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Animal welfare in New Zealand is safeguarded by the Animal Welfare Act 1999 (the Act). The Act requires those in charge of animals to provide adequately for their physical, health and behavioural needs. These needs include:

- proper and sufficient food and water;
- adequate shelter;
- opportunity to display normal patterns of behaviour;
- physical handling in a manner which minimises the likelihood of unreasonable or unnecessary pain or distress;
- protection from, and rapid diagnosis of, any significant injury or disease.

The nature of the research, testing or teaching (RTT) may mean that the general obligations under the Act cannot be met. For instance, some pain or distress to a small number of animals may occur. This may, however, result in significant benefits to people, other animals or the environment. For this reason, the use of animals in RTT is governed by a separate set of provisions (Part 6) within the Act.

Within the constraints of any project, all reasonable steps must be taken to ensure that the physical, health and behavioural needs of those animals are met in accordance with both good practice and scientific knowledge, as is stated in section 80(2)(a)(i) of the Act.

Traditional acceptance of the use of animals in RTT has come under increasing scrutiny. Both the public and the scientific community are concerned that standards of husbandry and care of animals in laboratories should at least parallel those required in other areas.

For this reason, the use of animals in RTT carries significant responsibilities and strict legislative obligations. Part 6 of the Act allows such activities only where there is good reason to believe:

- that the findings of the research or testing or the results of the teaching will enhance the understanding of human beings, animals, or ecosystems; the maintenance or protection of human or animal health or welfare; the management, protection, or control of ecosystems, plants, animals, or native fauna; the production and productivity of animals; or the achievement of educational objectives; and
- that the benefits derived from the use of animals in research, testing, and teaching are not outweighed by the likely harm to the animals.

### 1.1. Purpose of this Guide

It is intended that this guide will be useful to:

- people or organisations wishing to establish an AEC;
- code holders, or members of current AECs, in discharging their responsibilities;
- investigators, preparing to submit an application to an AEC;
- members of the public, wanting to understand more about how RTT is managed in New Zealand.
Additional intentions are that the guide will:
• promote the humane and responsible use of animals for scientific purposes;
• encourage the highest standard of husbandry and animal care;
• set guidelines for what constitutes “good practice” in managing of animals in the RTT environment.

1.2. Scope of this Guide

The guide covers the use of animals for scientific purposes in medicine, biology, agriculture, veterinary and other animal sciences, industry and teaching. It includes their use in research, teaching, field trials, and product testing.

The guide provides general principles for the care and use of animals and specifies the responsibilities of investigators and teachers. It also provides guidelines for the humane conduct of experiments, and for the acquisition of animals and their care.

Throughout this guide:
• “shall” means that there is a statutory requirement;
• “must” denotes a minimum standard in a code of welfare;
• “should” and “may” denote a recommendation.

1.3. Feedback on this Guide

Comments on this guide are invited and should be sent to the NAEAC Secretary:

NAEAC@mpi.govt.nz or

The Secretary
NAEAC
PO Box 2526
Wellington 6140
DEFINITIONS OF TERMS & ABBREVIATIONS USED IN THIS GUIDE


Analgesia: The temporary abolition or diminution of pain perception.

Anaesthesia: A state of controllable, reversible insensibility in which sensor perception and motor responses are both markedly depressed.

Animal: The Animal Welfare Act 1999 defines “animal” as:

Any live member of the animal kingdom that is:
- A mammal; or
- A bird; or
- A reptile; or
- An amphibian; or
- A fish (bony or cartilaginous); or
- Any octopus, squid, crab, lobster, or crayfish (including freshwater crayfish); or
- Any other member of the animal kingdom which is declared from time to time by the Governor-General, by Order in Council, to be an animal for the purposes of this Act; and
- Includes any mammalian foetus, or any avian or reptilian pre-hatched young, that is in the last half of its period of gestation or development; and
- Includes any marsupial pouch young.

It does not include any other prenatal, pre-hatched, larval or developmental stage.

AEC: Animal Ethics Committee.

Approved project: A project which has been formally approved following evaluation of a written proposal by a properly constituted AEC.

Cachexia: Severe generalised weakness, malnutrition and emaciation.

CEC: Code of ethical conduct.

Distress: Acute or chronic response of an animal caused by stimuli that produce observable biological stress as shown by abnormal physiological or behavioural responses.

Embryonated egg: An egg in the last half of incubation.

EPA: Environmental Protection Authority.

Euthanasia: The humane termination of life.

Experiment: Any test or trial for a scientific purpose, including any activity to test a hypothesis or demonstrate a known fact.

Foetus: An unborn mammal in the post-embryonic stage of development.

Genetic modification (GM): The deletion, change or moving of genes within an organism, or the transfer of genes from one organism to another, or the modification of existing genes or the construction of new genes and their incorporation into any organism.


IATA: International Air Transport Association.

IBSC: Institutional Biological Safety Committee.

IDAO: Internal Drug Administration Order.

Investigator: A person approved by an AEC to be responsible for the conduct of an approved project involving animals.

IOP: Institutional Operating Plan.

MPI: Ministry for Primary Industries.

Moribund: Approaching death; about to die.

Manipulation: This is defined by the Act in relation to any live animal as meaning interfering with the normal physiological, behavioural or anatomical integrity of the animal by deliberately

Subjecting it to a procedure which is unusual or abnormal when compared with that to which animals of that type would be subjected to under normal management or practice and which involves:

- Exposing it to any parasite, micro-organism, drug, chemical, biological product, radiation, electrical stimulation, or environmental condition; or
- Enforced activity, restraint, nutrition, or surgical intervention; or
- Depriving it of usual care;

The term defined by subsection (1) includes

- the killing of an animal (other than an animal in a wild state) for the purpose of interfering with the animal's body or its tissues in a manner specified in that subsection
- the breeding or production of an animal using any breeding technique (including genetic modification) that may result in the birth or production of an animal that is more susceptible to, or at greater risk of, pain or distress during its life as a result of the breeding or production.
It does not include:

- any therapy or prophylaxis necessary or desirable for the welfare of the animal; or
- the killing of an animal as the end point of research testing or teaching if the animal is killed in such a manner that it does not suffer unreasonable or unnecessary pain or distress; or
- the hunting or killing of any animal in a wild state by a method that is not an experimental method;
- any killing of an animal that is carried out by any person:
  - while exercising powers under the Biosecurity Act 1993 for the purposes specified in section 121(1A) of that Act; or
  - while exercising powers or performing functions for the purposes of a response activity carried out under the Biosecurity Act 1993.

**Organisation:** Any individual, company, institution or organisation that might apply for approval of a code. In the case of an organisation, the code holder is the organisation, not a person such as a chief executive. The code is not transferable to another organisation if companies are sold or merge, unless the Director-General of MPI gives specific approval to the transfer prior to the sale or merger.

**Pain:** An awareness of acute or chronic discomfort, occurring in varying degrees of severity, and resulting from injury, disease, or emotional distress as evidenced by biological or behavioural changes or both.

**Parenting:** Any person or organisation operating under a CEC held by another organisation.

**Project:** A series of related experiments that forms a discrete piece of research.

**Proposal:** A written outline of a research project put forward for consideration by an AEC.

**RTT:** Research, testing and teaching.

**Scientific purposes:** All those activities performed to acquire, develop or demonstrate knowledge or techniques in any scientific discipline, including activities for the purposes of teaching, research, diagnosis, product testing, and the production of biological products.

**Tranquillisers:** Drugs which are used to treat anxiety or produce sedation.
3.1. Functions of NAEAC

The National Animal Ethics Advisory Committee (NAEAC) is established under s62 of the Act to:

• to advise the Minister on ethical issues and animal welfare issues arising from RTT;
• to make recommendations to the Minister under section 3(3) (which relates to manipulation);
• to make recommendations to the Director-General under section 85 (which relates to restrictions on use of non-human hominids);
• to provide advice and information on the development and review of codes of ethical conduct;
• to make recommendations to the Director-General concerning the approval, amendment, suspension, or revocation of any code of ethical conduct (CEC);
• to make recommendations to the Minister concerning the setting of standards and policies for codes of ethical conduct;
• to provide information and advice to animal ethics committees;
• to recommend, for approval by the Director-General under section 109, such persons as are, in the opinion of the Committee, suitable for appointment as accredited reviewers;
• to consider the reports of independent reviews of code holders and animal ethics committees;
• to make recommendations to the Minister under section 118(3) (which relates to the power of the Minister to approve research or testing).

3.2. Membership of NAEAC

NAEAC consists of no more than 10 members, including a chairperson, appointed by the Minister. In appointing members other than the chairperson, the Minister must have regard to:

• the public interest in relation to the manipulation of animals in research, testing, and teaching; and
• the need for the Committee to possess knowledge and experience in the following areas;
  • veterinary science;
  • medical science;
  • biological science;
  • the commercial use of animals in research and testing;
  • ethical standards and conduct in respect of animals;
  • education issues, including the use of animals in schools;
  • the manipulation of animals in research, testing, and teaching;
  • environmental and conservation management;
  • animal welfare advocacy;
  • any other area the Minister considers relevant; and
• the need for a balance between those members who are currently involved in research, testing, and teaching and those members who are not so involved.
3.3. Keeping in Touch with NAEAC

Contact with NAEAC can be maintained in a number of ways:

- Write to:
  NAEAC@mpi.govt.nz or
  
  The Secretary
  NAEAC
  PO Box 2526
  Wellington 6140

- Read the NAEAC newsletter.
- Attend a NAEAC workshop. Watch the NAEAC newsletter for details.
- Site visits. NAEAC visits places where RTT is being carried out. These visits aim to:
  - familiarise NAEAC members with actual RTT work being carried out;
  - allow NAEAC members to understand the challenges and experiences of AEC members;
  - educate AEC members about the role of NAEAC.

The selection of the sites to visit are made in consultation with MPI. The visit is NOT an audit exercise. There is a broad range of experience on the committee and there may be opportunity to discuss various aspects of welfare, research or teaching strategy.

Individual NAEAC members may also, from time to time, ask to attend an AEC meeting as an observer. Such visits are for the member’s personal development only and are not made to critique the AEC’s operation. The NAEAC member will not participate in any discussion uninvited.
4.1. Requirement for a CEC

Any person who engages in RTT and wishes to use animals for such purposes shall:

• have an approved CEC, or
• work for a person who holds such a code, or
• have a formal arrangement to use another person’s code pursuant to section 84 of the Act.

Codes must be approved by the Director-General of MPI prior to commencing RTT activities. Approvals are for a maximum of five years.

An application would normally be made by the chief executive of the organisation requiring a code.

4.2. Recommended Content for a CEC

The CEC lists the responsibilities of the code holder, the steps to be undertaken in setting up an AEC, and the policies and procedures to be adopted and followed by the code holder and the AEC.

At a minimum the CEC must meet the requirements specified in section 88 of the Act. A system of keeping records and monitoring all activities involving animal use and husbandry is essential.

Every code holder must also comply with the Animal Welfare (Records and Statistics) Regulations 1999.

The NAEAC template for drafting a CEC can be found here: https://www.mpi.govt.nz/protection-and-response/animal-welfare/national-animal-ethics-advisory-committee/ Applicants can vary the content of the CEC to suit their organisation, while maintaining a consistent format, and meeting the requirements of the Act. Having a consistent format will assist the Director-General and NAEAC when assessing applications.

A checklist to ensure all important points are covered can be found in Appendix 1 – CEC Checklist. This (Good Practice) guide contains additional information useful to the person drafting a CEC.

4.3. Applying for a CEC

The following documents must be submitted to the Director-General of MPI:

• a draft CEC;
• a completed application form;
• independent references as specified in section 89(2)(b) of the Act.

If the application is for a renewal of an approved CEC, a report from an accredited reviewer must have been submitted to MPI and NAEAC by the reviewer.
4.4. Consideration of an Application to Approve a CEC

The Director-General must consult with NAEAC on each application for a CEC. The Director-General must notify the applicant of the decision within 40 working days of the application being lodged. This can be extended to 80 working days if the Director-General requires more information or needs to consult with the applicant. If the Director-General does not notify the applicant of the decision within the required time period, the Director-General is deemed to have refused to approve the proposed code.

4.5. New AECs – Where to Get Assistance

When a new AEC is being formed, NAEAC will, on request, arrange assistance from a local NAEAC member. This is intended to ensure the new committee:

- develops procedures that are both appropriate for the nature of RTT involved, and compliant with the Animal Welfare Act; and
- is familiar with the literature available to provide support.

Generally, the NAEAC member would attend an AEC meeting as an observer, for the purpose of advising, and may be able to arrange for members of the new committee to attend a meeting of another AEC to gain further experience (this is considered to be particularly valuable for ‘external’ members of AECs). Such a visit is made in an advisory capacity only and will not fulfil any review function.

This is an area in which NAEAC can provide valuable assistance to AECs, establishing a personal connection that will have long-lasting benefits.

4.6. Undertaking RTT without a CEC

Organisations or individuals that, for whatever reason, elect not to have their own CEC may arrange to use the CEC and AEC of an organisation that does have them. This is termed a parenting arrangement. It is the code holder’s decision as to whether external parties will be allowed to do this.

NAEAC recommends that where code holders do not permit parenting arrangements, they state this in their code.

If the code holder intends to allow use by external parties the following expectations apply:

- the CEC must detail the policies and procedures it will implement to manage such arrangements;
- the code holder must ensure that members of its AEC are qualified to evaluate all projects submitted by a parented organisation;
- the code holder must be satisfied that the personnel who will manipulate animals on behalf of that parented organisation are appropriately qualified;
- the AEC must have a review process in place for all relevant operating SOPs of the parented organisation, such as those used to manage animal facilities;
- AEC responsibilities for monitoring and oversight described below, apply equally to the work and facilities of parented organisations;
- The code holder may wish to appoint a nominee of the parented organisation to the AEC.
Note: The code holder must notify MPI of the arrangement in writing and before the external party begins any RTT using animals. Notification should include the (registered if appropriate) name of the person or organisation, postal and physical addresses of the parented organisation and contact details for the person to whom correspondence should be addressed.

4.7. Amendments, Suspension or Revocation of the CEC

4.7.1 Amendments

Establish a procedure to enable the AEC to recommend amendments of the CEC to the code holder. When it is intended to amend the code:

- **Minor Amendments** – must be notified to MPI annually (and no later than 31 March after the year in which they were made).
- **Significant Amendments** – must be approved by MPI, prior to implementation.
- **Suspension or Revocation** – an organisation, under section 95 of the Act, may elect to suspend or revoke its CEC if it decides to cease manipulating animals for RTT.

4.7.2 Suspension and Revocation

Irrespective of an application under section 95, the Director-General has the power to revoke a CEC as a response to a conviction under:

- the Animal Welfare Act 1999;
- the Animals Protection Act 1960;
- the Agricultural Compounds and Veterinary Medicines Act 1997;
- the Biosecurity Act 1993;
- the Companies Act 1993;
- the Crimes Act 1961;
- the Dog Control Act 1996;
- the Serious Fraud Office Act 1990;
- the Trade in Endangered Species Act 1989;
- the Veterinarians Act 2005.

The Director-General also has the power to suspend or revoke a CEC if the code holder:

- is no longer carrying out research, testing, or teaching or no longer wishes to enable research, testing, or teaching to be carried out by another person;
- no longer has the capability and skills necessary to carry out research, testing, or teaching or to enable research, testing, or teaching to be carried out by another person;
- has failed to comply in a material respect with the Act or any regulations made under the Act or the code of ethical conduct;
- has provided in or with the code holder’s application under section 87 information that was false in a material respect.

In either case, the welfare of animals under a current approval must be maintained and each approval must either be terminated or submitted to another AEC for approval.
4.8. CEC Expiry and Statutory Review

Approved CECs expire after a maximum of 5 years. In order to renew a CEC, a statutory review of the CEC and the AEC’s compliance with it, is required during the fifth year.

Key components of the review are:

- the review is conducted by an MPI-accredited reviewer who is independent of the organisation and MPI;
- the reviewer audits and reports on compliance with the Act and CEC; including reviewing how the AEC(s) constituted under the CEC operate;
- a formal report is provided to MPI and NAEAC;
- NAEAC reviews the report, assesses the adequacy of the submitted CEC and makes a recommendation to the Director-General of MPI, as to whether a new code should be approved.

The statutory review is a key part of the RTT control process. Evidence of AEC activities (e.g. minutes of meetings and monitoring information) provides a tangible basis on which code-compliance can be assessed, and consequently contributes to the process by which regulators, the Minister and the public are assured of the ethical scientific use of animals.

Any recommendations that the reviewer makes relating to the CEC should be considered when revising the CEC. The new CEC must be submitted to MPI. Submission must allow sufficient time for the review of the existing CEC to be assessed and the new CEC considered by NAEAC, within a timeframe that allows the new CEC to come in to force immediately after the previous code expires. If this is not done in a timely manner, the code holder risks having no current CEC, and therefore will be unable to undertake any animal manipulations until such time as a new code is approved.

It is NAEAC’s advice to have the review completed, and a new CEC drafted and submitted at least 4 months prior to expiry of any current CEC.

4.9. Animal Ethics Committee (AEC)

The use of animals under section 6 of the Act must be approved by an AEC. A code holder is permitted to set up one or more AECs to consider applications to carry out RTT.

Key responsibilities of an AEC are:

- considering applications to use animals for RTT;
- setting appropriate conditions on approved projects;
- monitoring approved protocols;
- approving & monitoring adherence to specific operating procedures (SOPs);
- reviewing the results of approved projects;
- monitoring of animal facilities used for RTT;
- monitoring routine animal husbandry and welfare.

The AEC has the power to inspect animals, their accommodation, and related experimental records at any time to satisfy itself that approved procedures are being properly carried out.

These responsibilities extend equally to all projects and facilities approved by the AEC.
Part 6 of the Act also sets out criteria that the AEC must consider when reviewing proposals. This includes examination of the opportunities for refinement of the method, reduction in animals used and replacement of animals with another process (the Three Rs).

### 4.9.1 Jurisdiction of AECs

**Authority is within New Zealand only**

AECs are constituted under the Act. This legislation has force only within New Zealand (which includes New Zealand’s territorial waters). An AEC cannot approve a project in a jurisdiction where it is unable comply with all relevant parts of the CEC under which it operates.

AECs must not issue formal approvals for work performed wholly in another country. However, where projects involve work undertaken both within and outside NZ jurisdiction, only animals used within NZ must be reported to MPI in annual statistics.

**Animal Work Performed across Several Code Holders**

Routinely, the person wishing to undertake RTT should apply to the AEC associated with their host institution. However, situations sometimes arise where more than one organisation is involved in a RTT project. For example:

- where a student from one organisation intends to carry out RTT with a second organisation;
- where two organisations work together on a project involving RTT;
- where one organisation (that has its own AEC) intends collaborative or contracting work to a second organisation that also has an AEC;
- where several AECs are established under a single CEC.

The AEC that is best placed to assess the application according to the criteria in s100 and monitor any approval in accordance with s99 of the Act must be the AEC which considers the application. This would usually be the organisation that is the primary host for the RTT. This AEC must be advised of names, affiliations and suitability of all researchers involved in the project including any not employed by the primary host.

Where a protocol is going to involve activities occurring at more than one organisation associated with an AEC, and there is no obvious primary host institution, the investigators may choose which AEC they use. It may help to examine the CEC from each organisation, looking for the best fit with respect to expertise of the AEC.

For all applications to an AEC, the applicant must be a suitable person with respect to s82 of the Act and the CEC for that AEC must allow for the arrangement.

### 4.10. AEC Membership

#### 4.10.1 Statutory Members

1. A senior member of staff, capable of evaluating the scientific value of proposals in accordance with s101(3) & (4) of the Act.

2. Veterinarian nominated by the New Zealand Veterinary Association. This person must not be in the employ of, or otherwise associated with, the code holder in accordance with s101(5) of the Act.
3. Nominee of an approved animal welfare organisation. The Royal New Zealand Society for the Prevention of Cruelty to Animals is the only approved organisation in New Zealand. This person must not be in the employ of, or otherwise associated with, the code holder in accordance with s101(6) & (7) of the Act.

4. Lay person nominated by the local territorial authority or regional council, not associated with the scientific community in accordance with s101(8) & (9) of the Act.

The phrase ‘associated with the scientific community’ is not easily defined and must be considered on a case by case basis. However, the following points should add clarity:

- the aim is to appoint a lay member whose views represent an unbiased member of the public;
- avoid appointing anyone whose present or past occupations could lead to public perceptions that scientific, financial or philosophical biases might influence the person’s role on the committee;
- some understanding and knowledge of science in a general context would not necessarily imply that the person was associated with the scientific community, and would clearly assist the appointee with understanding the complexities of RTT project applications.

If unsure of the suitability of a potential appointee, seek advice from NAEAC.

The Importance of Statutory External Members
The Act mandates a minimum of three non-institutional members on each AEC. Statutory external members bring different skills, but are all intended to bring credibility & transparency to RTT in New Zealand. These outcomes are maximized when the members fully participate in the decision process. The views of external members should be sought in each case, and fully considered.

Remuneration
Statutory external members should be remunerated for the time spent on AEC business. Refund of travel and other expenses is also expected.

4.10.2 Additional Members
Additional members may be appointed to the AEC by the code holder. Additional appointees should bring useful expertise to the committee.

4.10.3 AEC Chair
The code holder decides on how the Chair and Deputy Chair will be determined. The code holder may appoint them directly or allow the AEC to choose them. The process may be described in the Code.

4.10.4 Period of Appointment
The period of appointment for each type of AEC member should be specified in the CEC. It is useful to include the eligibility criteria for reappointment in the code.

4.10.5 Induction of New Members
A formal process should be followed to induct new members including at least:
- a copy of the CEC
- the NAEAC induction pack
• a summary of the key AEC approvals open at the time;
• encouragement to attend NAEAC workshops.

4.10.6 Protection of Members

Section 104 of the Act states no member of an AEC is personally liable for any act done or omitted by the member or the committee in good faith in the course of the operations of the committee.

4.11. AEC Meetings and Decision Making

NAEAC expects that all new applications and amendments to existing projects will be considered at face to face meetings (see section 4.11.8). However, NAEAC accepts that it may occasionally be necessary to consider a new application under urgency, and sometimes desirable to make amendment decisions between meetings as part of refinement, and thus consistent with the Three Rs. The process for managing such contingencies should be described in the CEC.

4.11.1 Public Access to Meetings

Some public organisations are subject to the Local Government Official Information and Meetings Act 1987. In this case, meetings must be open to the public, unless there is good reason for their exclusion. The committee must apply section 48(1)(a)(ii) of the Local Government Official Information and Meetings Act. This means sections 6, 7 and 9 of the Official Information Act 1982 apply including commercial sensitivity and legal professional privilege. However, section 9(2)(g)(i) of the Official Information Act (which relates to free and frank expression of opinion between members of an organisation) is not available.

4.11.2 Timing for Circulation of Agenda Items

There is a need to provide members of an AEC adequate time to assess any application. The CEC should state the minimum period allowed (prior to the meeting) for circulation of materials. Phrases such as ‘well in advance’ are vague and not suitable.

4.11.3 Frequency of and Attendance at Meetings

Meeting frequency should be stated. A minimum of one face to face meeting a year should be held to maintain continuity and the relationships between AEC members. Even if there are no applications to consider, there will be monitoring reports to consider, or monitoring visits to carry out.

The CEC should also document the expectation on members for meeting attendance.

4.11.4 Quorum

The requirements for a quorum must be clearly documented. NAEAC recommends that a quorum is at least 50% +1 of the membership, and must include at least two of the statutory external members.

4.11.5 Vacancies

The CEC must detail how vacancies and unexpected/prolonged absences are managed. While it appreciates the difficulties that lack of a quorum can pose, NAEAC requires CECs to detail how
this will be managed. If there are insufficient members present to constitute a meeting then NAEAC considers that those members present could, in effect, act as a subcommittee and be governed by the process outlined for subcommittees in the CEC. If vacancies render the AEC inquorate this would invalidate its actions. The CEC should outline how this will be avoided.

4.11.6 Consensus

NAEAC strongly recommends that decisions are made by consensus, with applications revised until all members are satisfied. If consensus cannot be reached the default should be to reject the application.

4.11.7 Conflict of Interest

It is suggested that CECs include direction as to how all conflicts of interest are declared, managed and recorded.

Members of AECs are expected to perform their functions in good faith, honestly and impartially, and to avoid situations that might compromise their integrity or otherwise lead to conflicts of interest. At the start of each meeting, a general declaration of whether conflict of interest exists or not, should be made by each member and held on record. It is recommended that members err on the side of caution in disclosing perceived or actual conflicts of interest to ensure transparency.

Members are expected to declare to any areas (including research or any other matter before the committee) where they believe there is a potential conflict of interest. This is to be raised with the AEC Chair in advance of any committee work on a matter where a conflict of interest could occur, or be perceived to occur for whatever reason. AEC has the discretion to consider how the conflict of interest is to be handled on a case-by-case basis, with members given an opportunity to provide suggestions on how any conflict could be handled.

As a minimum, the AEC membership should be advised of the conflict of interest and committee records should reflect the expression of a specific conflict of interest, what it is in relation to, and how the conflict is being handled.

Additional remedies, when the matter relating to the conflict of interest is discussed in committee, could include:

- the member with the conflict of interest participating in discussion but not participating in any vote or final decision; or
- the member with the conflict of interest not participating in discussions, but may observe discussions and answer questions; or
- the member with the conflict of interest withdrawing from any discussions or deliberations.

Where the Chair has the conflict of interest, then the Deputy Chair should assume the chair for the duration of the matter where conflict has been declared.

NAEAC members can provide guidance or training on the management of conflict of interest to AEC committees, where requested. Further guidance on conflicts of interest is available at: [http://www.oag.govt.nz/](http://www.oag.govt.nz/).

Accredited reviewers will review conflicts of interest as part of their normal auditing process, therefore AECs should record adequate documentation of declarations for this purpose.
4.11.8 Appropriate Use of Teleconferences and Videoconferences

AECs should have a policy on the use of teleconferences and videoconferences in their decision-making process.

NAEAC accepts that with adequate software, videoconferences can be considered the equivalent of face-to-face meetings for the purposes of considering new applications. All meeting procedures must be adhered to.

While circumstances may require inclusion of one or more (but always limited to a minority of) members by teleconference in order to ensure a quorum, NAEAC believes that face-to-face meetings for assessment of applications allows for greater robustness of debate on issues arising out of applications, particularly those in which higher impact manipulations are to be undertaken.

Accredited reviewers will review the use of teleconferencing as part of their normal auditing process, therefore, AECs should record adequate documentation of formation, discussion and decisions made in such circumstances for this purpose.

4.11.9 Consideration between Meetings and Subcommittees

The CEC should indicate how decisions will be made between meetings. AEC members may allow a subcommittee to make decisions on their behalf. If subcommittees are used, procedures should describe their appointment and function.

For consideration of a new application, the membership of a subcommittee must always include at least two of the statutorily appointed external members. Decisions made by a subcommittee are conditional on ratification at the next AEC meeting. NAEAC advises that interim decisions should only be made for fast-tracking studies that:
- have a legitimate requirement for urgency; and
- are for manipulations of grades A and B only.

Any fast-tracking procedures must not circumvent proper consideration of the application.

For consideration of amendments to an existing application, the CEC must state who will have delegated authority to make such decisions. Any subcommittee must always include at least one of the statutorily appointed external members. A subcommittee review is appropriate for alterations to current studies where:
- the changes do not involve a major departure from the approved study design;
- there is no change to the impact grading;
- any request to change numbers is the minimum necessary to retain the statistical validity of the original approval. (Larger changes in numbers and any increase over 10% of the original number requested must be agreed by a quorum of the committee).

Accredited reviewers will review the consideration between meetings as part of their normal auditing process, therefore AECs should record adequate documentation of discussion and decisions made in such circumstances for this purpose.

4.11.10 Confidentiality

CECs should describe how confidentiality will be managed. Commercially sensitive information may be included in applications to the AEC. Applications must contain sufficient information so the AEC can ensure that the review criteria set out in the Act are met.
4.11.11 Applicant Presence at Meetings
The code should describe whether an applicant may be present or not.

4.11.12 Secretarial Support
The code should describe how the AEC is supported with secretarial functions.

4.12. Applications to the AEC

Applications should be on a standard form to ensure no important information is missed. This will allow the AEC to maintain consistency in its consideration of the criteria specified in s100 of the Act. The application template (Appendix 2) includes the criteria NAEAC considers central for AECs when considering applications. AECs in New Zealand deal with animal models and facilities ranging from rodents in cages, to herds of livestock on pasture, to populations of animals in the wild. It is acknowledged that not all categories in the template will be applicable for all proposals.

The Act requires that RTT will be undertaken only by suitably qualified persons. The application shall detail the qualifications of all key personnel involved in the project.

4.12.1 Criteria for Consideration of Applications

The key principle underlying the assessment process is that the benefits that are likely to be derived from using animals in RTT must outweigh any foreseeable harm to those animals. The CEC should therefore contain policies and procedures to ensure that relevant criteria (below) are considered both by project applicants in writing applications, and by the AEC in evaluating applications. This information forms the basis for assessing likely benefit: harm of the work.

The CEC should describe the criteria used in the consideration of applications to the AEC. The Act specifies that in considering any application for approval of a project and in setting, varying or revoking conditions of approval, every AEC shall have regard to:

a. the scientific or educational objectives of the project; and

b. the harm to, or the distress felt by, the animals as a result of the manipulation, and the extent to which that harm or distress can be alleviated by any means (including, where the pain or distress cannot be held within reasonable levels, the abandonment of the manipulation or the humane destruction of animals).

4.12.2 Impact Grading

The AEC should document how it will assess impact grading of proposed RTT. It is recommended that AECs adopt the MPI grading guidelines.

4.12.3 Development of Standard Operating Procedures (SOPs) for Manipulation of Animals

NAEAC considers that the use of SOPs for routine manipulations will minimise technique ‘drift’ and the welfare impact of such manipulations. The AEC must have a review process in place for all SOPs used in the conduct of an RTT project.
4.12.4 Avoiding Duplication of Animal Research

Scientific experiments involving animals are sometimes repeated by the same or other research groups. In considering whether the repetition of experiments is desirable, it is essential to distinguish between ‘replication’, ‘repetition’, and ‘duplication’.

Replication refers to repetition of experiments or tests to increase the reliability (by replication within experiments) and generality (by replication of entire experiments) of the findings. Replication is a fundamental component of the scientific method. The degree of replication required will depend on the level of scientific precision demanded, the natural variation in the variables measured, and the range of circumstances in which findings are to be applied. The level of replication required to achieve the desired statistical precision can be predicted by power analyses of pilot data.

Repetition of product testing is sometimes required by regulatory agencies. Such repeat testing may occur:

• where routine testing of different batches of products is required;
• where product stability or shelf life is being determined;
• where ‘generic’ companies commence manufacture of products coming off-patent and are required by regulators to provide evidence of efficacy and safety using animal models, and
• where the claimed efficacy of products is extended (e.g. an existing parasiticide is shown to be effective against an additional parasite).

All cases entail, either or both, novelty and uncertainty. Therefore, repetition of the animal tests is necessary to ensure that products are fit for purpose (i.e. both efficacious and acceptably safe).

Duplication of animal experiments, where neither replication nor repetition (as defined above) are being carried out, would be pointless. If the outcomes are predictable from previous experiments, such duplication is unacceptable because animals are needlessly manipulated. However, it could occur when a researcher, and those scrutinising the researcher’s proposal, are unaware that the experiment or test has already been done because results have not been published. This may be because:

• the research ‘failed’ and was not considered worth publishing, or
• the research findings were commercially sensitive, or
• the research is still in progress.

The Animal Welfare Act 1999 permits ‘duplication’ only where the original experiment was found to be flawed, or for confirmation (i.e. as replication or repetition).

The following guidelines should be followed to avoid needless duplication of animal use:

• AECs need to be aware of the important distinction between ‘replication’, ‘repetition’, and ‘duplication’ as defined above.
• AECs need to avoid approving research that involves ‘duplication’ unless previous studies were flawed.
• Researchers should be asked by AECs to provide evidence, in research applications, of the efforts made to avoid duplication of past research. Evidence could include for example: the literature and patent databases searched including keywords used, and reference to previous searches.
• Researchers should be asked to consider the likelihood that the research is currently being undertaken elsewhere, and if so, to make every effort to ensure they are aware of, and not duplicating such research.
• Researchers should be encouraged by AECs to follow the ARRIVE Guidelines (see: https://www.nc3rs.org.uk/arrive-guidelines) when reporting the results of in vivo animal research.

4.12.5 Outcomes of Consideration

The CEC should state clearly the possible outcomes of the consideration process. These are commonly:
• approved with conditions;
• returned to the applicant for revision;
• rejected.

4.12.6 Conditions of Approval

The AEC may set conditions of approval and may vary or revoke such conditions (s99(1)(c)). Such conditions may include matters such as:
• the time period for which approval is granted;
• reporting requirements of project applicants to the AEC;
• monitoring requirements – See Occasional Paper Nos 4 & 5;
  – “Compliance monitoring: The University of Auckland approach”;
  – “Monitoring methods for animal ethics committees”.

4.12.7 Maximum Approval Period

The CEC should set the maximum approval periods for all types of AEC approvals. The CEC should specify that no animal shall be manipulated unless under a current approval.

4.12.8 Power to Suspend/Revoke/Vary Approvals

The AEC has the power to set, vary or revoke conditions of project approval and to suspend, or revoke approvals if required. The CEC should specify grounds for how these decisions may be made and how the processes are managed.

4.12.9 Compliance with Other Legislation

The AEC should be provided with evidence that, for any activity which must be approved by other regulatory authorities, such approval has been obtained.

This includes:
• confirmation that advice from the institution’s biological safety committee or equivalent (where it exists) has been sought and that appropriate measures for containment, disposal and decontamination have been established;
• evidence of compliance with the organisation’s Institutional Operating Plan (IDAOs etc).
5.1. Compliance Reporting

A procedure should be established to allow:

- staff members to raise a concern over the conduct of work;
- reporting of non-compliances;
- reporting of deviations from the intended protocol.

This provides a structured means of informing the AEC and key staff when things do not go to plan. Non-compliance may be considered justifiable in hindsight; on other occasions there may be a need to make changes to how work is conducted. The aim should be to firstly consider the action that may be needed to address any animal welfare concerns, and secondly to address procedural and personnel matters based on a clear understanding of the nature of and reasons for non-compliance. Serious cases of non-compliance should be addressed by disciplinary procedures, as determined by management of the host institution in conjunction with the AEC.

5.2. Project Monitoring

Monitoring includes a range of activities undertaken by various members of the committee and/or the Animal Welfare Officer. As a minimum NAEAC expects the AEC to actively monitor at least 10% of all projects graded A or B and all projects graded C-E.

Accredited reviewers will review monitoring of projects as part of their normal auditing process, therefore AECs should record adequate documentation for this purpose.

Monitoring may include the following monitoring activities:

- Observations at unscheduled times i.e. not premeditated for compliance monitoring. This could be conducted by any member of the AEC who happens to be visiting animal holding facilities and observes animal practices as an unintended consequence of their primary activity. Examples of this type of visit include, but are not limited to, a visit to an animal facility by an AEC member in their capacity as an investigator undertaking their own manipulations, the animal facilities member during their management activities or the Animal Welfare Officer (AWO) during a scheduled training session. Veterinary visits and facility checks lead to the most frequent unplanned observations as the AWO walks past investigators performing manipulations.

- Veterinary care: each facility must have an approved procedure for reporting of sick (ill, injured, distressed) animals – conditions which the investigator may not have anticipated during the planning phase and does not have covered by their approved monitoring and contingencies in their approved protocol. Veterinary care of animals with conditions which have not been anticipated during the planning phase ensures that information on models is transferred back to the AEC and can, often, be “fed back” into the model to improve and refine the model, reduce variability and ultimately perhaps animal usage.
A structured post-approval review of procedures relating to projects should be developed. This should include conditions set by the AEC at the time of approval for new models, new procedures and new personnel. This aspect of post-approval review focuses more on individual procedures or components of an approval, rather than review of the whole protocol. It ensures that new models have been set up properly and that personnel are trained appropriately for the work at hand, especially for new models.

- Reviews of projects and experiments where:
  - numbers of animals for which approval is given are large;
  - an investigator has been liable for welfare incidents or breaches of approval. If any such incidents occur this will prompt immediate review as part of the non-compliance procedure and then followed up in 12 months.

5.3. Interim Reports

Where projects run over a number of years, it is recommended that the lead investigator submits an annual interim report. This will allow the AEC to remain engaged and familiar with the progress of the project. Contents should at least:

- summarise the aims and projected benefits of the project;
- state the number of animals of each species approved;
- restate the expected grading for animal impact;
- summarise any variations to the original project approved by the AEC;
- summarise any deviations, non-compliances, or adverse events that have occurred to date;
- report the actual number used, and the actual impact grading;
- summarise the current status of the project, and any results in hand.

An AEC may also request interim reports during the course of an approval as a means of monitoring the progress of a new manipulation, or the ongoing welfare of the animals.

5.4. Changes to Approved Applications

There should be a policy to manage variations to approved projects. The process to handle minor and significant variations may vary, in which case they should be clearly stated. In the case of minor modifications this could be done by an AEC.

Accredited reviewers will review approval amendments and the process used as part of their normal auditing process, therefore AECs should record adequate documentation for this purpose.

5.5. Adverse Event Reporting

Adverse events (AEs) are unanticipated or atypical incidents that occur to an animal as a result of experimental manipulation, routine husbandry or diseases. Where unexpected adverse events or outcomes occur during RTT, rapid reporting is essential, primarily from the point of view of animal welfare.

Basic components of an AE system are:

- a system should be in place to record details of the AEs;
- reporting timelines should be established. These may vary depending on how serious the event was.
• project leader and AEC should be notified;
• actions in response should be determined. These could range from additional observations, review of procedure, through to termination of the project.

5.6. End of Approval Reports

An important step in the approval cycle is the examination of results of approved projects. This allows the AEC to determine whether the proposed benefits of the work were realised. The best way to achieve this is to require investigators to submit an end of project report. This report should:
• summarise the aims and projected benefits of the project;
• compare the number of animals of each species approved with the number actually used;
• compare the expected grading for animal impact with actual grading;
• summarise any variations to the original project approved by the AEC;
• summarise any amendments, non-compliances, or AEs that occurred during the project;
• report the actual number used, and the actual impact grading;
• summarise the results and actual value realised by the project;
• indicate if any further work is being considered.

Reports should be submitted within 3 months of the trial conclusion. If the AEC meets infrequently, reports should be reviewed either by the Chair or a subcommittee in a timely manner. The reviewer(s) should call a meeting of the full committee if matters of concern are identified.

Organisations should consider adopting the ARRIVE guidelines for improved reporting of research using animals (https://www.nc3rs.org.uk/arrive-guidelines).

5.7. Non-compliance

The CEC should detail processes for dealing with non-compliance with any AEC approved protocol, including Standard Operating Procedures.

5.8. Complaints Procedure

The code holder must ensure that there are procedures to deal promptly and fairly with complaints.

Procedures should cover:
• complaints from AEC members, staff members or members of the public;
• complaints against the Chairperson.

The procedure should specify how complaints, and the outcomes of the complaints process are recorded and stored.
5.9. Record Keeping

The code holder must retain records relevant to its approvals, operations and functions under the Animal Welfare (Records and Statistics) Regulations 1999. These records form part of the statutory review of CEC and must include a record of the actual “impact” of the manipulation on each animal used.

Records kept should include (but not limited to) project applications, approvals, policies, project monitoring, procedures, site visits, and complaints log. The CEC should indicate how these records are managed and stored.

5.10. Reporting of Statistics to MPI

Every year the code holder must inform MPI of the numbers of animals it has used in RTT. Returns are required to be submitted by 28 February for the preceding calendar year. For details, see the MPI document: “Animal Use Statistics – Guidance for Completing Statistical Returns” https://www.mpi.govt.nz/dmsdocument/1477
Facilities include the buildings, cages, tanks, aquaria, yards or paddocks in which animals are kept. Investigators, AECs and the institutions must ensure that facilities are appropriately designed, constructed, equipped, staffed and maintained to ensure the health and welfare of the animals and to fulfil scientific requirements.

The design and management of facilities will depend on the type of animals to be kept and the experiments to be undertaken. The overall condition and management of facilities should permit effective maintenance and servicing and be compatible with maintaining good health and welfare.

### 6.1. Monitoring of Animal Facilities

Under section 99 of the Act, a key function of the AEC is to monitor animal management practices and facilities to ensure compliance with the terms of the code of ethical conduct. This applies also to any facilities operated by an organisation parented by an AEC.

Accredited reviewers will review monitoring of animal facilities as part of their normal auditing process, therefore AECs should record adequate documentation for this purpose.

#### 6.1.1 Scheduled Visits

AEC members should visit each animal facility every year. These visits should be programmed and scheduled. The purpose and scope should be clear. This may vary from, for example, inspection of a specific aspect of animal husbandry practice or the adequacy of a particular building, to a complete assessment of all practices and facilities. Complete assessments are probably most beneficial at a point midway between scheduled statutory reviews. A checklist should be used.

#### 6.1.2 Non-scheduled Visits

Experience shows that a collaborative approach between the AEC and facility staff is key to genuine transparency in the use and care of animals. Non-scheduled monitoring visits should be considered where an AEC holds unresolved concerns over the conduct of any RTT.

#### 6.1.3 Periodic Review of SOPs

The scientific body of knowledge underpinning animal management practices is constantly expanding. A periodic review of SOPs is required to allow incorporation of improvements. Typically, review occurs at 3-year intervals. However, in rapidly evolving areas, or protocols with very high ethical cost, more frequent review may be appropriate.

#### 6.1.4 Animal Carer on the AEC

The most direct means for the AEC to monitor the day-to-day operation of an animal facility is through the membership of an animal carer of the host institution on the committee. This enables the AEC to gain insight into the culture, commitment, capability and effectiveness of the staff responsible for animal welfare. Many AECs have a regular part of meetings devoted to discussion of items raised by the animal care representative.
6.2. Facility Personnel

6.2.1 Facility Manager

Animal facilities used for RTT should be supervised by persons with appropriate veterinary or animal care qualifications or experience with the species involved. The facility manager should have ready access to institutional or consultant veterinary services seven days a week.

The facility manager should:

- ensure that veterinary care is available at all times, for animals held for both breeding, and for experimental manipulations;
- be responsible for the management of the day-to-day care of the animals;
- supervise the work of other personnel in the facility;
- liaise between investigators, teachers, and facility personnel;
- communicate with the AEC on facility management, and any adverse events;
- continually develop the institution’s animal care policies and procedures. See Standard Operating Procedures (SOPs) section 6.5;
- be knowledgeable regarding signs of pain, distress and illness specific to the species housed;
- ensure the well-being of all animals is regularly assessed. (After animals are allocated to an approved project, the investigator has primary responsibility for ensuring adequate monitoring of the animals’ well-being);
- ensure that all ill or injured animals are treated promptly;
- investigate the cause of all adverse events including when animals die unexpectedly;
- provide appropriate protective clothing to facility personnel;
- enforce high standards of personal hygiene, and ensure staff do not eat, drink or smoke in animal areas;
- establish an effective cleaning/sanitation schedule for housed animals;
- ensure that the equipment and facility is clean and well maintained;
- keep adequate records of:
  - the source, transport and use of animals received;
  - health status, genetic constitution and the physical environment of animals held, when definition of these is required;
  - fate or disposal details of all animals held;
  - fertility, fecundity, morbidity and mortality in animal breeding groups;
  - diseases developed and treatment provided.

Records related to the use of animals in RTT must be made available to accredited reviewers, the AEC, and its nominated representatives.

6.2.2 Personnel

Having sufficient numbers of well-trained, committed staff is key to achieving high standards of animal care.

Institutions should encourage and promote formal training in animal science or technology.

Personnel employed in the care of animals should:

- be trained in the care and maintenance of those animals;
- understand how their actions will affect the animals’ well-being, and the outcome of experiments;
• be able to recognise early stage changes in animal behaviour, performance and appearance;
• have clear instruction in their duties, and in institutional policy and procedures;
• have their training documented.

Personnel should be informed of allergy hazards and the important zoonotic diseases of animals under their care and of precautions that should be taken. Regular health checks and appropriate immunisation (e.g. against tetanus and other zoonoses) of all personnel who handle animals are recommended in the interests of both personnel and animals.

### 6.3. Routine Monitoring of Animal Health

As a minimum, NAEAC expects that all animals housed in a facility undergo daily visual health checks and more thorough assessment checks at least once a week as part of routine husbandry.

The advantages include:
• prevention of unnecessary impact on animal welfare;
• ensuring scientific data is not compromised by animals behaving or functioning abnormally;
• avoiding costly and disruptive disease outbreaks.

Animal carers must be appropriately trained, and capable of designing and implementing the monitoring programme. There is a large body of literature to assist this process.

### 6.4. Food and Water

As a minimum:
• provide species-appropriate, clean and nutritionally correct food;
• provide sufficient food to allow normal growth of immature animals, the maintenance of adult animals, and the requirements of pregnancy or lactation;
• remove uneaten perishable food promptly unless contrary to the needs of the species;
• alteration to dietary regimes should be gradual;
• allow sufficient trough space or feeding points when feeding groups. This will avoid undue competition for food which is especially important if feed is restricted;
• supply clean, fresh, and uncontaminated drinking water continuously.

### 6.5. Standard Operating Procedures (SOPs)

Facility management should develop SOPs to enable consistent performance of routine tasks. Where they relate to small animal holding and breeding facilities for animals used in RTT, SOPs should be submitted to the AEC for consideration.

Facility SOPs should cover:
• animal husbandry;
• animal transport;
• facility sanitation;
• health checking, disease diagnosis and treatment;
• breeding colony management;
• restraint and manipulation of animals;
• emergency management;
• euthanasia;
• reporting of non-compliance and adverse events;
• staff health and safety.

Where available, MPI codes of welfare should be followed, unless alternative procedures have been approved by the animal ethics committee.

SOPs should be reviewed regularly and resubmitted to the AEC if amended.

6.6. Acquisition of Animals

Animals should be obtained from breeding and supply facilities that maintain conditions consistent with this guide and/or a relevant industry code.

6.7. Animals Collected from their Natural Habitats

Most species of indigenous fauna are protected by law. The Department of Conservation must be consulted when these species are required. Permits are usually necessary to collect, keep, release or kill protected fauna, and further permits are usually required to import or export such species. Any conditions imposed on permits must be observed.

Endangered species must not be used unless the research will be of direct benefit to the conservation of that species or a closely related species and will not further endanger the species.

Animals should be taken from natural habitats only if animals bred in captivity are not available or are unsuitable for the specific scientific purpose.

The definition of a manipulation specifically excludes (in s3 (2) (d) of the Act) the hunting or killing of any animal in a wild state by a method that is not an experimental method. Nonetheless, capture and restraint is clearly stressful. Distress felt by the animals will varying depending on the species, physiological state and experience of contact with humans. Strategies should be employed to minimise distress during capture such as:

• careful choice of capture techniques;
• use of skilled operators;
• choice of appropriate and safe enclosures, devices or caging;
• monitoring for signs of distress following capture;
• treatment of any capture-induced trauma.

Where live traps are used, their operation must comply with section 36 of the Act, which states:

• traps must be inspected within 12 hours after sunrise every day beginning on the day immediately after the day on which the trap was set or, where a reliable remote monitoring system is used, be inspected within 24 hours after an animal is captured;
• at each inspection any living animal must be removed.

Fish may be caught using commercial harvesting practices.
6.8. Animals Obtained from Other Countries

Under the Biosecurity Act 1993, the exit and entry of animals or animal tissues can be restricted. Permits must be obtained from MPI for the importation of live animals, and their genetic material.

The IBSC must be consulted when genetically modified animals are to be imported and approvals obtained from EPA through the institutional committee. The housing and use of GM organisms and animals requires specialised transitional and containment facilities for vertebrate laboratory animals. MPI approval must first be obtained for the establishment of transitional/containment facilities. This approval requires an on-site inspection by MPI inspectors.

Permits must be obtained from MPI for the importation of specimens from dead animals.

Permits must be obtained from the Department of Conservation for the import/export of both live and dead specimens of all native New Zealand fauna and animals or plants subject to the Trade in Endangered Species Act 1989 or regulations.

6.9. Transport of Animals

Transportation can cause distress due to confinement, movement, noise and changes in the environment and personnel. The extent of any distress will depend on the animals’ health, temperament, species, age, sex, the number travelling together and their social relationships, the period without food or water, the duration, the mode of transport, environmental conditions (particularly extremes of temperature) and the care given during the journey.

The CEC should require that animals are transported under humane and hygienic conditions at all times. The Transport within New Zealand Code of Welfare can be found at the following link: https://www.mpi.govt.nz/document-vault/1407

6.10. Animal Housing

All animal housing should provide a safe and appropriate environment for the animals held.

A documented pest control programme should be in place to monitor and control vermin.

The sections below provide an overview of requirements. For greater detail refer to a recognised reference such as:

- the Universities Federation for Animal Welfare (UFAW);
- the Guide for the Care and Use of Laboratory Animals (The Guide);
- Canadian Council on Animal Care: Guide for the Care and Use of Experimental Animals.

6.11. Outdoor Holding Areas

Outdoor area for housing animals should:

- be compatible with the needs of the species;
- provide adequate shelter and water;
- protect the animals from predation;
• meet any species-specific needs;
• comply with established farm or zoological garden practice.

6.12. Indoor Housing

Buildings for the indoor housing of animals must:
• be compatible with the physical, health and behavioural needs of the species and stage of animal to be housed;
• suit the project undertaken;
• allow for free movement;
• provide for group contact where this is appropriate;
• be designed and operated to supply environmental factors suitable to the species;
• prevent cross-contamination between different groups/types of animals;
• allow for the delivery and adequate storage of food, water, bedding and appropriate enrichment;
• facilitate the entry of people and other animals;
• restrict access to the animal quarters to authorised persons only;
• be maintained in good repair;
• have durable, impervious and easily sanitised walls;
• be kept clean and tidy, and operated to achieve maximum possible hygiene;
• allow for effective and humane pest control;
• have a reticulated water supply and proper facilities for drainage.

6.13. Environmental Factors

Provide environmental conditions suitable for the behavioural and biological needs of each species held. Consider:
• air exchange;
• temperature;
• humidity;
• noise;
• light intensity and duration.

Effective ventilation is essential for the comfort of animals and the control of temperature, humidity and odours. Ventilation systems must distribute air uniformly and achieve adequate air exchange, both within cages and within a room.

Noxious odours, particularly ammonia must be kept to a level compatible with the health and comfort of the animals and personnel.

The adequacy of the ventilation system, the design, construction and placement of cages and containers, population densities both within cages and within a room, the effectiveness of the cleaning and the frequency of bedding changes will all influence the level of noxious gases.

Attention should be given to the balance between the need for cleanliness and the potential impact of cleaning procedures on the animals.

Environmental factors potentially affect the welfare of the animals and may affect the results of experiments. Investigators should be consulted prior to any planned changes to the environmental conditions of their animals.
## 6.14. Pens, Cages and Containers

The enclosure shall be designed and managed to meet species-specific needs, unless otherwise signed off as part of an AEC approved project.

Consider the following, for each species held:

- **Sufficient space allocation:**
  - Population density within enclosures, and the placement of these in rooms should ensure that acceptable social and environmental conditions for the species are maintained.
  - House animals only when appropriate for the species or if required by the experiment, e.g. during recovery from surgery or collection of samples.

- **Appropriate substrate:**
  - Wire floor cages for rodents should only be used when essential to the research protocol, and then only for brief periods.
  - Cages with wire floors should include a solid resting area.
  - Bedding (litter) should be appropriate to the species, comfortable, absorbent, dust-free, non-palatable, non-toxic, able to be sterilised if required by the research project.

- **Nesting/ sleeping areas:**
  - All animals should be routinely provided with bedding material.
  - Pregnant animals must be provided with nesting materials where appropriate.

- **Social interaction**
- **Opportunity to perform a species-specific behavioural repertoire**
- **Hiding/retiring zones**
- **Appropriate environmental conditions**
- **Ready access to food and water**

- **Cleaning:**
  - Cleaning should be frequent enough to ensure a healthy environment, without creating unnecessary disturbance.
  - Choice of detergents, disinfectants and pesticides should be made in consultation with investigators.
  - Deodorants designed to mask odours must not be used in animal facilities. They are not an acceptable substitute for good cleaning practices and may interfere with breeding cycles and metabolic processes.
  - Cleaning practices should be monitored regularly to ensure effective sanitation. Methods include visual inspection, monitoring water temperatures and microbiological testing of surfaces after cleaning.
  - Spot cleaning should be used where appropriate.

- **Measures to prevent the spread of pests and disease**
- **Requirements of the project**
- **Ability to observe the animals readily.**

### 6.14.1 Pens, Cages, Enclosure Requirements

Animal containers and enclosures must be:

- constructed of durable, impervious materials;
- kept clean;
- well maintained;
• secure and escape-proof;
• sized to allow animals to stretch out when recumbent and to stand upright and must:
  – protect the animals from climatic extremes;
  – prevent injury to the animals.

6.14.2 Farm Animals (Special Considerations)
As well the above criteria, the following points must be noted when housing farm animals:
• housing and management practices shall meet the requirements of the code of welfare for the species concerned;
• material used for indoor facilities should be impervious to moisture, insects and vermin. Concrete and metal are the preferred building materials;
• pipes supplying drinking water should not be copper or galvanised;
• any paint/sealer applied to wood where cleaning and disinfection is required must be non-toxic to livestock;
• floors (paved or concrete) surfaces should be ‘textured’ to prevent slipping;
• fencing must be maintained to prevent escape or injury;
• ruminants require a resting area either in a well-drained outside area or bedded shelter;
• plan how to manage ventilation, temperature, relative humidity, air velocity, moisture, dust, light, gas accumulation, odours, space and manure. These become increasingly important when large animals are housed indoors.

6.15. Enrichment and Environmental Complexity
Most animals used in RTT are housed in unnatural environments. Wherever possible such animals must be provided with an environment that can accommodate the behavioural and physiological needs of the species.

Almost all the species of animals used in RTT have well-defined social structures and prefer to live in groups, although care must be taken to ensure that the animals are socially compatible. Individual housing is stressful for social animals, therefore social isolation should be avoided whenever possible and limited to meet the specific research objectives as approved by an AEC. The effects of physical isolation should be minimised where possible by the use of non-contact communication, whether visual, auditory or olfactory. Judicious use of mirrors can also be helpful, as can an environment of increased complexity.

The living areas of the animals must be set up and provisioned with the means that will enable them to perform a behavioural repertoire appropriate to the species.

6.16. Admission of New Animals into Holding Areas
Consider the following general points for new animals:
• quarantine and inspect by a qualified person;
• evaluate their health, and treat if required;
• determine suitability for the proposed experiments;
• hold for sufficient time to acclimatise to the facility and personnel;
• experimental manipulations are not normally permitted while animals are held in quarantine. This may vary, depending on the nature of the project. Animals may be bred while in quarantine;
• animals that do not adapt to their new environment should not be kept.
6.17. Routine Husbandry Procedures

Routine husbandry procedures should meet any code of welfare for the species involved and be performed by competent personnel.

Procedures for the care of normal healthy breeding stock and supply of animals are viewed as routine husbandry and fall outside the definition of manipulation.

When special breeding requirements are integral to a research or teaching project such as the creation of a genetically modified animal, then procedures applicable to breeding must be regarded as a manipulation and should be included in the proposal to the AEC.

Variations to normal procedure as part of an experimental project must receive prior AEC approval.

6.17.1 Identification of Animals

The method of identification should be reliable, appropriate for the species and project, and cause the least possible stress to the animal. Available methods include:

- tattoo;
- neck-band;
- individual tag;
- electronic numbering device;
- physical mark;
- box/enclosure label.

Less invasive methods should be chosen when suitable. Invasive identification procedures should be performed, or closely supervised, by an experienced practitioner.

6.18. Euthanasia

The method and procedures used for killing an animal must be humane and:

i. avoid pain or distress and produce rapid loss of consciousness until death occurs.

ii. be compatible with the purpose and aims of the project or activity.

iii. be appropriate to the species, age, developmental stage and health of the animal.

iv. require minimum restraint of the animal.

v. be reliable, reproducible and irreversible.

vi. ensure that animals are killed in a quiet, clean environment away from other animals.

vii. ensure that death is established before disposal of the carcass, foetuses, embryos and fertilised eggs.

Dependent offspring of animals to be killed must be cared for or humanely killed.

NAEAC recommends AECs refer to the American Veterinary Medical Association (AVMA) Guidelines for the Euthanasia of Animals.
6.18.1 Disposal of Animal Carcasses and Waste

Provide for prompt and sanitary disposal of animal carcasses and waste material in accordance with Hazardous Substances and New Organisms (HSNO) and EPA legislation, local council by-laws and community standards.


Every animal facility should have a documented emergency recovery plan, to cover such emergencies as flooding and fire, or the breakdown of lighting, heating, cooling or ventilation. The AEC should:

- work with the organisation to ensure adequate contingency arrangements for the care of animals are in place.
- ensure adequate consideration is given to special classes of animals. For example, does containment of genetically modified (GM) animals requires special attention?
- ensure provision is made for emergency euthanasia of animals.
- require facility staff to be trained against the emergency plan.
People who use animals for scientific purposes have an obligation to treat the animals humanely and to consider their welfare and the Three Rs as essential factors when planning and conducting experiments.

Investigators have direct and ultimate responsibility for all matters related to the welfare of the animals under their control, including the general husbandry and housing of those animals as well as the specific manipulations. They should act in accordance with their specific AEC approval.

The responsibility of investigators extends over all facets of the care and use of animals in projects approved by the AEC, beginning when the animal is allocated to the approved project and ending with its fate at the end of the project.

Investigators are responsible for the standard of animal care and use by all other persons involved in the project. They should ensure that the extent of supervision is compatible with the level of competence of each person and the responsibilities they are given.

In many institutions the daily animal husbandry is delegated to professional animal care personnel. In this case, the investigator must implement strategies for effective communication with the facility manager.

Investigators have a legal and ethical responsibility to ensure that animals on a project are manipulated using medical and surgical techniques which are consistent with the principles of good practice and scientific knowledge in the discipline of laboratory animal veterinary medicine. Investigators should consult with veterinarians whenever unexpected adverse effects occur in order that appropriate veterinary care and treatment regimens are immediately implemented. This responsibility is consistent with the public’s duty of care to seek veterinary management of any sick animals in their charge.

Where an application is likely to involve or impinge on indigenous species, the applicant should discuss these aspects with local iwi or hapu. The application should include evidence that these discussions have taken place and that Māori perspectives are not compromised. If there is any uncertainty, the applicant should approach local Māori representatives for clarification.

7.1. Responsibilities of Teachers

Animals should only be used for teaching activities when there are no suitable alternatives for achieving all of the educational objectives. Students should be given the opportunity to discuss the ethical, legal, social and scientific issues involved in the use of animals for scientific purposes, including teaching.

7.1.1 All New Zealand Schools

Use of animals in all New Zealand schools must comply with the approved CEC of the New Zealand Association of Science Educators. This CEC covers early childhood centres,
kindergartens, schools (both teachers and students) home-schooled students and their families. It is administered by the association on behalf of the Ministry of Education.

An animal ethics committee is established under the code. It approves appropriate projects and is able to provide advice on when AEC approval is necessary.

Any animals that are housed at schools must be well cared for at all times, including during weekends and holidays. The standard of care must meet the requirements of this guide or the relevant code of welfare for the species. Students should not be allowed to take animals home unless there is a clear, written undertaking from a parent or guardian that the animals will be cared for adequately and responsibly.

There are two resources of particular relevance to schools:

1. Ethical guidelines for students in laboratory classes involving the use of animals or animal tissues, an ANZCCART New Zealand publication.

### 7.1.2 Tertiary Institutions

The person in charge of animals used to achieve educational objectives must:

- accept ultimate responsibility for ensuring that the care and use of the animals is in accordance with this guide and all relevant legislation;
- have relevant training and qualifications;
- incorporate into the proposed activities any methods for the replacement, reduction or refinement in the use of animals, provided such methods are compatible with the educational objectives;
- obtain prior AEC approval for use of all animals for the entire course and ensure that activities are conducted as directed and approved by the AEC;
- instruct students appropriately in the care and use of animals before those students participate in experiments with live animals and, where possible, use alternative methods in that preparation;
- ensure that there is close, competent supervision of all students;
- allow students to anaesthetise animals or carry out surgery only if it is essential for their training;
- ensure that in the event of injury to animals, prompt treatment is provided;
- be responsible for the humane killing of the animals, if required.

### 7.2. AEC Approval

Before any project begins, investigators shall receive approval from the AEC. This requires a formal application to the AEC that:

- explains the value of the project;
- explains why non-animal models cannot be used;
- documents the names, roles and qualifications of the key personnel;
- specifies the numbers, types, and stages of animals to be used;
- specifies the procedures that will occur;
• estimates the impact on the animals;
• stipulates what will be done to mitigate the impact;
• describes the fate of the animals which demonstrates that the project will comply with the Act.

Moreover, the investigators must satisfy the AEC of their competence to conduct the techniques described in the experiment.

Species requiring special permission from other organisations (e.g. Department of Conservation) must not be obtained and held before such approval is granted.

If approval is granted, investigators must:
• supervise all aspects of the project to ensure good animal welfare standards are met;
• comply with all conditions imposed by the AEC;
• document the numbers of animals used, and the actual impact grading experienced;
• alert the AEC to any adverse events;
• inform the AEC when the project is completed or discontinued;
• provide a written report to the AEC at the conclusion of the project.

7.3. Planning Projects

It is essential that investigators carefully plan their experimental protocol. This guide raises important issues for consideration which may lead to adjustments to the final protocol for submission to the AEC.

7.3.1 Choice of Species

Ensure the choice of species is appropriate for the purpose of the project. Requirements for known genetic constitution, freedom from specific diseases, documented health, nutritional and environmental histories, and other relevant factors should be taken into account. When the definition of the biological status of animals is necessary, investigators must ensure that the supplier can provide adequate proof that any requirements can be met. Where relevant, species and individual animals should be chosen on the basis that the proposed experiments will result in the least pain and distress. In making this decision, all aspects of the biological nature of the animals, including their behavioural characteristics and their cognitive development, should be taken into account.

7.3.2 Monitoring of Animals on Approval

Observe intensively managed animals as frequently as circumstances require, but at least daily, to assess their health and welfare. Ensure satisfactory arrangements are made for contacting key personnel in the event of emergencies. Any unexpected adverse effects that impact on animal welfare must be reported to the AEC.

7.3.3 Record-keeping

Investigators must record details of animal husbandry routine, environmental conditions, and other potential non-experimental variables that may affect the study. Monitoring records, including methods of assessment of health and welfare shall also be maintained.

Animal use records must meet the statistical reporting requirements of the Animal Welfare (Records and Statistics) Regulations 1999, as detailed in MPI's publication “Animal Use Statistics”. 
7.3.4 Consultation

Investigators should consult other experienced scientists, veterinarians, or laboratory animal, livestock or wildlife specialists when necessary.

7.3.5 Checklist

When planning is completed, the investigator should recheck the protocol to ensure that the following points have been adequately covered and the Three Rs (replacement, refinement and reduction) have been considered.

a. Do the potential benefits outweigh any ethical concerns about the impact on animal welfare?

b. Can the aims be achieved without using animals (replacement)?

c. Are there better ways of achieving the same ends (refinement)?

d. Are suitable holding facilities, equipment and competent personnel available (refinement)?

e. Have all involved personnel been informed of the planned experimental and other procedures?

f. Has the most appropriate species of animal been selected?

g. Is the biological status (genetic, nutritional, microbiological, general health) of the animals appropriate?

h. Are the environmental conditions (including caging or pen type, noise, photoperiod, temperature, humidity, ventilation, density of housing and social structures) appropriate (refinement)?

i. Are the experiments designed so that statistically valid results can be obtained or the educational objectives achieved using the minimum necessary number of animals (reduction)?

j. If the potential impact on the animal is unknown, is it appropriate to incorporate a pilot study into the project design to allow a staged assessment of the impact on animal welfare and how it will be managed (refinement)?

k. If the scientific activity could cause the animals any pain or distress, what will be done to minimise or avoid this (refinement)?

l. What arrangements will be made to monitor the animals adequately, in terms of both their general health and welfare and their response to manipulation (refinement)?

m. If any of the experiments have been performed previously, why should they be repeated (reduction)? Is this required for quality control or legislative reasons (repetition, replication)? Have you checked to see if similar work has been done before so that you can avoid duplicating previous work and thus using animals unnecessarily?

n. If any animals are to be used repeatedly, what will be done to minimise the cumulative effects of such use (refinement)?
7.4. Conduct of Experiments

7.4.1 Limiting Pain and Distress

Pain and distress cannot always be adequately evaluated in animals and investigators must therefore assume that animals experience pain in a manner similar to humans. Decisions regarding their welfare in experiments must be based on this assumption unless there is evidence to the contrary.

The investigator should anticipate any potentially adverse effects of a manipulation and take all possible steps to avoid or minimise pain and distress.

These steps should include:

- choosing the most appropriate and humane method for manipulations;
- ensuring the technical skills and competence of all persons involved in animal care and use;
- using pre-emptive analgesia when pain is anticipated;
- ensuring that animals are adequately monitored to allow prompt alleviation of pain or distress;
- developing a plan to manage any adverse effects of a manipulation e.g. increase in frequency of observation, consultation with a veterinarian, administration of appropriate medication, removal from the project or humane euthanasia;
- using anaesthetic, analgesic and tranquillising agents appropriate to the species and the experimental purposes;
- developing study endpoints that minimise pain and distress;
- conducting projects over the shortest time practicable; and
- using appropriate methods of euthanasia.

The use of local or general anaesthetics, analgesics or tranquillisers must be appropriate to the species, and should meet the criteria generally accepted in current medical, laboratory animal or veterinary practice.

Manipulations which are liable to cause pain of a kind and degree for which anaesthesia would normally be used in medical or veterinary practice must be carried out under anaesthesia.

Distress can sometimes be avoided or minimised by non-pharmacological means. Before an experiment begins, animals should be appropriately conditioned to the experimental environment and procedures and be familiar with animal care personnel. During and after experiments appropriate nursing procedures to minimise pain and distress and to promote the well-being of the animals should be provided.

If animals develop signs of severe pain or distress despite the precautions outlined above, they should have the pain or distress alleviated promptly or must be killed humanely and without delay. Veterinary consultants involved in the animal care programme should be informed immediately. Alleviation of such pain or distress takes precedence over continuing or finishing the experiment. If in doubt, investigators must always seek a professional veterinary opinion before continuing an experiment.

o. Are there any permits that must be obtained for the importation, capture, use, destruction or release of the animals?

p. What arrangements have been made for the fate of all healthy animals at the completion of the project?
Unexpected deaths occurring during a project must be properly investigated by a veterinarian or other qualified person to determine the cause and initiate remedial action. If the deaths are due to manipulations, these must cease. The AEC must be notified and the project protocol resubmitted with appropriate modification.

7.4.2 Animal Welfare Monitoring of Pain or Distress

Investigators should be familiar with the normal behaviour of the animal species chosen, be knowledgeable of signs of pain or distress specific to that species, and must regularly monitor their animals for these signs.

Animals should be monitored to allow detection of deviations from normal behaviour patterns. Such deviations are often the first indications that animals are experiencing pain or distress. Any changes in patterns of sleeping, feeding, drinking, grooming, exploratory behaviour, performance in learning or discriminatory tasks, reproduction or social behaviour should be noted, assessed and acted on if appropriate. Animals must be monitored appropriately for clinical signs of acute pain or distress. These may include one or more of the following:

- abnormal behaviour;
- abnormal stance or movements;
- abnormal sounds;
- altered cardiovascular and/or respiratory function;
- abnormal appetite;
- rapid decline in body weight;
- altered body temperature; and
- vomiting and abnormal defecation or urination.

Animal welfare monitoring score sheets should be used to document the observations and collection of data listed above. For example, variations in body weight, water intake, and grooming behaviour can be recorded every 24 hours or more often as needed during the immediate post-operative period for surgical manipulations.

7.4.3 Study Endpoints

For all but the most minor of manipulations, the investigator should develop humane study endpoints when preparing a project application. These can be used to judge when an animal should be removed from a project or euthanased in order to promote animal welfare.

Death as an endpoint is generally unacceptable and should be fully justified. All animals found in a moribund state must be euthanased unless specifically justified as above and approved by the AEC. It is generally accepted that moribund animals give unreliable research data because they are frequently in a state of multiple organ failure. Attention to good practice indicates that endpoints earlier than the moribund condition should always be used. Typically these are based on changes in body weight, tumour size, and/or body temperature combined with abnormal clinical condition.

For example, animals should be killed when:

- they have lost more than 20% of their pre-study body weight; or
- they have lost more than 10% in 24 hours; or
- a tumour grows to more than 10% of the animal’s weight; or
- body temperature falls below a preset level (as determined by pilot studies which indicate that the level set is predictive of death); or
• animals self-mutilate limbs and feet; or
• animals develop abscesses.

7.4.4 Repeated Use of Animals in Experiments (Reuse)

Individual animals should not be used in more than one experiment, either in the same or different projects, without the express approval of the AEC. However, appropriate reuse of animals may reduce the total number of animals used in a project, result in better design of experiments, and reduce stress or prevent pain in other animals.

The benefits of reusing animals must be balanced against any adverse effects on their well-being, taking into account the lifetime experience of the individual animal.

Applications involving the reuse of animals should take into account:
• the pain or distress and any potential long-term or cumulative effects caused by any previous manipulations;
• the total time taken;
• the pain or distress likely to be caused by the next and subsequent manipulations; and
• whether the animal has recovered fully from the first experiment before further experiments are carried out.

7.4.5 Duration of Experiments

Experimental duration should be the minimum required to provide answers to the questions asked.

Experiments which involve any pain or distress, should be as brief as practicable. AEC approval must be sought for the continued long-term use of individual animals. The decision to continue must be based on the clinical well-being of the animal and the absence of aversion to the experimental situation.

7.4.6 Handling and Restraining Animals

Animals should be handled only by persons instructed and competent in methods which minimise pain, injury or distress.

When the use of restraint devices is necessary for the welfare of the animal and the safety of the handler, they should be used to the minimum extent, for the minimum period required to accomplish the purpose of the experiment, and be appropriate for the animal.

Tranquillisers or anaesthetics may aid restraint but may prolong recovery from the procedure. When these agents have been used, greater attention may be required in assessing the recovery of the animals.

Periods of prolonged restraint or confinement should be avoided. However, where prolonged restraint or confinement is proposed, such as housing livestock in metabolism cages, consideration should be given to their biological needs, including their behavioural requirements and the need for appropriate exercise. Such animals must be assessed regularly by a veterinarian or other qualified person not otherwise involved in the project. If any negative impact on an animal is detected, the animal should be removed from the restraint or the method modified to minimise the impact.
7.4.7 Fate at Completion of Projects

Upon completion of the project, animals must be returned promptly to either normal husbandry conditions or, if appropriate and permitted, to their natural habitat, or, when necessary, be euthanased.

7.4.8 Rehoming

Opportunities to rehome animals should be considered wherever possible, especially when the project or manipulation has had minimal impact on the welfare of the animal. Rehoming should be the preferred choice if the physiological condition and behavioural attributes of the animal indicate that it can be introduced to a new environment with little, or no, transient impact on its well-being.

An animal should not be released to a person at the conclusion of its use unless:

• the AEC has approved such release;
• safeguards are in place and approved by the AEC to ensure the ongoing well-being of the animal;
• transport of animals between sites is appropriate.

7.4.9 Tissue Sharing

Investigators and animal carers should ensure that, if practicable, tissue samples from animals that have died or been humanely killed are provided or made available to other investigators for their work or deposited in a tissue bank for subsequent distribution.

7.4.10 Autopsy

An autopsy should be performed when animals die unexpectedly or are euthanased due to unforeseen complications. The autopsy should be carried out by a person with appropriate qualifications.

Investigators should consider the value of an autopsy for all animals that die for reasons other than approved euthanasia during a project. Post-mortem evaluation may identify one or more non-experimental variables which could compromise the remaining, or future, research subjects.

7.4.11 Pre-operative Planning

Surgical success can be improved by careful attention to the following.

• The use of animals that are fit for purpose will ensure more reliable research data. Investigators should consult the institutional veterinarian or other qualified person to assist in obtaining such animals.
• Pre-operative physical examination can often identify potential problems, such as increased anaesthetic risk, which may compromise the surgical procedure. Animals that are not in an appropriate state of health should be rejected.
• Pre-surgical fasting is necessary for many species to minimise complications of anaesthetic administration.
• Pre-operative antibiotic administration should be considered. This can ensure maximal blood levels of drug during the surgical procedure. Additional post-operative antibiotic treatment may be required.
• Pre-experimental practice on cadavers enables investigators to familiarise themselves with anatomical landmarks and streamline the experimental surgical procedures, thereby reducing the quantity of anaesthetic required, reducing operative time and minimising tissue damage. This will speed post-operative recovery and promote animal well-being.

• For any surgical procedure, a pain management plan aimed at prevention or alleviation of pain and which is appropriate for the procedures and the species should be developed, implemented and reviewed if necessary.

• For any surgical procedure, a pain management plan appropriate for the procedures and the species should be developed, implemented and reviewed if necessary.

7.4.12 Surgery

Anaesthesia and surgery should be performed by competent personnel with appropriate training and experience. Training should only be given by competent personnel with relevant expertise in surgery and anaesthesia.

Surgical procedures should be carried out under appropriate local or general anaesthesia. There should be adequate monitoring of the depth of anaesthesia and effects such as hypothermia, and cardiovascular and respiratory depression.

The choice and administration of anaesthetic, analgesic and tranquillising agents should be suitable for the species and appropriate for the purpose of the experiment. The use of such agents should parallel that used in current medical, laboratory animal or veterinary practice.

Investigators should consider the value of a limited anaesthetic trial to familiarise themselves with new anaesthetic or analgesic drug combinations. Species and strain variation in drug metabolism can result in unexpected morbidity and mortality when dosages are extrapolated from published data. A limited trial, when combined with a non-survival surgical practice session, can provide invaluable information and promote surgical success and animal well-being in subsequent study animals.

When more than one surgical procedure is to be performed the animal must have recovered to good general health between each procedure. Every effort should be made to reduce the total number of procedures on one animal and the AEC should be informed specifically of the need for more than one procedure.

When the animal is not to recover from the surgery, it must be unconscious for the whole procedure, with euthanasia either by overdose of the general anaesthetic or by inducing brain death by a variety of methods e.g. exsanguination or disruption of the thorax.

When the animal is to recover from the anaesthetic, surgical procedures must conform to accepted standards in laboratory animal and veterinary practice. All tissues should be handled with care and particular attention should be given to haemostasis. Aseptic technique should be used for animals which undergo any survival surgery which involves disruption of the skin’s barrier function. Aseptic technique includes aseptic preparation of the surgical field, use of sterilised instruments, wearing of sterile surgical gloves, gowns, caps, and face masks.

The use of post-operative antibiotics should not be a substitute for correct aseptic technique.

7.4.13 Post-operative Care

The comfort of animals must be promoted throughout the post-operative period. Attention should
be given to warmth, hygiene, fluid and food intake, and control of infection. Consideration of pain relief is paramount in post-operative care. The use of analgesics and tranquillisers may be needed to minimise post-operative pain or distress. Care should be taken that animals recovering from anaesthesia are housed to prevent injury and that conditions are such that they are not disturbed, attacked or killed by other animals in the same enclosure.

Investigators should ensure that animals are adequately monitored. They should ensure that they, or other experienced personnel, are fully informed of the animals’ condition. The duties of all personnel must be clearly defined and ways of dealing with emergencies established.

Appropriate clinical records, including observations and administration of any drugs, fluids or other treatments, should be kept, and made accessible to all involved in the post-operative care of the animal.

Regular observation of surgical wounds is essential to check the progress of healing. Any problems must be attended to immediately.

Any post-operative animal observed to be in a state of severe pain or distress that cannot be alleviated quickly must be killed humanely immediately and the consultant veterinarian informed without delay.

7.4.14 Modifying Animal Behaviour

Procedures used to modify an animal’s behaviour or to induce it to perform specific tasks depend on motivation of the animal. The preferred inducement is positive reinforcement, but in some cases the inducement may need to be some form of biological stress. This stress should be as mild as possible. Severe deprivation of water, food, social interaction or sensory stimuli should not be used.

The level and duration of painful or noxious stimuli should be minimised and escape from the stimuli should be available. Behaviour can usually be modified using procedures that involve no more of a stressor than that normally experienced by the species. When noxious stimuli are used to modify behaviour the AEC should be aware of the duration and possibility of escape from the stimuli.

7.4.15 Withholding Food or Water

Projects involving the withholding or severe restriction of food or water should produce no continuing detrimental effect on the animals. In these experiments, the fluid balance and/or body weight must be monitored, recorded and maintained within the limits approved by the AEC.

7.5. Guidelines for Specific Manipulations

7.5.1 Production of Genetically Modified Animals

The breeding or production of an animal using genetic modification that may result in the birth or production of an animal that is more susceptible to, or at greater risk of, pain or distress during its life as a result of the breeding or production is considered a manipulation under the Act.

Genetic modification is defined as the deletion, change, or moving of genes within an organism, or the transfer of genes from one organism to another, or the modification of existing genes, or
the construction of new genes and their incorporation into any organism (as defined in the Royal Commission Report 2001).

Reproductive or therapeutic cloning is the genetic modification of animals, in particular livestock or rodents, to establish desired traits for food production (e.g. meat quality), livestock health, or medicine production (e.g. induce cows to produce medical compounds in milk). Alternatively, animals can be produced with genetic mutations to better understand disease process. Cloning uses a process called somatic cell nuclear transfer (SCNT). Scientists take an immature egg or oocytes from a female animal, often by inducing superovulation. Instead of combining it with sperm, they remove the nucleus (which contains the oocyte’s genes). This leaves behind the other components necessary for the initial stages of embryo development. Scientists then add the nucleus or cell from the donor animal that has the desirable or modified traits they wish to copy. The donor nucleus is fused with the ooplast (the oocyte whose nucleus has been removed), and an embryo begins to form. The embryo is then implanted in the uterus of a surrogate dam which carries it to term.

7.5.2 Animal Models of Disease

The scientific validity of animal models of human diseases rests in part on how closely a given model resembles a particular disease. Thus, the attendant pain and distress of the human disease may also occur in the animal. Special care must be taken in selecting the appropriate species, and the investigator must accept responsibility for ensuring that any pain or distress is minimised and that the AEC is informed of the potential effects of the disease on the animals.

Investigators must not allow the experiments to proceed to the painful or distressful or lingering death of animals unless no other experimental endpoint is feasible and the goals of the experiments are the prevention, alleviation, treatment or care of a life-threatening disease or situation in human beings or animals.

Investigators must avoid using death as an experimental endpoint whenever possible. When death as an endpoint cannot be avoided, the experiments must be designed to result in the deaths of as few animals as possible.

7.5.3 Experimental Induction of Neoplasia

The site for induction of tumours must be chosen carefully. Subcutaneous sites on the back or flank should be chosen when possible. Prior to the use of footpad, brain and eye sites, specific justification as to the lack of any alternative should be made to the AEC.

Investigators should monitor their animals regularly for signs of pain or distress, especially sudden changes in body condition and signs that tumour growth is impacting on the well-being of the animals.

Animals with experimentally induced tumours should be euthanased before predictable death occurs, cachexia becomes advanced or the tumour becomes large enough to cause ulceration or severe limiting of normal behaviour. Animals should be used when tumours are at the minimum size necessary to obtain valid results.

In tumour therapy experiments, the endpoints chosen should be as early as possible, compatible with reliable assessment of the therapy. Weight changes should be monitored closely. Death from the tumour should not be an endpoint.
7.5.4 Production of Monoclonal Antibodies

*In vitro* methods should be used for the routine amplification of hybridomas for the production of monoclonal antibodies. Investigators wishing to use the *in vivo* (ascites) method should provide in their proposal to the AEC recent laboratory evidence to show that *in vitro* methods are unsuitable for the specific monoclonal antibody that is the subject of the proposal.

In the immunisation phase, investigators must ensure the minimisation of pain and distress to animals from factors such as the type, volume, site and frequency of the injection of adjuvants, and the methods and frequency of blood sampling.

With ascitic tumours, including hybridomas, investigators should ensure that the volume of ascitic fluid does not cause gross abdominal distension, and the volumes of solid tumours and cachexia do not become distressful to the animals. Careful monitoring is necessary because body weight loss can be difficult to discern in the presence of ascites and abdominal tumours.

7.5.5 Lesions of the Central Nervous System

Anatomical or chemical lesions of the central nervous system have been widely used to study its structure and function in health and disease. These experiments demand special consideration when the lesion produces loss of function, including impairment of limb or trunk movements, loss of sensibility to touch, temperature or pain, impairment of the animal's awareness of its surroundings or impairment of appetite or injury mechanisms. Special animal care, caging, and other facilities may be needed and the AEC, in approving such experiments, has a particular responsibility to ensure that these facilities are available and that the condition of the animals is closely monitored.

7.5.6 Implanted Devices

Investigators should be aware of the need for strict attention to aseptic technique when foreign bodies are surgically implanted. Contamination of prosthetic devices frequently requires their removal after antibiotic therapy has failed.

Skilled and specialised attention is required in the care of animals following an operation in which monitoring or sampling devices have been implanted, or a fistula created. Regular observation is essential to determine signs of distress, pain or infection which must be treated immediately.

7.5.7 Organ and Tissue Transplantation

Skilled and specialised attention is required for animals following organ or tissue transplantation. Animals must be assessed frequently for any signs of pain, distress, infection and tissue rejection and treated immediately if these occur. Special attention should be given to the management of immunosuppression and the disease hazards and adverse outcomes that may be associated with organ and tissue transplantation between species (xenotransplantation). Death as an endpoint is unacceptable when determining recipient survival times.

7.5.8 Neuromuscular Paralysis

Neuromuscular blocking agents may only be used with adequate general anaesthesia. When these agents are used, specialist advice on anaesthesia should be obtained.

Immobilisation of an animal solely with a neuromuscular blocking agent is not acceptable.
When these agents are used with general anaesthesia, special care must be taken to ensure the maintenance of an adequate plane of anaesthesia. Since the usual criteria such as character of respiration and corneal and flexor withdrawal reflexes cannot be used, continuous or frequent intermittent monitoring of physiological variables such as heart rate, blood pressure, oxygen saturation, pupil size and electroencephalogram is necessary, together with the effects of these of mild sensory stimuli. Care is required to ensure that drugs used in the procedures do not interfere with this monitoring.

7.5.9 Electro-immobilisation

Electro-immobilisation must not be used as an alternative to analgesia or anaesthesia. When its use is proposed for the restraint of animals, AECs must carefully evaluate published evidence to assess whether it may cause distress. If so, an alternative restraint method must be used.

7.5.10 Toxicological Experiments

If suitable non-animal tests are available, they shall be used. In particular, \textit{in vitro} methods must be used as an initial screening test wherever possible.

Investigation of the safety of agents intended for use in human beings, animals, the household or the environment, or of naturally occurring toxins, or of agents to be used as poisons for pest control should be performed by persons with appropriate training.

The endpoint of such experiments must be as early as is compatible with reliable assessment of toxicity, and must minimise the extent of any pain and distress.

When death as an endpoint cannot be avoided, the experiments must be designed to result in the deaths of as few animals as possible.

All such tests shall be undertaken in accordance with internationally-recognised guidelines such as an OECD Guideline for the testing of chemicals.

Investigators must not allow the painful, distressing or lingering deaths of animals unless no other endpoint is feasible and the goals of the project are the prevention, alleviation or cure of a life-threatening disease or situation in humans or animals.

7.5.11 Foetal Experimentation

Under the Act, the definition of an animal includes mammalian, avian and reptilian prenatal or pre-hatched young in the second half of their developmental stage. Any projects involving such animals require AEC approval.

When foetal experimentation or surgery compromises the ability of the neonate to survive and be without pain or distress, it must be euthanased before or immediately following birth unless such pain or distress can be relieved.

Although there is increasing evidence that foetuses do not feel pain under normal conditions, investigators should ensure adequate anaesthesia for both mother and foetus when the latter is undergoing surgical or other manipulation \textit{in utero}.

During surgery of the mother consideration must be given to any special requirements for anaesthesia of the foetus.
Eggs must be destroyed before hatching, unless hatching is a requirement of the experiment. The AEC must approve the arrangements made for hatchlings.

7.5.12 Research on Pain Mechanisms and the Relief of Pain

In experiments in which unanaesthetised animals are to be subjected to stimuli designed to produce pain or when pain is to be inflicted on animals as part of normal management, investigators should:

- ensure that these stimuli limit pain at all times to levels comparable to those which do not distress human beings
- ensure that the animals are exposed to the minimum pain necessary for the purpose of the experiment
- provide treatment for the relief of pain, or allow self-administration of analgesics, or escape from repetitive, painful stimuli when possible.

7.5.13 Animal Welfare and Animal Health Research

When studying ways of improving the health and welfare of animals, investigators may need to design experiments that replicate a problem such as injury, trauma, nutritional disorder, physical exertion, disease or environmental stress.

Thus, the attendant pain and distress may also be replicated. When such experiments are necessary, the investigator must ensure that:

- the principal aim of the project is to improve animal health or welfare;
- alternative methods, such as the use of animals already experiencing the problem, are not possible;
- all possible steps are taken to minimise any pain or distress;
- the experiments do not proceed to the painful or distressful or lingering death of animals unless no other experimental endpoint is feasible and the goals of the experiments are the prevention, alleviation, treatment or care of a life-threatening disease or situation. When death as an endpoint cannot be avoided, the experiments must be designed to result in the deaths of as few animals as possible.

7.5.14 Experiments Involving Hazards to Humans or Other Animals

Hazards may arise from sources that include:

- viruses;
- bacteria;
- fungi;
- parasites;
- radiation;
- radioactivity;
- corrosive substances;
- toxins;
- allergens;
- carcinogens;
- recombinant DNA;
- anaesthetic gases; and
- physical injuries.
Any potential pathogenic effects of these hazards when used in experiments must be explained as far as possible to all personnel. Tests before, during and after the experiments may be required for personnel.

Protocols submitted to the institution’s AEC should include a description of any intended use of hazardous compounds or organisms. They should describe specific safety measures and disposal protocols used to prevent contamination of caging, other animals, research personnel and students.

Investigators must not allow the experiments to proceed to the painful or distressful or lingering death of animals unless no other experimental endpoint is feasible and the goals of the experiments are the prevention, alleviation, treatment or care of a life-threatening disease or situation in human beings or animals. When death as an endpoint cannot be avoided, the experiments must be designed to result in the deaths of as few animals as possible.

Precautions, security and emergency plans to contain hazardous agents must be appropriate to a “worst-case” situation.
APPENDIX 1 – CEC CHECKLIST

COVER PAGE
Name of institution/code holder

Act Reference
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TABLE OF CONTENTS

TABLE OF ABBREVIATIONS

1. INTRODUCTION
[Background on the activities of the applicant]

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2. ESTABLISHMENT OF ANIMAL ETHICS COMMITTEE
All members of the AEC should be required to read and understand the CEC.

Statutory membership
Code holder or senior member of the organisation
A veterinarian nominated by the New Zealand Veterinary Association
A person nominated by an approved animal welfare organisation
A person nominated by a territorial authority or regional council
Reappointment of external members at the expiry of their term must be done through a formal nomination by the relevant body
External member appointments
Remuneration for AEC members
The term of appointment
Induction of new members
Appointment and endorsement of Chairperson

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3. AEC PROCEDURES
Meetings
Public access to meetings
Timing for circulation of agenda items
Frequency of meetings
Quorum
Vacancies
– Impact on quorum
– Prolonged absence
Decision process
Conflict of interest
Ensuring effective input of all members
Use of teleconference/videoconference
Consideration between meetings

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S101 (3) S101 (4)
S100 (5)
S101 (6) S101 (7)
S101 (8) S101 (9)
S101 (10)
• Confidentiality
• Applicant present or not
• Secretarial duties

4. CONSIDERATION OF PROTOCOLS BY THE AEC

The function of the AEC is to ensure that the highest ethical standards are observed by ... and all persons associated with it in relation to the manipulation and use of animals

• Criteria for consideration
• Impact grading
• SOPs
• Avoiding duplication
• Outcomes of consideration
• Conditions of approval
• Maximum approval period
• Power to suspend/revoke/vary project approvals
• Protection of members
• Compliance with other legislation

5. RESPONSIBILITIES

• Compliance reporting
• Project monitoring
• Interim reporting
• Changes to approved manipulations
• Adverse event reporting
• End of approval reporting
  – End of approval grading
  – End of approval statistics
• Non-compliance
• Record keeping and storage of information
  – Security/access to information
  – Duration of storage
  – Storage of information
  – Reporting of statistics to MPI

6. ANIMAL FACILITIES AND PRACTICES

• Management of facilities
• Staffing facilities
• Development of SOPs
• Sourcing of animals
• Transport of animals
• Housing of animals
• Dealing with sick or injured animals
• Euthanasia for tissue collection/dissection
7. **MONITORING**

The animal ethics committee has the power to inspect animals, their accommodation, and related experimental records at any time to satisfy itself that approved procedures are being properly carried out.

- Frequency of approval monitoring
- Monitoring of manipulations graded A&B
- Monitoring of manipulations graded C-E
- Monitoring on behalf of the AEC by a nominated person

8. **ARRANGEMENTS FOR EXTERNAL PARTIES TO USE THE CODE AND AEC**

9. **COMPLAINTS PROCEDURES**

10. **AMENDMENT, SUSPENSION AND REVOCATION OF A CEC**
# APPENDIX 2 – EXAMPLE OF A TEMPLATE FOR ANIMAL USE APPLICATIONS

## 1. Project description

<table>
<thead>
<tr>
<th>A</th>
<th>Lay summary</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Lay summary of the project</td>
</tr>
<tr>
<td></td>
<td>Lay summary of the proposed manipulations, including animal numbers and manipulation gradings (NAEAC impact scale details)</td>
</tr>
<tr>
<td></td>
<td>Application of the Three Rs (reduction, replacement, refinement)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>B</th>
<th>Background</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Justification for the need for animals including consideration of suitable alternatives and previous relevant studies</td>
</tr>
<tr>
<td></td>
<td>Projected benefits to animals or humans vs cost to the animals</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>C</th>
<th>Study design</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Objectives</td>
</tr>
<tr>
<td></td>
<td>Methods (items in 2-5 could be incorporated in this section) Sample sizes and justification, randomisation, blinding</td>
</tr>
</tbody>
</table>

## 2. Animals, husbandry and welfare

<table>
<thead>
<tr>
<th>A</th>
<th>Species, sex, strain, age and why they are appropriate</th>
</tr>
</thead>
<tbody>
<tr>
<td>B</td>
<td>Source of animals</td>
</tr>
<tr>
<td>C</td>
<td>Transport requirements</td>
</tr>
<tr>
<td>D</td>
<td>Animal housing and locations</td>
</tr>
<tr>
<td>E</td>
<td>Husbandry to ensure general health and welfare</td>
</tr>
<tr>
<td>F</td>
<td>Health status assessment of animals before, during and after manipulation</td>
</tr>
<tr>
<td>G</td>
<td>Specific requirements of animals bred with potential for compromised welfare e.g. transgenic animals</td>
</tr>
<tr>
<td>H</td>
<td>Harm or distress that may be experienced by animals resulting from manipulations, whether this can be alleviated and, if so, the methods to be used</td>
</tr>
<tr>
<td>I</td>
<td>Previous RTT use of the animals, and consideration of the possible cumulative effects</td>
</tr>
</tbody>
</table>
3. Surgical and other manipulations

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Anaesthetics used, dosing rates and frequency*</td>
</tr>
<tr>
<td>B</td>
<td>How depth of anaesthesia will be monitored during the procedure*</td>
</tr>
<tr>
<td>C</td>
<td>Adverse effects and their management – pain management* – degree of harm, multiple procedures/surgeries on one animal, peri-surgical support to maximise welfare outcomes e.g. use of fluid therapy, heat pads</td>
</tr>
<tr>
<td>D</td>
<td>Other manipulations with the potential to impact on welfare e.g. use of muscle relaxants</td>
</tr>
<tr>
<td>E</td>
<td>Environmental manipulation including lack of enrichment, single housing of social species, confinement e.g. in metabolic crates</td>
</tr>
</tbody>
</table>

4. Euthanasia

<p>| | |</p>
<table>
<thead>
<tr>
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</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Methods, drugs, routes*</td>
</tr>
<tr>
<td>B</td>
<td>Personnel involved</td>
</tr>
</tbody>
</table>

*Note: the AEC may not require some or all of this detail in the main document and instead require attachment of supporting documentation of authorised use of veterinary medicines (e.g. if the applicant already holds an approved Institutional Drug Administration Order (IDAO), this could be attached).

5. Endpoints

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
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</thead>
<tbody>
<tr>
<td>A</td>
<td>Study endpoints – at what point will the study end for animals? What will happen to them on completion?</td>
</tr>
<tr>
<td>B</td>
<td>Humane endpoints – where animals will be euthanased within the study to avoid suffering, how will certainty of outcome be balanced against animal welfare considerations?</td>
</tr>
<tr>
<td>C</td>
<td>Death as an endpoint – if death is the required endpoint, this must be fully justified</td>
</tr>
</tbody>
</table>

6. Project personnel details

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Names and contact details – including emergency contact details</td>
</tr>
<tr>
<td>B</td>
<td>Training and experience</td>
</tr>
<tr>
<td>C</td>
<td>How additional training will be provided, if required</td>
</tr>
</tbody>
</table>

7. Animals used in teaching

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Preparation of students for animal use, especially consideration of ethics e.g. cost versus benefit</td>
</tr>
<tr>
<td>B</td>
<td>Extent of student supervision</td>
</tr>
<tr>
<td>C</td>
<td>Relevant experience of students</td>
</tr>
<tr>
<td>D</td>
<td>Copy of laboratory handout</td>
</tr>
</tbody>
</table>
## 8. Other requirements

<table>
<thead>
<tr>
<th>A</th>
<th>Confirmation that appropriate investigations (e.g. literature searches, patent searches, personal networks) have been undertaken to avoid unnecessary duplication.</th>
</tr>
</thead>
<tbody>
<tr>
<td>B</td>
<td>Has the application been considered by another AEC?</td>
</tr>
<tr>
<td>C</td>
<td>How will the findings be promoted or published?</td>
</tr>
<tr>
<td>D</td>
<td>Signatures of approval (e.g. science reviewer, biometrician, animal facility manager, head of department or equivalent)</td>
</tr>
<tr>
<td>E</td>
<td>Project leader to be responsible for ensuring that key project personnel read and understand the approval</td>
</tr>
<tr>
<td>F</td>
<td>Need for formal amendment to the approval if changes are required</td>
</tr>
<tr>
<td>G</td>
<td>Reporting to the AEC by the project leader required</td>
</tr>
<tr>
<td>H</td>
<td>Monitoring of the approved project to be conducted by the AEC</td>
</tr>
<tr>
<td>I</td>
<td>Other conditions of approval specific to the particular application</td>
</tr>
</tbody>
</table>