interim reports

(1) The director-general may ask a quality assurance committee to prepare a report on its activities before it completes an assessment and evaluation under section 36.

(2) A report prepared in response to a request under subsection (1) must include the following:

(a) details of the health services that are being assessed and evaluated;

(b) details of how the assessment and evaluation is progressing;

(c) details of any conclusions the committee may have reached;

(d) the committee’s recommendations (if any).
(3) A report under subsection (1) must be in writing and may include sensitive information.

Note Sensitive information—see s 124.

39 Giving health service reports to CEO or director-general

(1) This section applies to a quality assurance committee if it prepares a health service report.

(2) The quality assurance committee must give a copy of the report to—

(a) if the committee is a health facility QAC for a health facility—the CEO of the health facility; or

(b) if the committee is a health professional organisation QAC for a health professional organisation—the CEO of the health professional organisation; or

(c) if the committee is a special purpose QAC—the director-general.

Note The report must be given as soon as possible (see Legislation Act, s 151B).

40 Monitoring implementation of recommendations

If a quality assurance committee makes a recommendation in a health service report, the committee may monitor the implementation of the recommendation.
Division 4.4 Quality assurance committees—reporting

41 Annual quality assurance committee report to Minister

(1) A quality assurance committee must, for each financial year, prepare a report (a ministerial report) about the committee’s operation during the year.

(2) The ministerial report must include information for the financial year about—

   (a) the committee’s functions under division 4.3 (Assessment and evaluation of health services); and

   (b) how the committee’s functions were facilitated by the members, and other people mentioned in section 34, being protected from liability under section 34 (Quality assurance committees—protection of members etc from liability); and

   (c) why it was in the public interest for part 8 (Secrecy) to apply to information held by the committee members.

(3) The ministerial report must comply with any requirements prescribed by regulation for this section.

(4) The ministerial report must not include sensitive information.

   *Note* Sensitive information is defined in s 124.

(5) The ministerial report must be given to the Minister not later than 3 months after the end of the financial year.
42 Other quality assurance committee reports

(1) A quality assurance committee must prepare a report prescribed by regulation for this section.

(2) The report must include the following information about the operation of the committee—

(a) how the committee’s functions were facilitated by the members, and other people mentioned in section 34, being protected from liability under section 34 (Quality assurance committees—protection of members etc from liability); and

(b) why it was in the public interest for part 8 (Secrecy) to apply to information held by the committee members.

(3) The report must not include sensitive information.

Note Sensitive information is defined in s 124.

Division 4.5 Quality assurance committees—information sharing

43 Quality assurance committees—giving information to the Coroner’s Court

A quality assurance committee may give protected information to the Coroner’s Court if the committee is satisfied that giving the information would be likely to facilitate the improvement of health services provided in the ACT.

Note Protected information includes sensitive information (see s 123).
44 Quality assurance committees—giving information to other quality assurance committees

A quality assurance committee may give protected information to another quality assurance committee if the committee is satisfied that giving the information would be likely to facilitate the improvement of health services provided in the ACT.

Note Protected information includes sensitive information (see s 123).

45 Quality assurance committees—giving information to health board and health services commissioner

(1) A quality assurance committee may give protected information to a health board if the committee is satisfied that giving the information would be likely to facilitate the improvement of health services provided in the ACT.

(2) If a quality assurance committee gives protected information to a health board under subsection (1), the committee must also give the information to the health services commissioner.

Note Protected information includes sensitive information (see s 123).

46 Quality assurance committees—giving information to Minister

A quality assurance committee may give protected information to the Minister if the committee is satisfied that giving the information would be likely to facilitate the improvement of health services provided in the ACT.

Note Protected information includes sensitive information (see s 123).

47 Quality assurance committees—admissibility of evidence

(1) The following are not admissible as evidence in a proceeding before a court:

(a) an oral statement made in a proceeding before a quality assurance committee;
Quality assurance committees—information sharing

(b) a document given to a quality assurance committee, but only to the extent that it was prepared only for the committee;

(c) a document prepared by a quality assurance committee.

(2) In this section:

court includes a tribunal, authority or person with power to require the production of documents or the answering of questions.
Part 5  Reviewing scope of clinical practice

Section 50

50 Definitions—pt 5

In this part:

CEO, of a health facility—see section 53.

Chief executive officer, Calvary means the person engaged to exercise the functions of the position of chief executive officer (however described) of Calvary Health Care ACT Limited (Public Division) under the rules of Calvary Health Care ACT Limited.

dentist, for a health facility—see section 52.

doctor, for a health facility—see section 52.

eligible midwife, for a health facility—see section 52.

hospital includes a day hospital.

review, in relation to scope of clinical practice—see section 55.

scope of clinical practice, of a doctor, dentist or eligible midwife for a health facility—see section 54.

scope of clinical practice executive decision notice—see section 70.

scope of clinical practice report—see section 67.

Note Scope of clinical practice committee is defined for the Act in s 51.

51 What is a scope of clinical practice committee?

In this Act:

scope of clinical practice committee means a committee approved under section 56 as a scope of clinical practice committee.
Who is a doctor, dentist or eligible midwife for a health facility?

In this Act:

dentist, for a health facility, means a dentist who—
(a) provides health services at the health facility; or
(b) uses the equipment or other facilities of the health facility to provide health services elsewhere.

doctor, for a health facility, means a doctor who—
(a) provides health services at the health facility; or
(b) uses the equipment or other facilities of the health facility to provide health services elsewhere.

eligible midwife, for a health facility, means a midwife who—
(a) is an eligible midwife within the meaning of the Health Insurance Act 1973 (Cwlth), section 21 (Meaning of eligible midwife); and
(b) either—
   (i) provides health services at the health facility; or
   (ii) uses the equipment or other facilities of the health facility to provide health services elsewhere.

Who is the CEO of a health facility?

In this part:

CEO, of a health facility, means—
(a) for a health facility operated by the Territory—the director-general; or
(b) in any other case—the person with overall responsibility for the control of the health facility.
Part 5  
Reviewing scope of clinical practice

Section 54

54 What is scope of clinical practice?
In this part:

scope of clinical practice, of a doctor, dentist or eligible midwife for a health facility, means the rights of the doctor, dentist or eligible midwife established by agreement between the doctor, dentist or eligible midwife and the health facility—

(a) to treat patients or carry out other procedures at the health facility; or

(b) to use the equipment or other facilities of the health facility.

55 Meaning of review scope of clinical practice
In this part:

review, in relation to the scope of clinical practice, includes assess and evaluate the scope of clinical practice.

56 Approval of scope of clinical practice committees
(1) The Minister may approve a committee as a scope of clinical practice committee in accordance with section 57.

(2) An approval is a notifiable instrument.

Note A notifiable instrument must be notified under the Legislation Act.

57 Scope of clinical practice committees—criteria for approval
The Minister may approve a committee as a scope of clinical practice committee under section 56 only if satisfied that—

(a) the committee’s functions would be facilitated by the members, and other people mentioned in section 63, being protected from liability under section 63 (Scope of clinical practice committees—protection of members etc from liability); and
(b) it is in the public interest for part 8 (Secrecy) to apply to information held by the committee members.

58 Scope of clinical practice committees—revocation of approval

The Minister may revoke the approval of a committee as a scope of clinical practice committee if the Minister is not satisfied about 1 or both of the criteria mentioned in section 57 in relation to the committee.

Note Power to make a statutory instrument includes power to amend or repeal the instrument. The power to amend or repeal the instrument is exercisable in the same way, and subject to the same conditions, as the power to make the instrument (see Legislation Act, s 46).

59 Scope of clinical practice committees—functions

(1) A scope of clinical practice committee has the following functions:

(a) to decide—

(i) whether to credential a doctor, dentist or eligible midwife for a health facility; and

(ii) the terms on which a doctor, dentist or eligible midwife is credentialled;

(b) to define, and review, the scope of clinical practice of a doctor, dentist or eligible midwife credentialled for a health facility;

(c) to review the scope of clinical practice of a doctor, dentist or eligible midwife if the CEO of a health facility refers the doctor’s, dentist’s or eligible midwife’s scope of clinical practice to the committee under section 69 (5);

(d) to immediately withdraw or amend the scope of clinical practice of a doctor, dentist or eligible midwife credentialled for a health facility in accordance with this Act;

(e) any other function given to the committee under this Act.
(2) A reference in this section to credentialling a doctor, dentist or eligible midwife includes re-credentialling the doctor, dentist or eligible midwife.

(3) A scope of clinical practice committee must, as far as practicable, exercise its functions under subsection (1) (a), (b) and (c) in accordance with the Standard.

(4) In this section:

credential, in relation to a doctor, dentist or eligible midwife, means endorse the doctor, dentist or eligible midwife (the practitioner) to provide health services based on verification and assessment of the practitioner’s qualifications, experience, skill, professional standing and any other relevant professional attributes.


60 Scope of clinical practice committees—appointment of members

The director-general must appoint the members of a scope of clinical practice committee.

Note 1 For the making of appointments (including acting appointments), see the Legislation Act, pt 19.3.

Note 2 In particular, an appointment may be made by naming a person or nominating the occupant of a position (see Legislation Act, s 207).

Note 3 A person may be reappointed to a position if the person is eligible to be appointed to the position (see Legislation Act, s 208 and dict, pt 1, def appoint).
61 Scope of clinical practice committees—disclosure of interests

(1) Section 190 (Disclosure of interests by committee members) applies to scope of clinical practice committees.

(2) If a person acting under the direction of a scope of clinical practice committee has a material interest in an issue being considered, or about to be considered, by the committee, the person must disclose the nature of the interest at a committee meeting as soon as practicable after the relevant facts come to the person’s knowledge.

(3) In this section:

material interest—see section 190 (4).

62 Scope of clinical practice committees—procedure

(1) In exercising its functions, a scope of clinical practice committee—

(a) must comply with the rules of natural justice; and

(b) is not bound by the rules of evidence but may inform itself of anything in the way it considers appropriate; and

(c) may do whatever it considers necessary or convenient for the fair and prompt conduct of its functions.

(2) A scope of clinical practice committee may, by resolution, determine the procedures for carrying out its functions.

63 Scope of clinical practice committees—protection of members etc from liability

(1) A relevant person for a scope of clinical practice committee is not personally liable for anything done or omitted to be done honestly and without recklessness—

(a) in the exercise of a function under this Act; or
(b) in the reasonable belief that the act or omission was in the exercise of a function under this Act.

Note A reference to an Act includes a reference to the statutory instruments made or in force under the Act, including any regulation (see Legislation Act, s 104).

(2) Any civil liability that would, apart from this section, attach to a relevant person for a scope of clinical practice committee attaches instead to the Territory.

(3) In this section:

relevant person, for a scope of clinical practice committee—

(a) means a person who is, or has been, a member of the committee; and

(b) includes anyone engaging in conduct under the direction of a person who is a member of the committee.

64 Scope of clinical practice committees—obtaining information

(1) A scope of clinical practice committee carrying out a function under this Act may ask anyone to give the committee information, including protected information, that is relevant to the committee carrying out the function.

Note The identity of a person who gives information to a committee under this section is protected (see pt 8).

(2) When asking anyone for information, the committee must tell the person that giving false or misleading information is an offence against the Criminal Code, section 338 (Giving false or misleading information).
(3) If someone gives information honestly and without recklessness to a scope of clinical practice committee under subsection (1)—
   (a) the giving of the information is not—
      (i) a breach of confidence; or
      (ii) a breach of professional etiquette or ethics; or
      (iii) a breach of a rule of professional conduct; and
   (b) the person does not incur civil or criminal liability only because of giving the information.

65 Scope of clinical practice committee must give doctor, dentist or eligible midwife opportunity to explain

(1) This section applies to a scope of clinical practice committee if—
   (a) the committee is reviewing the scope of clinical practice of a doctor, dentist or eligible midwife for a health facility; and
   (b) the committee proposes to recommend in a scope of clinical practice report that—
      (i) the scope of clinical practice of the doctor, dentist or eligible midwife should be amended or withdrawn; or
      (ii) the terms of engagement of the doctor, dentist or eligible midwife by the health facility should be amended; or
      (iii) the engagement of the doctor, dentist or eligible midwife by the health facility should be suspended or ended.

Note Scope of clinical practice reports are prepared under s 67.

(2) The committee must give the doctor, dentist or eligible midwife a written notice (a recommendation notice) stating—
   (a) the committee’s proposed recommendation; and
   (b) the reasons for the committee’s proposed recommendation; and
Part 5  
Reviewing scope of clinical practice

Section 66

(c) that the doctor, dentist or eligible midwife may, not later than 21 days after the day the recommendation notice is given to the doctor, dentist or eligible midwife, make a submission to the committee about the proposed recommendation.

(3) A recommendation notice must not include sensitive information.

Note  Sensitive information—see s 124.

(4) The committee must consider any submission made by the doctor, dentist or eligible midwife to the committee in accordance with the notice.

66  
Interim and emergency withdrawal or amendment of scope of clinical practice by committee

(1) If at any time a scope of clinical practice committee forms the view that the clinical practice of a doctor, dentist or eligible midwife at a health facility poses a threat to the safety of members of the public, the committee may withdraw or amend the scope of clinical practice of the doctor, dentist or eligible midwife with immediate effect.

(2) The scope of clinical practice committee may take action under subsection (1) before the completion of a review by the committee of the doctor’s, dentist’s or eligible midwife’s scope of clinical practice under section 65.

(3) Any withdrawal or amendment under this section has effect until a decision of the CEO of a health facility on the scope of clinical practice report in relation to the doctor, dentist or eligible midwife takes effect under section 71 (When CEO decision on scope of clinical practice report takes effect).

(4) If a scope of clinical practice committee withdraws or amends the scope of clinical practice of a doctor, dentist or eligible midwife under subsection (1), the committee must tell the director-general and the chief executive officer, Calvary (the executive officers) of the committee’s decision and the date of the decision, in writing, as soon as possible.
(5) If an executive officer is told about the withdrawal or amendment of the scope of clinical practice of a doctor, dentist or eligible midwife under this section, the executive officer must tell appropriate officers under their authority or direction of the committee’s decision so that proper effect can be given to the decision.

Examples—appropriate officers
- general manager of the health facility
- clinical unit director
- head of department at health facility
- immediate supervisor of doctor, dentist or eligible midwife
- human resource personnel

67 Preparing scope of clinical practice reports

(1) This section applies to a scope of clinical practice committee if—

(a) the committee has reviewed the scope of clinical practice of a doctor, dentist or eligible midwife for a health facility; and

(b) if the committee has given the doctor, dentist or eligible midwife a recommendation notice—the committee has considered any submission made by the doctor, dentist or eligible midwife in accordance with the notice; and

(c) the committee has completed the review.

(2) The scope of clinical practice committee must prepare a report (a scope of clinical practice report) about the review.

Note The report must be prepared as soon as possible (see Legislation Act, s 151B).

(3) The scope of clinical practice report must include the committee’s recommendations about whether—

(a) the scope of clinical practice of the doctor, dentist or eligible midwife should stay the same, be amended or be withdrawn; and
(b) the terms of engagement of the doctor, dentist or eligible midwife by the health facility should be amended; and

(c) the engagement of the doctor, dentist or eligible midwife by the health facility should be suspended or ended.

(4) In this section:

recommendation notice—see section 65 (2).

68 Giving scope of clinical practice reports to CEO of health facility and doctor, dentist or eligible midwife

If a scope of clinical practice committee prepares a scope of clinical practice report about a doctor, dentist or eligible midwife for a health facility, the committee must give a copy of the report to—

(a) the CEO of the health facility; and

(b) the doctor, dentist or eligible midwife.

Note The report must be given as soon as possible (see Legislation Act, s 151B).

69 CEO may make interim or emergency decision on scope of clinical practice

(1) If the CEO of a health facility has concerns about a doctor, dentist or eligible midwife for a health facility of sufficient seriousness to warrant the immediate amendment or withdrawal of the scope of clinical practice of the doctor, dentist or eligible midwife, the CEO may, by notice in writing, amend or withdraw the scope of clinical practice of the doctor, dentist or eligible midwife with immediate effect.

(2) The CEO may take action under subsection (1) even if a scope of clinical practice committee has not reported on, or is not currently investigating, the scope of clinical practice of the doctor, dentist or eligible midwife.
(3) Any amendment or withdrawal of the scope of clinical practice of a doctor, dentist or eligible midwife under this section has effect from the day and time the notice is given to the doctor, dentist or eligible midwife—

(a) if a scope of clinical practice report is prepared under section 67 in relation to the doctor, dentist or eligible midwife—until a decision on the scope of clinical practice report takes effect under section 71; or

(b) in any other case—until the CEO, by notice in writing, revokes the amendment or withdrawal.

(4) Subsection (5) applies if—

(a) the CEO amends or withdraws the scope of clinical practice of a doctor, dentist or eligible midwife under subsection (1); and

(b) the scope of clinical practice of the doctor, dentist or eligible midwife is not the subject of an investigation by a scope of clinical practice committee.

(5) The CEO must immediately refer the scope of clinical practice of the doctor, dentist or eligible midwife to a scope of clinical practice committee.

(6) If the CEO amends or withdraws the scope of clinical practice of a doctor, dentist or eligible midwife under subsection (1), the CEO must, in writing, notify—

(a) the doctor, dentist or eligible midwife; and

(b) if the CEO is not the director-general—the director-general; and

(c) if the CEO is not the chief executive officer, Calvary—the chief executive officer, Calvary; and

(d) the relevant health board for the doctor, dentist or eligible midwife; and

(e) the health services commissioner; and
(f) the CEO of any other health facility at which the doctor, dentist or eligible midwife is engaged; and

(g) if a scope of clinical practice committee submitted a report about the doctor, dentist or eligible midwife under section 68 to the CEO—the scope of clinical practice committee that submitted the report; and

(h) all appropriate officers under the CEO’s authority or direction of the committee’s decision so that proper effect can be given to the decision.

Examples—appropriate officers
- general manager of the health facility
- clinical unit director
- head of department at health facility
- immediate supervisor of doctor, dentist or eligible midwife
- human resource personnel

70 CEO must make decision on scope of clinical practice report

(1) This section applies if the CEO of a health facility is given a scope of clinical practice report about a doctor, dentist or eligible midwife for the health facility.

(2) The CEO must—

(a) consider the recommendations in the scope of clinical practice report; and

(b) decide whether to take—

(i) the action recommended in the scope of clinical practice report; or
(ii) any other action that the committee could have recommended under section 67 (3) that the CEO considers appropriate.

Note 1   The CEO must consider the recommendations and make a decision as soon as possible (see Legislation Act, s 151B).

Note 2   A decision of the CEO under this section is a reviewable decision (see pt 10).

(3) After the CEO has made a decision under subsection (2), the CEO must give the following people notice in writing (a scope of clinical practice executive decision notice) of the decision:

(a) each doctor, dentist or eligible midwife for the health facility whose scope of clinical practice or engagement will be affected by the CEO’s decision;

(b) the scope of clinical practice committee that prepared the scope of clinical practice report;

(c) all appropriate officers under the CEO’s authority or direction so that proper effect can be given to the decision.

Examples—appropriate officers
- general manager of the health facility
- clinical unit director
- head of department at health facility
- immediate supervisor of doctor, dentist or eligible midwife
- human resource personnel

(4) A scope of clinical practice executive decision notice in relation to a doctor, dentist or eligible midwife must include the following information:

(a) if the doctor’s, dentist’s or eligible midwife’s scope of clinical practice is to stay the same—a statement to that effect;

(b) if the doctor’s, dentist’s or eligible midwife’s scope of clinical practice is to be amended—how the scope of clinical practice is being amended;
(c) if the doctor’s, dentist’s or eligible midwife’s scope of clinical practice is to be withdrawn—a statement to that effect;

(d) if the term of engagement of the doctor, dentist or eligible midwife by a health facility is to be amended—how the term is being amended;

(e) if the engagement of the doctor, dentist or eligible midwife by a health facility is to be suspended—the period for which the engagement is being suspended;

(f) if the engagement of the doctor, dentist or eligible midwife by a health facility is to be ended—a statement to that effect;

(g) if the doctor, dentist or eligible midwife was the subject of a decision of the CEO under section 69—a statement to that effect;

(h) when the decision takes effect.

(5) The scope of clinical practice review notice must be in accordance with the requirements for a reviewable decision notice.

Note  The requirements for reviewable decision notices are prescribed under the ACT Civil and Administrative Tribunal Act 2008.

71 When CEO decision on scope of clinical practice report takes effect

(1) A decision of the CEO of a health facility under section 69 or section 70 in relation to a doctor, dentist or eligible midwife for the health facility takes effect on the later of the following:

(a) the day stated in the scope of clinical practice review notice for the decision;

(b) the day the scope of clinical practice review notice is given to the doctor, dentist or eligible midwife.
(2) For subsection (1) (b), if the notice cannot be given to the doctor, dentist or eligible midwife in person, the notice is taken to be given to the doctor, dentist or eligible midwife 7 days after the day it is posted to his or her last known home address.

72 CEO may give information about decision to health facility outside ACT

(1) If the CEO of a health facility makes a decision under section 69 or section 70 to amend or withdraw the scope of clinical practice of a doctor, dentist or eligible midwife, the CEO may tell the CEO of a health facility that is outside the ACT (the other CEO) about the amendment or withdrawal.

(2) However, the CEO may not tell the other CEO about the amendment or withdrawal, unless the other CEO asks, in writing, for information about the scope of clinical practice of the doctor, dentist or eligible midwife.

73 Request for information by health facility outside ACT

(1) This section applies if a health facility outside the ACT (the requesting facility) asks the CEO of a health facility for clinical practice information about a doctor, dentist or eligible midwife that has been the subject of a scope of clinical practice review at the health facility.

(2) The CEO must—

(a) if the request for information is in writing—forword the request within 7 days of receiving it to the scope of clinical practice committee that reviewed the doctor’s, dentist’s or eligible midwife’s scope of clinical practice; or

(b) if the request is not in writing—tell the requesting facility as soon as practicable that the request must be made in writing.
(3) A scope of clinical practice review committee that receives a request from a CEO under subsection (2) (a) may give the requesting facility—

(a) the following information if the information formed part of the committee’s review of the doctor, dentist or eligible midwife, and is relevant to the information asked for by the requesting facility:

(i) particulars of the complaint against the doctor, dentist or eligible midwife;

(ii) particulars about any patients treated by the doctor, dentist or eligible midwife;

(iii) health facility medical records;

(iv) reports from other providers of health services; and

(b) a summary of the committee’s review report into the doctor’s, dentist’s or eligible midwife’s scope of clinical practice.

(4) However, any information given to a requesting facility under subsection (3) must be given in a form that does not allow a person mentioned in the information, other than the doctor, dentist or eligible midwife reviewed by the committee, to be identified.

(5) In this section:

*clinical practice information*, about a doctor, dentist or eligible midwife, means information relating to the clinical competency and standards of professional conduct of the doctor, dentist or eligible midwife.
74 Scope of clinical practice committees—giving information to health board and health services commissioner

(1) A scope of clinical practice committee may give protected information to a health board if the committee is satisfied that giving the information would be likely to facilitate the improvement of health services provided in the ACT.

Note Protected information includes sensitive information (see s 123).

(2) If a clinical practice committee gives protected information to a health board under subsection (1), the committee must give the information to the health services commissioner.

(3) A scope of clinical practice committee must tell the relevant health board, and the health services commissioner, if the committee is satisfied that the clinical practice of a doctor, dentist or eligible midwife has failed to meet a required standard of practice, or that the doctor, dentist or eligible midwife does not satisfy the suitability to practise requirements.

Note The *Health Practitioner Regulation National Law (ACT)*, pt 8, div 2 imposes an obligation to report misconduct or impairment.

75 Scope of clinical practice committees—admissibility of evidence

(1) The following are not admissible as evidence in a proceeding before a court:

   (a) an oral statement made in a proceeding before a scope of clinical practice committee;

   (b) a document given to a scope of clinical practice committee, but only to the extent that it was prepared only for the committee;

   (c) a document prepared by a scope of clinical practice committee.
Part 5  Reviewing scope of clinical practice

Section 76

(2) In this section:

*court* includes a tribunal, authority or person with power to require the production of documents or the answering of questions.

76 Sharing information with other committees

A scope of clinical practice committee may share the following information, including protected information, with another scope of clinical practice committee or a quality assurance committee:

(a) any information that comes before the committee in the course of its functions;

(b) a decision of a CEO under section 69 or section 70 that related to a recommendation made by the committee.

77 Sharing information with 3rd parties

(1) This section applies if—

(a) the CEO of a health facility makes a decision, under section 69 or section 70, to amend or withdraw the scope of clinical practice of a doctor, dentist or eligible midwife; and

(b) a person asks for information about the decision.

(2) The CEO may give the person information about the decision, but may not disclose the identity of the doctor, dentist or eligible midwife or any other sensitive information.

*Note*  *Sensitive information*—see s 124.
78 Complainants to remain anonymous

If a person makes a complaint about a doctor, dentist or eligible midwife and the matter is referred to a scope of clinical practice committee, the committee—

(a) must not disclose the identity of the complainant to the doctor, dentist, eligible midwife or any other person who is not a member of the committee; and

(b) if the committee provides any information to a person about a complaint—may provide information in a way that protects the identity of the complainant unless required to do otherwise by this Act or any other Territory law.
Part 6 Abortions

Division 6.1 Abortions—generally

Section 80

80 Definitions—pt 6

(1) In this part:

*abortifacient* means a medicine, drug or other substance that causes a pregnancy to end prematurely.

*abortion* means a medical abortion or surgical abortion.

*approved medical facility* means a medical facility approved under section 84.

*surgical abortion* means a surgical procedure or any other procedure or act (other than the administration or supply of an abortifacient) that causes a pregnancy to end prematurely.

(2) In this section:

*medical abortion* means the prescription, supply or administration of an abortifacient.

81 Offence—unauthorised supply or administration of abortifacient

(1) A person commits an offence if—

(a) the person supplies or administers an abortifacient to another person; and

(b) the abortifacient is supplied or administered by the person for the purpose of ending a pregnancy; and

(c) the person is not a doctor.

Maximum penalty: imprisonment for 5 years.
(2) Subsection (1) does not apply to—
   (a) a pharmacist supplying an abortifacient in accordance with a prescription; or
   (b) a person assisting a pharmacist in supplying an abortifacient in accordance with a prescription.

(3) For this section, it does not matter whether or not—
   (a) the other person was pregnant; or
   (b) the abortifacient supplied or administered was sufficient to end a pregnancy.

(4) In this section:
   
   prescription—see the Medicines, Poisons and Therapeutic Goods Act 2008, dictionary.

82 Offence—unauthorised surgical abortion

(1) A person commits an offence if the person—
   (a) carries out a surgical abortion; and
   (b) is not a doctor.

   Maximum penalty: imprisonment for 5 years.

(2) Subsection (1) does not apply to a person assisting a doctor to carry out a surgical abortion.

83 Surgical abortion to be carried out in approved medical facility

A person commits an offence if the person carries out a surgical abortion other than in an approved medical facility.

Maximum penalty: 50 penalty units, imprisonment for 6 months or both.
**84 Approval of facilities**

(1) A person may apply to the Minister to have a medical facility, or a part of a medical facility, approved to carry out surgical abortions.

(2) The Minister must approve the application if reasonably satisfied the medical facility is suitable.

(3) An approval is a notifiable instrument.

*Note* A notifiable instrument must be notified under the *Legislation Act*.

**84A Conscientious objection**

(1) Subject to subsection (2), an authorised person may refuse to prescribe, supply or administer an abortifacient, or carry out or assist in carrying out a surgical abortion, on religious or other conscientious grounds (*a conscientious objection*).

(2) An authorised person must not refuse, only because of a conscientious objection—

(a) to carry out, or assist in carrying out, a surgical abortion in an emergency where an abortion is necessary to preserve the life of the pregnant person; or

(b) to provide medical assistance or treatment to a person requiring medical treatment because of an abortion.

(3) There is no breach of duty (by contract or by statutory or other legal requirement) or contravention of a territory law if an authorised person refuses to prescribe, supply or administer an abortifacient, or carry out or assist in carrying out a surgical abortion, because of a conscientious objection.

(4) However, if an authorised person refuses to prescribe, supply or administer an abortifacient, or carry out or assist in carrying out a surgical abortion because of a conscientious objection, the authorised person must tell a person requesting the abortifacient or abortion that the authorised person refuses because of the objection.
(5) In this section:

authorised person means a doctor or nurse.

Division 6.2 Patient privacy in protected areas

85 Definitions—div 6.2

(1) In this division:

capture visual data—a person captures visual data of another person if the person captures moving or still images of the other person by a camera or any other means in such a way that—

(a) a recording is made of the images; or

(b) the images are capable of being transmitted in real time with or without retention or storage in a physical or electronic form; or

(c) the images are otherwise capable of being distributed.

prohibited behaviour, in a protected area around a protected facility, means any of the following:

(a) the harassment, hindering, intimidation, interference with, threatening or obstruction of a person, including by the capturing of visual data of the person, in the protected period that is intended to stop the person from—

(i) entering the protected facility; or

(ii) having an abortion, providing a surgical abortion or prescribing, supplying or administering an abortifacient in the protected facility;

(b) an act that—

(i) can be seen or heard by anyone in the protected period; and

(ii) is intended to stop a person from—

(A) entering the protected facility; or
Part 6
Division 6.2
Patient privacy in protected areas

Section 86

(B) having an abortion, providing a surgical abortion or prescribing, supplying or administering an abortifacient in the protected facility;

(c) a protest, by any means, in the protected period in relation to a person doing any of the things mentioned in paragraph (b) (ii) (A) or (B).

protected area means an area declared under section 86.

protected facility means an approved medical facility or other place around which a protected area has been declared under section 86.

(2) For this section, protected period, in relation to a protected facility, means the period between 7 am and 6 pm on each day the facility is open or any other period declared by the Minister.

(3) A declaration is a disallowable instrument.

Note A disallowable instrument must be notified, and presented to the Legislative Assembly, under the Legislation Act.

86 Declaration of protected area

(1) The Minister must declare an area around an approved medical facility to be a protected area.

(2) The Minister may declare an area around a place where an abortifacient is prescribed, supplied or administered to be a protected area.

(3) In making the declaration, the Minister must be satisfied that the area declared is—

(a) not less than 50m at any point from the protected facility; and

(b) sufficient to ensure the privacy and unimpeded access for anyone entering, trying to enter or leaving the protected facility; but

(c) no bigger than necessary to ensure that outcome.
A declaration is a disallowable instrument.

Note A disallowable instrument must be notified, and presented to the Legislative Assembly, under the Legislation Act.

87 Prohibited behaviour in or in relation to protected area

(1) A person commits an offence if the person—

(a) is in a protected area; and

(b) engages in prohibited behaviour.

Maximum penalty: 25 penalty units.

(2) A person commits an offence if—

(a) the person publishes captured visual data of a person (the recorded person) entering or leaving, or trying to enter or leave, a protected facility; and

(b) the person does so with the intention of stopping a person from—

(i) having an abortion; or

(ii) providing a surgical abortion; or

(iii) prescribing, supplying or administering an abortifacient; and

(c) the recorded person did not consent to the publication.

Maximum penalty: 50 penalty units, imprisonment for 6 months or both.
(3) In this section:

*publish*, captured visual data—

(a) means communicate or distribute visual data in a way or to an extent that makes it available to, or likely to come to the notice of, the public or a section of the public or anyone else not lawfully entitled to the visual data; and

(b) includes—

(i) entering into an agreement or arrangement to do a thing mentioned in paragraph (a); and

(ii) attempting to do a thing mentioned in paragraph (a) or subparagraph (i).
Part 7  

VMO service contracts

100 Definitions for pt 7

In this part:

authorised representative means an entity authorised as a representative under section 105.

core conditions means conditions determined under section 102.

entity means a corporation or an unincorporated association.

negotiating agent means an entity approved as a negotiating agent under section 104.

negotiating period—see section 103 (2).

practice corporation, of a VMO, means a corporation that is controlled or conducted by the VMO and by which the VMO conducts his or her practice as a doctor or dentist.

service contract means a contract for services, between the Territory and a VMO (or the VMO’s practice corporation), under which the VMO is to provide health services to or for the Territory.

VMO (visiting medical officer) means a doctor or dentist who is engaged, or who the Territory proposes to engage, under a service contract.

101 Service contracts

(1) The Territory must not enter into a service contract unless it includes the core conditions that apply to the contract.

(2) A service contract entered into in contravention of subsection (1) is void.

(3) A condition of a service contract that is inconsistent with a core condition that applies to the contract is void to the extent of the inconsistency.
102 Core conditions

(1) The Minister may determine core conditions for service contracts.

(2) The Minister must not determine a condition as a core condition unless the condition has been—
   (a) agreed in collective negotiations under section 103; or
   (b) decided by arbitration under section 106.

(3) A determination of core conditions is a notifiable instrument.

   Note A notifiable instrument must be notified under the Legislation Act.

103 Collective negotiations

(1) The Territory may negotiate with a negotiating agent, or negotiating agents, to establish proposed core conditions for service contracts.

(2) Before beginning collective negotiations, the Minister must determine a period (the negotiating period) for the negotiations.

(3) A negotiating period must not be shorter than 3 months unless the parties to the negotiations agree to a shorter negotiating period.

(4) A determination of a negotiating period is a notifiable instrument.

   Note A notifiable instrument must be notified under the Legislation Act.

104 Negotiating agents

(1) The Minister may, in writing, approve an entity as a negotiating agent.

(2) The Minister must not approve an entity as a negotiating agent unless the Minister is satisfied that—
   (a) the entity is the authorised representative of at least 50 VMOs who, between them, belong to at least 3 of the following categories:
      (i) physician;
      (ii) surgeon;
(iii) obstetrician and gynaecologist;
(iv) anaesthetist;
(v) general practitioner or other doctor or dentist; and

(b) the entity is not disqualified under subsection (3); and

(c) the entity is otherwise suitable to be a negotiating agent having regard to anything that may reasonably influence that decision, including the following:

(i) any criminal or civil court proceedings in which the entity or an executive officer of the entity has been concerned in the previous 10 years;
(ii) any levy of execution against the entity or an executive officer of the entity that is not satisfied;
(iii) whether an executive officer of the entity has ceased to carry on business, or has been involved in the management of an entity that has ceased to carry on business, with the result that creditors were not fully paid or are unlikely to be fully paid.

(3) For subsection (2) (b), an entity is disqualified if—

(a) the entity, or an executive officer of the entity, has been convicted, in the ACT or elsewhere, of—

(i) an offence punishable by imprisonment for longer than 1 year; or

(ii) an offence that involves dishonesty and is punishable by imprisonment for 3 months or longer; or

(b) the entity has a receiver, receiver and manager, or provisional liquidator appointed over part or all of its affairs, or is otherwise under external administration; or
(c) the entity is bankrupt or personally insolvent; or

Note Bankrupt or personally insolvent—see the Legislation Act, dict, pt 1.

(d) the executive officer of the entity is disqualified from managing corporations under the Corporations Act, part 2D.6 (Disqualification from managing corporations).

(4) In this section:

executive officer, of an entity, means a person, by whatever name called, and whether or not the person is a director of the entity, who is concerned with or takes part in the management of the entity.

105 Authorised representatives

(1) A VMO may, in writing, authorise 1 entity to represent the VMO in collective negotiations under section 103.

Note If a form is approved under s 194 for an authorisation, the form must be used.

(2) The authorisation must nominate 1 of the categories mentioned in section 104 (2) (a) as the category to which the VMO belongs.

106 Arbitration

(1) This section applies if agreement is not reached in collective negotiations between the Territory and a negotiating agent or negotiating agents in relation to a matter before the end of the negotiating period.

(2) Unless resolved by mediation beforehand, the matter must be decided by arbitration.

(3) The arbitration must be conducted under the Commercial Arbitration Act 2017 and in accordance with principles and rules determined by the Minister.
(4) That Act applies to the arbitration as if the determined principles and rules were an arbitration agreement between the Territory and the negotiating agent or negotiating agents.

(5) The principles and rules—

(a) must be determined by the Minister having regard to the objective of improving the efficiency, effectiveness and quality of health services, and other public interest considerations; and

(b) must include a requirement that the arbitrator—

(i) be a person with experience in determining industrial awards or a barrister with mediation experience; and

(ii) have appropriate experience to enable the arbitrator to carry out the arbitrator’s role; and

(c) must be fair and reasonable.

(6) A determination of principles and rules for arbitration is a notifiable instrument.

Note A notifiable instrument must be notified under the Legislation Act.

107 Competition and Consumer Act authorisation

For the Competition and Consumer Act 2010 (Cwlth) and the Competition Code of the ACT, the following are authorised:

(a) collective negotiations between the Territory and an approved negotiating agent, or approved negotiating agents, under this part;

(b) the conditions agreed in those negotiations;

(c) service contracts containing core conditions;

(d) everything done under a service contract.
Part 8  Secrecy

120  Definitions—pt 8

In this part:

*divulge*—see section 121.

*information holder*—see section 122.

*protected information*—see section 123.

*Note*  *Sensitive information* is defined for the Act in s 124.

121  When is information *divulged*?

In this part:

*divulge* includes communicate.

122  Who is an *information holder*?

For this part, a person is an *information holder* if—

(a)  the person is or has been—

(i)  a member of a quality assurance committee; or

(ii)  a member of a scope of clinical practice committee; or

(iii)  someone else exercising a function under part 4 (Quality assurance) or part 5 (Reviewing scope of clinical practice); or

(iv)  someone else engaged in the administration of part 4 (Quality assurance) or part 5 (Reviewing scope of clinical practice); or
(b) the person has been given information under this Act by a person mentioned in paragraph (a).

Note Information may be given to people under various provisions of pt 4 and pt 5, including:

- s 39 (Giving health service reports to CEO or director-general)
- s 43 (Quality assurance committees—giving information to the Coroner’s Court)
- s 44 (Quality assurance committees—giving information to other quality assurance committees)
- s 45 (Quality assurance committees—giving information to health board and health services commissioner).
- s 74 (Scope of clinical practice committees—giving information to health board and health services commissioner).

123 What is protected information?

(1) For this part, information is protected information about a person if it is information about the person that is disclosed to, or obtained by, an information holder because of the exercise of a function under this Act by the information holder or someone else.

(2) Without limiting subsection (1), protected information includes sensitive information.

124 What is sensitive information?

In this Act:

Sensitive information means information that—

(a) identifies a person who—

(i) has received a health service; or

(ii) is a health service provider; or
Part 8  Secrecy

Section 125

(iii) has provided information to a quality assurance committee under section 35 (Quality assurance committees—obtaining information) or otherwise in the course of the committee carrying out the committee’s functions under this Act; or

(iv) has provided information to a scope of clinical practice committee under section 64 (Scope of clinical practice committees—obtaining information) or otherwise in the course of the committee carrying out the committee’s functions under this Act; or

(b) would allow the identity of the person to be worked out.

125 Offence—secrecy of protected information

(1) An information holder commits an offence if—

(a) the information holder—

(i) makes a record of protected information about someone else; and

(ii) is reckless about whether the information is protected information about someone else; or

(b) the information holder—

(i) does something that divulges protected information about someone else; and

(ii) is reckless about whether—

(A) the information is protected information about someone else; and

(B) doing the thing would result in the information being divulged to another person.

Maximum penalty: 50 penalty units, imprisonment for 6 months or both.
(2) This section does not apply to the making of a record or the divulging of information if the record is made or the information divulged—
   (a) under this Act; or
   (b) in the exercise of a function, as an information holder, under this Act.

(3) This section does not apply to the making of a record or the divulging of information if—
   (a) the protected information is not sensitive information; and
   (b) the record is made or the information divulged—
      (i) under another territory law; or
      (ii) in the exercise of a function, as an information holder, under another territory law.

(4) This section does not apply to the divulging of protected information about someone with the person’s agreement.

(5) An information holder must not divulge protected information to a court, or produce a document containing protected information to a court, unless it is necessary to do so for this Act.

   *Note* A quality assurance committee may give protected information to the Coroner’s Court (see s 43).

(6) In this section:

   *court* includes a tribunal, authority or person with power to require the production of documents or the answering of questions.

   *produce* includes allow access to.
126 Information may be given to Chief Executive Medicare

(1) The CEO of a health facility may give protected information about a health service provided by a health service provider for the health facility to—

(a) Chief Executive Medicare; or

(b) the auditor-general.

Note Protected information includes sensitive information (see s 123).

(2) However, the CEO must not give the information unless—

(a) the CEO is satisfied that the giving of the information will help the prevention or detection of fraud; and

(b) the Minister agrees, in writing, to the giving of the information.

(3) In this section:

CEO, of a health facility—see section 22.

Chief Executive Medicare—see the Human Services (Medicare) Act 1973 (Cwlth), section 3.
Part 8A  Offence—provision of health services by non-health practitioners

127  Provision of regulated health service by person not health practitioner

(1)  A person commits an offence if—

   (a)  the person intentionally provides a regulated health service; and
   
   (b)  the person is not a health practitioner.

Maximum penalty:  50 penalty units, imprisonment for 6 months or both.

Example—someone providing a regulated health service to someone in the ACT when not a health practitioner

A person (Dr W) provides a medical service by a video link from an island in the south Pacific to Mary Smith in the ACT. Dr W advises Mary that she needs to have her tonsils removed. Dr W is not a health practitioner. Dr W contravenes this subsection.

(2)  This section does not apply to—

   (a)  a health service provided in an emergency; or

   (b)  the provision, by mail order, or over the internet or by other electronic means, of manufactured aids to rehabilitation or surgical prosthetics and orthotics; or

   (c)  a health service ordinarily provided in the ordinary course of business by people other than health practitioners.

Example—par (b)
dental restorative or corrective devices

(3)  In this section:

regulated health service means a health service ordinarily provided by a health practitioner.
Part 10  Review of decisions

Section 130

Part 10  Review of decisions

130  Review of decisions

(1) A doctor, dentist or eligible midwife for a health facility may apply to the ACAT for review of a decision of the CEO of the health facility under section 69—

(a) to amend or withdraw the scope of clinical practice of the doctor, dentist or eligible midwife; or

(b) to amend the terms of engagement of the doctor, dentist or eligible midwife; or

(c) to suspend or end the engagement of the doctor, dentist or eligible midwife.

(2) An applicant under section 84 may apply to the ACAT for review of a decision of the Minister to refuse approval of a medical facility, or a part of a medical facility, to carry out surgical abortions.

131  Pt 10 obligations—no contracting out

To remove any doubt, for section 130 this part applies in relation to a doctor, dentist or eligible midwife for a health facility despite anything to the contrary in a term of the doctor’s, dentist’s or eligible midwife’s engagement.
Part 15  Miscellaneous

189  Protection of doctor, dentist or eligible midwife from liability in emergency

(1) A doctor, dentist or eligible midwife for a health facility does not incur personal civil liability for an act done or omission made that falls outside the doctor’s, dentist’s or eligible midwife’s scope of clinical practice at the health facility if done or made honestly and without recklessness to assist, or give advice about the assistance to be given to, a person who is apparently—

(a) injured or at risk of being injured; or

(b) in need of emergency medical assistance.

(2) However, the protection does not apply if—

(a) there is in force a professional indemnity insurance arrangement that covers the liability; or

(b) the doctor’s, dentist’s or eligible midwife’s capacity to exercise appropriate care and skill was, at the relevant time, significantly impaired by a recreational drug.

(3) In this section:

recreational drug means a drug consumed voluntarily for non-medicinal purposes, and includes alcohol.
190*Disclosure of interests by committee members*

(1) If a member of a committee to which this section applies has a material interest in an issue being considered, or about to be considered, by the committee, the member must disclose the nature of the interest at a committee meeting as soon as practicable after the relevant facts come to the member’s knowledge.

*Note 1* This section applies to a quality assurance committee (see s 32) and a scope of clinical practice committee (see s 61).

*Note 2* Material interest is defined in s (4). The definition of indirect interest in s (4) applies to the definition of material interest.

(2) The disclosure must be recorded in the committee’s minutes and, unless the committee otherwise decides, the member must not—

(a) be present when the committee considers the issue; or

(b) take part in a decision of the committee on the issue.

Example
Adam, Ben and Charlotte are members of a quality assurance committee. They have an interest in an issue being considered at a committee meeting and they disclose the interest as soon as they become aware of it. Adam’s and Ben’s interests are minor but Charlotte has a direct financial interest in the issue. The committee considers the disclosures and decides that because of the nature of the interests:

- Adam may be present when the committee considers the issue but not take part in the decision
- Ben may be present for the consideration and take part in the decision.

The committee does not make a decision allowing Charlotte to be present or take part in the committee’s decision. Accordingly, since Charlotte has a material interest she cannot be present for the consideration of the issue or take part in the decision.

(3) Any other committee member who also has a material interest in the issue must not be present when the committee is considering its decision under subsection (2).
(4) In this section:

*associate*, of a person, means—

(a) the person’s business partner; or
(b) a close friend of the person; or
(c) a family member of the person.

*executive officer*, of a corporation, means a person (however described) who is concerned with, or takes part in, the corporation’s management, whether or not the person is a director of the corporation.

*indirect interest*—without limiting the kinds of indirect interests a person may have, a person has an *indirect interest* in an issue if any of the following has an interest in the issue:

(a) an associate of the person;
(b) a corporation if the corporation has not more than 100 members and the person, or an associate of the person, is a member of the corporation;
(c) a subsidiary of a corporation mentioned in paragraph (b);
(d) a corporation if the person, or an associate of the person, is an executive officer of the corporation;
(e) the trustee of a trust if the person, or an associate of the person, is a beneficiary of the trust;
(f) a member of a firm or partnership if the person, or an associate of the person, is a member of the firm or partnership;
(g) someone else carrying on a business if the person, or an associate of the person, has a direct or indirect right to participate in the profits of the business.
material interest—a committee member has a material interest in an issue if the member has—

(a) a direct or indirect financial interest in the issue; or

(b) a direct or indirect interest of any other kind if the interest could conflict with the proper exercise of the member’s functions in relation to the committee’s consideration of the issue.

191 References to Health and Community Care Service

(1) In any Act, instrument made under an Act, contract or other document, a reference to the Health and Community Care Service is, for the application of that Act, instrument, contract or other document after the commencement of this section, a reference to the Territory.

(2) In this section:

Health and Community Care Service means the Australian Capital Territory Health and Community Care Service established by the Health and Community Care Services Act 1996 (repealed).

192 Determination of fees

(1) The Minister may determine fees for this Act.

Note The Legislation Act contains provisions about the making of determinations and regulations relating to fees (see pt 6.3)

(2) Without limiting subsection (1), the Minister may determine fees in relation to the provision of health and community care services.

(3) A determination is a disallowable instrument.

Note A disallowable instrument must be notified, and presented to the Legislative Assembly, under the Legislation Act.
(4) A determination may adopt a Commonwealth law or a health benefits agreement (or a provision of a Commonwealth law or health benefits agreement) as in force from time to time.

Note 1 The text of an applied, adopted or incorporated law or instrument, whether applied as in force from time to time or at a particular time, is taken to be a notifiable instrument if the operation of the Legislation Act, s 47 (5) or (6) is not disapplied (see s 47 (7)).

Note 2 A notifiable instrument must be notified under the Legislation Act.

(5) In this section:

*Commonwealth law* means a Commonwealth Act, or any regulations, rules, ordinance or disallowable instrument under a Commonwealth Act.

*disallowable instrument*, for a Commonwealth Act, means a disallowable instrument under the *Acts Interpretation Act 1901* (Cwlth), section 46A.

*health benefits agreement* means an agreement between the Territory and an entity that provides health benefits to contributors of a health benefits fund conducted by the entity.

### 193 Payment of fees and interest

(1) A fee is payable to the Territory on or before the payment date.

(2) If an amount for a fee remains unpaid after the payment date, in addition to that amount, interest calculated on the aggregate amount at the rate determined by the Minister is payable to the Territory in relation to every month or part of a month that the aggregate amount remains unpaid.

(3) A determination is a disallowable instrument.

Note A disallowable instrument must be notified, and presented to the Legislative Assembly, under the Legislation Act.
(4) In this section:

**aggregate amount**, for a month, means the total of—

(a) the amount of the fee; and

(b) the amount of interest;

remaining unpaid at the end of the previous month.

**payment date**, for a fee, means the 28th day after the day when the account for the fee was issued.

194 **Approved forms**

(1) The Minister may approve forms for this Act.

(2) If the Minister approves a form for a particular purpose, the form must be used for that purpose.

*Note* For other provisions about forms, see **Legislation Act**, s 255.

(3) An approved form is a notifiable instrument.

*Note* A notifiable instrument must be notified under the **Legislation Act**.

196 **Regulation-making power**

The Executive may make regulations for this Act.

*Note* Regulations must be notified, and presented to the Legislative Assembly, under the **Legislation Act**.
Dictionary

(see s 2)

Note 1  The Legislation Act contains definitions and other provisions relevant to this Act.

Note 2  For example, the Legislation Act, dict, pt 1 defines the following terms:

- ACAT
- Act
- ACT
- appoint
- Commonwealth
- Coroner’s Court
- director-general (see s 163)
- doctor
- entity
- exercise
- found guilty
- function
- health practitioner
- in relation to
- interest
- make
- Minister (see s 162)
- notifiable instrument (see s 10)
- penalty unit (see s 133)
- pharmacist
- proceeding
- territory law
- the Territory
- tribunal.

abortifacient, for part 6 (Abortions)—see section 80 (1).

abortion, for part 6 (Abortions)—see section 80 (1).
approved medical facility, for part 6 (Abortions)—see section 80 (1).

authorised representative, for part 7 (VMO service contracts)—see section 100.

capture visual data, for division 6.2 (Patient privacy in protected areas)—see section 85.

CEO—

(a) of a health facility, for part 4 (Quality assurance)—see section 22; and

(b) of a health professional organisation, for part 4 (Quality assurance)—see section 23; and

(c) of a health facility, for part 5 (Reviewing scope of clinical practice)—see section 53.

chief executive officer, Calvary, for part 5 (Reviewing scope of clinical practice)—see section 50.

core conditions, for part 7 (VMO service contracts)—see section 100.

day hospital means a facility where a person is admitted for surgical or medical treatment and discharged on the same day.

dental technical work means work involving the making, altering, repairing or maintaining of dental prosthetic appliances.

Example—dental technical work
shade-taking for dental prosthetic appliances

dental technician means a person who does dental technical work and either—

(a) is a graduate of a course of education in dental technical work accredited by the Council of Regulating Authorities for Dental Technicians and Dental Prosthetists (CORA); or
(b) has—

(i) completed a course of education or training in dental prosthetic work outside Australia that is accredited by CORA; and

(ii) passed an exam in dental prosthetic work accredited by CORA.

dentist, for a health facility, for part 5 (Reviewing scope of clinical practice)—see section 52.
divulge, for part 8 (Secrecy)—see section 121.
doctor, for a health facility, for part 5 (Reviewing scope of clinical practice)—see section 52.
eligible midwife, for a health facility, for part 5 (Reviewing scope of clinical practice)—see section 52.
engage in conduct means—

(a) do an act; or

(b) omit to do an act.

dentist, for a health facility, for part 5 (Reviewing scope of clinical practice)—see section 52.
divulge, for part 8 (Secrecy)—see section 121.
doctor, for a health facility, for part 5 (Reviewing scope of clinical practice)—see section 52.
eligible midwife, for a health facility, for part 5 (Reviewing scope of clinical practice)—see section 52.
engage in conduct means—

(a) do an act; or

(b) omit to do an act.

dentist, for a health facility, for part 5 (Reviewing scope of clinical practice)—see section 52.
divulge, for part 8 (Secrecy)—see section 121.
doctor, for a health facility, for part 5 (Reviewing scope of clinical practice)—see section 52.
eligible midwife, for a health facility, for part 5 (Reviewing scope of clinical practice)—see section 52.
engage in conduct means—

(a) do an act; or

(b) omit to do an act.

entity, for part 7 (VMO service contracts)—see section 100.
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health facility—see section 6.
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health professional organisation QAC, for a health professional organisation, for part 4 (Quality assurance)—see section 20.
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**health service report**, for part 4 (Quality assurance)—see section 38.

**hospital**, for part 5 (Reviewing scope of clinical practice)—see section 50.

**information holder**, for part 8 (Secrecy)—see section 122.

**ministerial report**, for part 4 (Quality assurance)—section 41.

**negotiating agent**, for part 7 (VMO service contracts)—see section 100.

**negotiating period**, for part 7 (VMO service contracts)—see section 103 (2).

**practice corporation**, for part 7 (VMO service contracts)—see section 100.

**prohibited behaviour**, for division 6.2 (Patient privacy in protected areas)—see section 85.

**protected area**, for division 6.2 (Patient privacy in protected areas)—see section 85.

**protected facility**, for division 6.2 (Patient privacy in protected areas)—see section 85.

**protected information**, for part 8 (Secrecy)—see section 123.

**quality assurance committee**—see section 24.

**review**, in relation to the scope of clinical practice, for part 5 (Reviewing scope of clinical practice)—see section 55.

**scope of clinical practice**, of a doctor, dentist or eligible midwife, for a health facility, for part 5 (Reviewing scope of clinical practice)—see section 54.

**scope of clinical practice committee**—see section 51.

**scope of clinical practice executive decision notice**, for part 5 (Reviewing scope of clinical practice)—see section 70.
**scope of clinical practice report**, for part 5 (Reviewing scope of clinical practice)—see section 67.

**sensitive information**—see section 124.

**service contract**, for part 7 (VMO service contracts)—see section 100.

**special purpose QAC**, for part 4 (Quality assurance)—see section 20.

**surgical abortion**, for part 6 (Abortions)—see section 80 (1).

**VMO**, or visiting medical officer, for part 7 (VMO service contracts)—see section 100.
Endnotes

1 About the endnotes

Amending and modifying laws are annotated in the legislation history and the amendment history. Current modifications are not included in the republished law but are set out in the endnotes.

Not all editorial amendments made under the Legislation Act 2001, part 11.3 are annotated in the amendment history. Full details of any amendments can be obtained from the Parliamentary Counsel’s Office.

Uncommenced amending laws are not included in the republished law. The details of these laws are underlined in the legislation history. Uncommenced expiries are underlined in the legislation history and amendment history.

If all the provisions of the law have been renumbered, a table of renumbered provisions gives details of previous and current numbering.

The endnotes also include a table of earlier republications.

2 Abbreviation key

A = Act
AF = Approved form
am = amended
amdt = amendment
AR = Assembly resolution
ch = chapter
CN = Commencement notice
def = definition
DI = Disallowable instrument
dict = dictionary
disallowed = disallowed by the Legislative Assembly
div = division
exp = expires/expired
Gaz = gazette
hdg = heading
IA = Interpretation Act 1967
ins = inserted/added
LA = Legislation Act 2001
LR = legislation register
LRA = Legislation (Republication) Act 1996
mod = modified/modification
NI = Notifiable instrument
o = order
om = omitted/repealed
ord = ordinance
orig = original
par = paragraph/subparagraph
pres = present
prev = previous
(prev...) = previously
pi = part
r = rule/subrule
reloc = relocated
renum = renumbered
R[X] = Republication No
RI = reissue
sch = schedule
s = section/subsection
sdv = subdivision
SL = Subordinate law
sub = substituted
underlining = whole or part not commenced or to be expired

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3 Legislation history

Health Act 1993 A1993-13
notified 1 March 1993 (Gaz 1993 No S23)
commenced 1 March 1993 (s 2)

as amended by

Health (Amendment) Act 1994 A1994-23
notified 20 May 1994 (Gaz 1994 No S87)
commenced 20 May 1994 (s 2)

notified 30 June 1994 (Gaz 1994 No S121)
s 1, s 2 commenced 30 June 1994 (s 2 (1))
sch 1 pt 44 commenced 1 July 1994 (s 2 (2) and Gaz 1994 No S142)

Administrative Appeals (Consequential Amendments) Act 1994 A1994-60 sch 1
notified 11 October 1994 (Gaz 1994 No S197)
s 1, s 2 commenced 11 October 1994 (s 2 (1))
sch 1 commenced 14 November 1994 (s 2 (2) and see Gaz 1994 No S250)

notified 1 July 1996 (Gaz 1996 No S130)
commenced 1 July 1996 (s 2)

Health (Amendment) Act 1998 A1998-50
notified 16 November 1998 (Gaz 1998 No S205)
commenced 16 November 1998 (s 2)

Statute Law Revision (Penalties) Act 1998 A1998-54 sch
notified 27 November 1998 (Gaz 1998 No S207)
s 1, s 2 commenced 27 November 1998 (s 2 (1))
sch commenced 9 December 1998 (s 2 (2) and Gaz 1998 No 49)
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Legislation (Consequential Amendments) Act 2001 A2001-44 pt 175
notified 26 July 2001 (Gaz 2001 No 30)
s 1, s 2 commenced 26 July 2001 (IA s 10B)
pt 175 commenced 12 September 2001 (s 2 and see Gaz 2001 No S65)

Statute Law Amendment Act 2001 (No 2) 2001 No 56 pt 1.3
notified 5 September 2001 (Gaz 2001 No S65)
s 1, s 2 commenced 5 September 2001 (IA s 10B)
amdts 1.3-1.8, 1.10-1.13, 1.15, 1.16, 1.17, 1.35 commenced 12 September 2001 (s 2 (2))
pt 1.3 remainder commenced 5 September 2001 (s 2 (1))

Health and Community Care Services (Repeal and Consequential Amendments) Act 2002 A2002-47 pt 1.2
notified LR 20 December 2002
s 1, s 2 commenced 20 December 2002 (LA s 75 (1))
pt 1.2 commenced 31 December 2002 (s 2)

Statute Law Amendment Act 2003 A2003-41 sch 1 pt 1.1
notified LR 11 September 2003
s 1, s 2 commenced 11 September 2003 (LA s 75 (1))
sch 1 pt 1.1 commenced 9 October 2003 (s 2 (1))

Health Amendment Act 2003 A2003-43
notified LR 29 September 2003
s 1, s 2 commenced 29 September 2003 (LA s 75 (1))
remainder commenced 30 September 2003 (s 2)

Nurse Practitioners Legislation Amendment Act 2004 A2004-10 pt 2
notified LR 19 March 2004
s 1, s 2 commenced 19 March 2004 (LA s 75 (1))
pt 2 commenced 27 May 2004 (s 2 and CN2004-9)

Health Professionals Legislation Amendment Act 2004 A2004-39
sch 1 pt 1.3
notified LR 8 July 2004
s 1, s 2 commenced 8 July 2004 (LA s 75 (1))
sch 1 pt 1.3 commenced 7 July 2005 (s 2 and see Health Professionals Act 2004 A2004-38, s 2 and CN2005-11)
Health Legislation Amendment Act 2005 A2005-28 amdt 1.70
notified LR 6 July 2005
s 1, s 2 commenced 6 July 2005 (LA s 75 (1))
amdt 1.70 commenced 7 July 2005 (s 2)

notified LR 27 October 2005
s 1, s 2 commenced 27 October 2005 (LA s 75 (1))
sch 1 pt 1.24 commenced 24 November 2005 (s 2)

notified LR 21 December 2005
s 1, s 2 commenced 21 December 2005 (LA s 75 (1))
sch 3 pt 3.10 commenced 11 January 2006 (s 2 (1))

Health Legislation Amendment Act 2006 A2006-27 pt 2, sch 1
notified LR 14 June 2006
s 1, s 2 commenced 14 June 2006 (LA s 75 (1))
pt 2, sch 1 commenced 14 December 2006 (s 2 and LA s 79)

Health Legislation Amendment Act 2006 (No 2) A2006-46 sch 2 pt 2.8
notified LR 17 November 2006
s 1, s 2 commenced 17 November 2006 (LA s 75 (1))
sch 2 pt 2.8 commenced 18 November 2006 (s 2 (2))

Statute Law Amendment Act 2007 (No 2) A2007-16 sch 3 pt 3.18
notified LR 20 June 2007
s 1, s 2 taken to have commenced 12 April 2007 (LA s 75 (2))
sch 3 pt 3.18 commenced 11 July 2007 (s 2 (1))

Medicines, Poisons and Therapeutic Goods Act 2008 A2008-26
sch 2 pt 2.12
notified LR 14 August 2008
s 1, s 2 commenced 14 August 2008 (LA s 75 (1))
sch 2 pt 2.12 commenced 14 February 2009 (s 2 and LA s 79)

ACT Civil and Administrative Tribunal Legislation Amendment
Act 2008 (No 2) A2008-37 sch 1 pt 1.53
notified LR 4 September 2008
s 1, s 2 commenced 4 September 2008 (LA s 75 (1))
sch 1 pt 1.53 commenced 2 February 2009 (s 2 (1) and see ACT Civil
and Administrative Tribunal Act 2008 A2008-35, s 2 (1) and CN2009-2)
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Health Practitioner Regulation National Law (ACT) Act 2010 A2010-10 sch 2 pt 2.7
notified LR 31 March 2010
s 1, s 2 commenced 31 March 2010 (LA s 75 (1))
sch 2 pt 2.7 commenced 1 July 2010 (s 2 (1) (a))
as modified by

Health Practitioner Regulation National Law (ACT) (Transitional Provisions) Regulation 2010 (No 2) SL2010-39 s 3 and sch 1
notified LR 11 October 2010
s 1, s 2 commenced 11 October 2010 (LA s 75 (1))
s 3 and sch 1 commenced 12 October 2010 (s 2)
as amended by

Fair Trading (Australian Consumer Law) Amendment Act 2010 A2010-54 sch 3 pt 3.11
notified LR 16 December 2010
s 1, s 2 commenced 16 December 2010 (LA s 75 (1))
sch 3 pt 3.11 commenced 1 January 2011 (s 2 (1))
Statute Law Amendment Act 2011 A2011-3 sch 3 pt 3.22
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s 1, s 2 commenced 22 February 2011 (LA s 75 (1))
sch 3 pt 3.22 commenced 1 March 2011 (s 2)
Health Amendment Act 2011 A2011-11
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s 1, s 2 commenced 12 April 2011 (LA s 75 (1))
s 4, s 5 commenced 1 July 2011 (s 2 (2))
remainder commenced 13 April 2011 (s 2 (1))
Administrative (One ACT Public Service Miscellaneous Amendments) Act 2011 A2011-22 sch 1 pt 1.76
notified LR 30 June 2011
s 1, s 2 commenced 30 June 2011 (LA s 75 (1))
sch 1 pt 1.76 commenced 1 July 2011 (s 2 (1))
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Statute Law Amendment Act 2011 (No 2) A2011-28 sch 3 pt 3.18
notified LR 31 August 2011
s 1, s 2 commenced 31 August 2011 (LA s 75 (1))
sch 3 pt 3.18 commenced 21 September 2011 (s 2 (1))

Statute Law Amendment Act 2013 A2013-19 sch 1 pt 1.1
notified LR 24 May 2013
s 1, s 2 commenced 24 May 2013 (LA s 75 (1))
sch 1 pt 1.1 commenced 14 June 2013 (s 2)

Statute Law Amendment Act 2013 (No 2) A2013-44 sch 1 pt 1.2,
sch 3 pt 3.11
notified LR 11 November 2013
s 1, s 2 commenced 11 November 2013 (LA s 75 (1))
sch 1 pt 1.2, sch 3 pt 3.11 commenced 25 November 2013 (s 2)

Veterinary Surgeons Act 2015 A2015-29 sch 2 pt 2.4
notified LR 20 August 2015
s 1, s 2 commenced 20 August 2015 (LA s 75 (1))
sch 2 pt 2.4 commenced 1 December 2015 (s 2 (1) and CN2015-22)

Health Legislation Amendment Act 2016 A2016-11 pt 3
notified LR 1 March 2016
s 1, s 2 commenced 1 March 2016 (LA s 75 (1))
pt 3 commenced 2 March 2016 (s 2)

Health (Patient Privacy) Amendment Act 2015 A2015-43
notified LR 4 November 2015
s 1, s 2 commenced 4 November 2015 (LA s 75 (1))
remainder commenced 22 March 2016 (s 2 and CN2016-4)

Statute Law Amendment Act 2017 A2017-4 sch 3 pt 3.14
notified LR 23 February 2017
s 1, s 2 commenced 23 February 2017 (LA s 75 (1))
sch 3 pt 3.14 commenced 9 March 2017 (s 2)

Commercial Arbitration Act 2017 A2017-7 sch 1 pt 1.3
notified LR 4 April 2017
s 1A, s 1B commenced 4 April 2017 (LA s 75 (1))
sch 1 pt 1.3 commenced 1 July 2017 (s 1B and CN2017-1)
Health (Improving Abortion Access) Amendment Act 2018 A2018-37
notified LR 27 September 2018
s 1, s 2 commenced 27 September 2018 (LA s 75 (1))
remainder commenced 1 July 2019 (s 2 (1) and CN2019-10)

Statute Law Amendment Act 2018 A2018-42 sch 1 pt 1.2, sch 3 pt 3.17
notified LR 8 November 2018
s 1, s 2 taken to have commenced 1 July 2018 (LA s 75 (2))
sch 1 pt 1.2, sch 3 pt 3.17 commenced 22 November 2018 (s 2 (1))
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renum as s 13 R4 LA (see A2001-56 amdt 1.36)
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def **health service** ins A2006-27 s 10  
def **health service provider** sub A2004-39 amdt 1.16; A2006-27 s 10  
def **health service report** ins A2006-27 s 10  
  sub A2006-27 s 10  
def **hospital** ins A2006-27 s 10  
  sub A2011-11 s 27  
def **information holder** ins A2006-27 s 10  
def **local hospital network** ins A2011-11 s 28  
  om A2017-4 amdt 3.68  
def **ministerial report** ins A2006-27 s 10  
def **negotiating agent** ins A2003-43 s 6  
  sub A2006-27 s 10  
def **negotiating period** ins A2003-43 s 6  
  sub A2006-27 s 10  
def **practice corporation** ins A2003-43 s 6  
  sub A2006-27 s 10  
def **prescribed body** ins A2001-56 amdt 1.35  
  om A2006-27 s 10  
def **private day hospital facility** ins A2001-56 amdt 1.35  
  om A2006-27 s 10  
def **prohibited behaviour** ins A2015-43 s 6
Endnotes

Amendment history

4

def protected area ins A2015-43 s 6
def protected facility ins A2018-37 s 25
def protected information ins A2006-27 s 10
sub A2011-28 amdt 3.130

def quality assurance activities om A2001-56 amdt 1.5
def quality assurance activity ins A2001-56 amdt 1.5
om A2006-27 s 10

def quality assurance committee ins A2006-27 s 10
def review ins A2006-27 s 10
sub A2011-11 s 29

def scope of clinical practice ins A2011-11 s 24
def scope of clinical practice committee ins A2011-11 s 24
def scope of clinical practice executive decision notice ins
A2011-11 s 24

def scope of clinical practice report ins A2011-11 s 24
def sensitive information ins A2006-27 s 10
def service contract ins A2003-43 s 6
sub A2006-27 s 10

def special purpose QAC ins A2006-27 s 10
am A2018-42 amdt 3.56

def surgical abortion ins A2018-37 s 25
def tribunal om A1994-60 sch 1
def VMO ins A2003-43 s 6
sub A2006-27 s 10
5 **Earlier republications**

Some earlier republications were not numbered. The number in column 1 refers to the publication order.

Since 12 September 2001 every authorised republication has been published in electronic pdf format on the ACT legislation register. A selection of authorised republications have also been published in printed format. These republications are marked with an asterisk (*) in column 1. Electronic and printed versions of an authorised republication are identical.

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Expired transitional or validating provisions

This Act may be affected by transitional or validating provisions that have expired. The expiry does not affect any continuing operation of the provisions (see Legislation Act 2001, s 88 (1)).

Expired provisions are removed from the republished law when the expiry takes effect and are listed in the amendment history using the abbreviation ‘exp’ followed by the date of the expiry.

To find the expired provisions see the version of this Act before the expiry took effect. The ACT legislation register has point-in-time versions of this Act.
7 Renumbered provisions

This Act was renumbered under the *Legislation Act 2001*, in R4 (see A2001-56). Details of renumbered provisions are shown in endnote 4 (Amendment history). For a table showing the renumbered provisions, see R4.