Risk Assessment and Risk Management Plan

for

**DIR 154**

Limited and controlled release of a GM vaccine for chickens, Vaxsafe® ILT

**Applicant** - Bioproperties Pty Ltd

PAGE INTENTIONALLY LEFT BLANK

# Summary of the Risk Assessment and Risk Management Plan

**for**

**Licence Application No. DIR 154**

# Decision

The Gene Technology Regulator (the Regulator) has decided to issue a licence for this application for the intentional release of a genetically modified organism (GMO) into the environment. The licence authorises conduct of experiments, transport and disposal of a GM vaccine to protect chickens against infectious laryngotracheitis for the purpose of field trials.

A Risk Assessment and Risk Management Plan (RARMP) for this application was prepared by the Regulator in accordance with the requirements of the *Gene Technology Act 2000* (the Act) and corresponding State and Territory legislation, and finalised following consultation with a wide range of experts, agencies and authorities, and the public. The RARMP concludes that the field trials pose negligible risks to human health and safety and the environment and that any risks posed by the dealings can be managed by imposing conditions on the release.

Veterinary medicines must be approved by the Australian Pesticides and Veterinary Medicines Authority (APVMA), which provides a national registration scheme for agricultural and veterinary chemical products under the *Agricultural and Veterinary Chemicals Code Act 1994* (AgVet Code). The APVMA has issued a permit to Bioproperties to supply and use the GM vaccine for the purpose of animal research.

# The application

|  |  |
| --- | --- |
| Application number | DIR 154 |
| Applicant | Bioproperties Pty Ltd |
| Project Title | Limited and controlled release of a GM vaccine for Chickens, Vaxsafe® ILT |
| Parent organism | Infectious laryngotracheitis virus (ILTV) CSW-1 strain |
| Modified genes | Deletion of gene encoding glycoprotein G protein from the ILTV genome |
| Proposed release date | August 2017 – August 2022 |
| Proposed duration | 5 years |
| Proposed locations | Selected chicken farms in rural Victoria and New South Wales |
| Purpose | To study the efficacy and safety of a GM vaccine against infectious laryngotracheitis disease in farmed broiler chickens. |

The proposed field trials would assess the efficacy and safety of the GM vaccine under field conditions, including likelihood of challenge with a range of distinct field strains. The field trials are proposed to take place at up to 40 selected broiler farms, potentially including free range farms, in rural Victoria and NSW. Up to 2,000,000 chickens would be inoculated with the GM vaccine over a 5 year period. As is common in veterinary vaccine trials, the vaccinated chickens could enter general commerce, including use in human food or animal feed. At an appropriate time, the chickens inoculated by the GM vaccine would be transported from farms to poultry processing plants.

# Risk assessment

The risk assessment concludes that risks to the health and safety of people, or the environment, from the proposed dealings, either in the short or long term, are negligible. No specific risk treatment measures are required to manage these negligible risks.

The risk assessment process considers how the genetic modifications and proposed activities conducted with the GM vaccine might lead to harm to people or the environment. Risks are characterised in relation to both the seriousness and likelihood of harm, taking into account information in the application (including proposed controls), relevant previous approvals and current scientific/technical knowledge. Both the short and long term impact are considered.

Credible pathways to potential harm that were considered included exposure of people or susceptible birds to the GMO, potential for recombination and establishment of the GMO outside the trial limits. Potential harms that were considered in relation these pathways included disease, toxicity or allergenicity to people and adverse impacts to desirable species in the environment.

The principal reasons for the conclusion of negligible risks are the attenuated phenotype of the GMO, ILTV’s limited host range, APVMA permit conditions for the use of the GM vaccine, local council and state requirements for broiler farms, and suitability of the controls proposed by the applicant.

# Risk management plan

The risk management plan describes measures to protect the health and safety of people and to protect the environment by controlling or mitigating risk. The risk management plan is given effect through licence conditions.

As the level of risk is considered negligible, specific risk treatment is not required. However, since this is a limited and controlled release, the licence includes limits on the size, location and duration of the release, as well as a range of controls to minimise the potential for the GMO to spread in the environment. In addition, there are several general conditions relating to ongoing licence holder suitability, auditing and monitoring, and reporting requirements which include an obligation to report any unintended effects.

Table of contents

[**SUMMARY OF THE RISK ASSESSMENT AND RISK MANAGEMENT PLAN** III](#_Toc489267258)

[DECISION III](#_Toc489267259)

[THE APPLICATION III](#_Toc489267260)

[RISK ASSESSMENT IV](#_Toc489267261)

[RISK MANAGEMENT PLAN IV](#_Toc489267262)

[**TABLE OF CONTENTS** V](#_Toc489267263)

[**ABBREVIATIONS** VII](#_Toc489267264)

[**CHAPTER 1 RISK ASSESSMENT CONTEXT** 1](#_Toc489267265)

[SECTION 1 BACKGROUND 1](#_Toc489267266)

[SECTION 2 REGULATORY FRAMEWORK 1](#_Toc489267267)

[2.1 Interface with other regulatory schemes 2](#_Toc489267268)

[SECTION 3 BACKGROUND TO THE DIR APPLICATION 3](#_Toc489267269)

[SECTION 4 THE PROPOSED FIELD TRIALS 3](#_Toc489267270)

[4.1 The proposed limits of the field trials (duration, scale, location and people) 4](#_Toc489267271)

[4.2 The proposed controls to restrict the spread and persistence of the GMO in the environment 4](#_Toc489267272)

[4.3 Details of the proposed activities 4](#_Toc489267273)

[SECTION 5 PARENT ORGANISM 9](#_Toc489267274)

[5.1 Basic Biology 10](#_Toc489267275)

[5.2 Host range 11](#_Toc489267276)

[5.3 Clinical signs 11](#_Toc489267277)

[5.4 Latency 12](#_Toc489267278)

[5.5 Shedding 13](#_Toc489267279)

[5.6 Transmission 13](#_Toc489267280)

[5.7 ILTV vaccines 14](#_Toc489267281)

[5.8 ILTV classes and recombination between types 14](#_Toc489267282)

[5.9 Recent outbreaks in Australia 16](#_Toc489267283)

[5.10 Environmental stability and decontamination methods 16](#_Toc489267284)

[SECTION 6 THE GMO – NATURE AND EFFECT OF GENETIC MODIFICATIONS 17](#_Toc489267285)

[6.1 The genetic modification 17](#_Toc489267286)

[6.2 Glycoprotein G 18](#_Toc489267287)

[6.3 Characterisation of the GMO 18](#_Toc489267288)

[SECTION 7 RECEIVING ENVIRONMENT 20](#_Toc489267289)

[7.1 Background on broiler farming 20](#_Toc489267290)

[7.2 Biosecurity 26](#_Toc489267291)

[7.3 Waste management 29](#_Toc489267292)

[7.4 Site of release 33](#_Toc489267293)

[7.5 Related viral species in the receiving environment 33](#_Toc489267294)

[7.6 Potential hosts in the environment 33](#_Toc489267295)

[SECTION 8 PREVIOUS AUTHORISATIONS 34](#_Toc489267296)

[8.1 Australian authorisations 34](#_Toc489267297)

[8.2 International authorisations and experience 34](#_Toc489267298)

[**CHAPTER 2 RISK ASSESSMENT** 35](#_Toc489267299)

[SECTION 1 INTRODUCTION 35](#_Toc489267300)

[SECTION 2 RISK IDENTIFICATION 35](#_Toc489267301)

[2.1 Postulated risk scenarios 36](#_Toc489267302)

[SECTION 3 UNCERTAINTY 53](#_Toc489267303)

[SECTION 4 RISK EVALUATION 54](#_Toc489267304)

[**CHAPTER 3 RISK MANAGEMENT PLAN** 55](#_Toc489267305)

[SECTION 1 BACKGROUND 55](#_Toc489267306)

[SECTION 2 RISK TREATMENT MEASURES FOR SUBSTANTIVE RISKS 55](#_Toc489267307)

[SECTION 3 GENERAL RISK MANAGEMENT 55](#_Toc489267308)

[3.1 Licence conditions to limit and control the release 55](#_Toc489267309)

[3.2 Other risk management considerations 59](#_Toc489267310)

[SECTION 4 ISSUES TO BE ADDRESSED FOR FUTURE RELEASES 60](#_Toc489267311)

[SECTION 5 CONCLUSIONS OF THE RARMP 60](#_Toc489267312)

[**REFERENCES**  62](#_Toc489267313)

[**Appendix A** 68](#_Toc489267314)

**TABLE OF FIGURES**

[Figure 1. Summary of parameters used to establish the risk assessment context 1](#_Toc482087663)

[Figure 2. Organisation of ILTV genome 10](#_Toc482087664)

[Figure 3. Construction of the GM virus 17](#_Toc482087665)

[Figure 4. The risk assessment process 35](#_Toc482087666)

[Figure 5. Components of a risk scenario 36](#_Toc482087667)

Abbreviations

|  |  |
| --- | --- |
| ACEC | Animal Care and Ethics Committee |
| ACMF | Australian Chicken Meat Federation |
| AgVet Code | *Agricultural and Veterinary Chemicals Code Act* *1994* |
| APVMA | Australian Pesticides and Veterinary Medicines Authority |
| BHV | *Bovine herpesvirus* |
| BOD | biological oxygen demand |
| bp | base pairs |
| cDNA | complementary DNA |
| CEK | chicken embryo kidney |
| CEO | chicken embryo origin |
| CMA | catchment management authorities |
| DAWR | Department of Agriculture and Water Resources |
| DIR | Dealings involving Intentional Release |
| DNA | deoxyribonucleic acid |
| eGFP | enhanced green fluorescent protein |
| EHV | *Equine herpesvirus* |
| ELISA | enzyme-linked immunosorbent assay |
| EPA | Environment Protection Authority |
| EP&A Act | *Environment Planning and Assessment Act 1979* |
| FeHV | *Feline herpesvirus* |
| FPV | *Fowlpox virus* |
| FSANZ | Food Standards Australia New Zealand |
| gC | glycoprotein C |
| gG | glycoprotein G |
| GM | genetically modified |
| GMO | genetically modified organism |
| GMP | Good Manufacturing Practice |
| GTTAC | Gene Technology Technical Advisory Committee |
| HACCP | Hazard Analysis of Critical Control Points |
| HEPA | High-Efficiency Particulate Air filter |
| HSV | *Herpes simplex virus* |
| HVT | *Herpesvirus of turkeys* |
| IBC | Institutional biosafety committee |
| ILTV | *Infectious laryngotracheitis virus* |
| IR | internal repeat |
| kb | kilo base pairs |
| L | litres |
| LMH | leghorn chicken hepatocellular carcinoma |
| LTS | Land Transport Standards |
| m | metres |
| ml | millilitres |
| mm | millimetres |
| mRNA | messenger ribonucleic acid |
| NLRD | Notifiable Low Risk Dealings |
| NSW | New South Wales |
| NICNAS | National Industrial Chemicals Notification and Assessment Scheme |
| OGTR | Office of the Gene Technology Regulator |
| Ori | origin of replication |
| PCR | polymerase chain reaction |
| PC2 | Physical containment 2 |
| PFU | plaque forming unit |
| POEO Act | *Protection of the Environment Operations Act 1997 (NSW)* |
| PsHV | *Psittacid herpesvirus* |
| RARMP | Risk Assessment and Risk Management Plan |
| Regulations | Gene Technology Regulations 2001 |
| Regulator | Gene Technology Regulator |
| RFLP | Restriction Fragment Length Polymorphism |
| RMIT | Royal Melbourne Institute of Technology |
| qPCR | quantitative polymerase chain reaction |
| TCO | tissue culture origin |
| TGA | Therapeutic Goods Administration |
| the Act | *Gene Technology Act 2000* |
| TR | terminal repeat |
| UL | unique long |
| US | unique short |
| US | United States |
| vCKBP | virus-encoded chemokine binding protein |
| v/v | volume/volume |

1. Risk assessment context
	1. Background
2. An application has been made under the *Gene Technology Act 2000* (the Act) for a licence to conduct Dealings involving the Intentional Release (DIR) of genetically modified organisms (GMOs) into the Australian environment.
3. The Act in conjunction with the Gene Technology Regulations 2001 (the Regulations), an inter-governmental agreement and corresponding legislation in States and Territories, comprise Australia’s national regulatory system for gene technology. Its objective is to protect the health and safety of people, and to protect the environment, by identifying risks posed by or as a result of gene technology, and by managing those risks through regulating certain dealings with GMOs.
4. This chapter describes the parameters within which potential risks to the health and safety of people or the environment posed by the proposed release are assessed. The risk assessment context is established within the regulatory framework and considers application-specific parameters (Figure 1).

PREVIOUS RELEASES

RECEIVING ENVIRONMENT

Environmental conditions

Presence of related species

Presence of similar genes

PARENT ORGANISM

Origin and taxonomy

Biological characterisation

PROPOSED DEALINGS

Proposed activities involving the GMO

Proposed limits of the release

Proposed control measures

GMO

Introduced or deleted genes (genotype)

Novel traits (phenotype)

**RISK ASSESSMENT CONTEXT**

LEGISLATIVE REQUIREMENTS

(including Gene Technology Act and Regulations)

RISK ANALYSIS FRAMEWORK

OGTR OPERATIONAL POLICIES AND GUIDELINES

Figure 1. Summary of parameters used to establish the risk assessment context

* 1. Regulatory framework
1. Sections 50, 50A and 51 of the Act outline the matters which the Gene Technology Regulator (the Regulator) must take into account, and who must be consulted, when preparing the Risk Assessment and Risk Management Plans (RARMPs) that inform the decisions on licence applications. In addition, the Regulations outline further matters the Regulator must consider when preparing a RARMP.
2. In accordance with Section 50A of the Act, this application is considered to be a limited and controlled release application, as its principal purpose is to enable the applicant to conduct experiments and the applicant has proposed limits on the size, location and duration of the release, as well as controls to restrict the spread and persistence of the GMOs and their genetic material in the environment. Therefore, the Regulator was not required to consult with prescribed experts, agencies and authorities before preparation of the RARMP.
3. Section 52 of the Act requires the Regulator to seek comment on the RARMP from the States and Territories, the Gene Technology Technical Advisory Committee, Commonwealth authorities or agencies prescribed in the Regulations, the Minister for the Environment, relevant local council(s), and the public. The advice from the prescribed experts, agencies and authorities and how it was taken into account is summarised in Appendix A. No submissions from the public were received.
4. The *Risk Analysis Framework* (OGTR 2013) explains the Regulator’s approach to the preparation of RARMPs in accordance with the legislative requirements. Additionally, there are a number of operational policies and guidelines developed by the Office of the Gene Technology Regulator (OGTR) that are relevant to DIR licences. These documents are available from the [OGTR website](http://www.ogtr.gov.au/internet/ogtr/publishing.nsf/Content/home-1).
	* 1. Interface with other regulatory schemes
5. Gene technology legislation operates in conjunction with other regulatory schemes that regulate GMOs or genetically modified (GM) products in Australia. Dealings conducted under a licence issued by the Regulator may also be regulated by the Therapeutic Goods Administration (TGA), Food Standards Australia New Zealand (FSANZ), the Australian Pesticides and Veterinary Medicines Authority (APVMA), the National Industrial Chemicals Notification and Assessment Scheme (NICNAS) and the Department of Agriculture and Water Resources (DAWR). Dealings may also be subject to the operation of State legislation declaring areas to be GM, GM-free, or both, for marketing purposes.
6. To avoid duplication of regulatory oversight, risks that have been considered by other regulatory agencies are generally not assessed by the Regulator.
7. The APVMA provides a national registration and permit scheme for agricultural and veterinary chemical products. It administers the provisions of the *Agricultural and Veterinary Chemicals Code Act 1994* (AgVet Code). For registration, the APVMA assesses whether a new veterinary vaccine meets the criteria set out in the AgVet Code before it is registered in the Register of Agricultural and Veterinary Chemical Products. A new veterinary vaccine that is not registered may be legally used, such as in animal trials, by obtaining a permit from the APVMA. As part of the permit process, the APVMA assesses the quality, safety and efficacy of the vaccine. Quality aspects could include batch-to-batch consistency in vaccine composition, purity and potency. The APVMA audits the Good Manufacturing Practice (GMP) record of the applicant. Safety aspects include the toxicological profile of the vaccine and its residues, including metabolites and degradation products. The APVMA approves the label, handling and directions for use of veterinary vaccines to ensure safe use. The APVMA may also impose conditions on a permit for the use of veterinary vaccines for research purposes.
8. The Regulator notes that as part of their safety assessment, the APVMA considers viral shedding and transmission to other susceptible birds not included in the field trials, as well as the potential for recombination. The Regulator does not assess vaccine excipients and would not assess manufacturing by-products and impurities unless they are GM products.
9. FSANZ develops the food standards in the Food Standards Code with advice from other government agencies and input from stakeholders. The Standards in the Food Standards Code are legislative instruments and the Food Standards cover the composition of some foods, such as dairy, meat and beverages. FSANZ is also responsible for labelling of packaged and unpackaged food, including specific mandatory warnings or advisory labels.
10. Food Standards are enforced by the states and territories (usually their health or human services departments) or, in some cases, by local government. These authorities regularly check food products for compliance with the Food Standards Code.
11. FSANZ has developed the Primary Production and Processing (PPP) Standard for Poultry Meat (Standard 4.2.2) (FSANZ 2010). PPP Standards (which only apply in Australia) aim to strengthen food safety and traceability throughout the food supply chain from paddock to plate. The standard introduces new legal safeguards for growing live poultry and requires poultry growers to identify and control food safety hazards associated with poultry growing. Poultry processors are also required to identify and control food safety hazards associated with poultry processing (which includes the slaughtering process) and verify the effectiveness of the control measures.
	1. Background to the DIR application
12. Bioproperties Pty Ltd (Bioproperties) proposes to conduct field trials using a live attenuated GM infectious larygotracheitis virus (ILTV) vaccine to inoculate broiler chickens. The GM vaccine to be trialled has a product name of Vaxsafe® ILT. This vaccine has been developed to protect chickens against infectious laryngotracheitis disease.
13. The APVMA has issued a permit to Bioproperties to supply and use the GM vaccine for the purpose of animal research[[1]](#footnote-2). The GM vaccine is a new veterinary chemical product that has never been used previously as a registered veterinary product in Australia or elsewhere in the world.
14. Broiler farms, potentially including free range farms, in rural Victoria and NSW would be selected to participate in the field trials. Up to 2,000,000 chickens would be inoculated with the GM vaccine over a 5 year period.
15. The most likely route for administration of the GM vaccine would be via drinking water, although the option of delivery by eye drop has also been included in the application. The GM vaccine would only be administered by a suitably trained person such as a farm manager under the supervision of a registered veterinarian or qualified personnel.
16. As is common in veterinary vaccine trials, unless otherwise indicated on the APVMA permit, treated production animals would be allowed to enter the food chain. At an appropriate time, the chickens inoculated by the GM vaccine would be transported from farms to poultry processing plants. The processed chickens would normally be used for human and animal consumption.
	1. The proposed field trials
17. Bioproperties proposes to conduct field trials to assess the efficacy of the GM vaccine for protection of chickens from infectious laryngotracheitis disease under field conditions, including likelihood of challenge with a range of distinct field strains. The field trials would also assess the safety of the vaccine including the capacity for transmission and recombination with other available live ILTV strains.
18. The dealings assessed by the Regulator are:
* conduct of experiments with the GMO;
* transport the GMO;
* disposal of the GMO; and

the possession, supply or use of the GMO for the purposes of, or in the course of, any of the above.

* + 1. The proposed limits of the field trials (duration, scale, location and people)
1. The field trials are proposed to take place at approximately 40 selected broiler farms in rural Victoria and NSW, where intensive poultry production are concentrated. The trials would run over a 5 year period from the date of issue of the licence until the trials have completed assessment of the efficacy and safety of the vaccine. Up to 2 million chickens are expected to be vaccinated. The GM vaccine would be administered by appropriately trained farm personnel in accordance with trial protocols and under the supervision of a registered veterinarian or qualified personnel.
	* 1. The proposed controls to restrict the spread and persistence of the GMO in the environment
2. The applicant has proposed a number of controls to restrict the spread and persistence of the GMO in the environment. These include:
* only vaccinating broiler chickens on commercial chicken farms, excluding layers and breeders
* employing strict biosecurity measures that commercial broiler farms typically follow, such as supplying and wearing overalls and high rubber boots to all shed visitors and workers, and disinfecting hands and boots when entering and exiting the shed
* controlling access and movement of vehicles and people at the farm
* disinfecting all contaminated equipment and materials such as bottles, vials, droppers, feed containers, water lines and tanks after use
* cleaning and disinfecting the shed after removal of a vaccinated flock and before another unvaccinated flock is introduced into the shed
* disposing litter and dead chickens by composting, burial, rendering or landfill following State/Territory and/or local council requirements.
1. In addition to the above controls, the APVMA permit also has a number of conditions to restrict the spread and persistence of the GMO in the environment, such as managing populations of pests (e.g. dogs, cats, rodents, wild birds and darkling beetles), and disinfecting sheds, vehicles and equipment after use.
	* 1. Details of the proposed activities
			1. Selection of chicken farms
2. The field trials would take place in rural and semi-rural Victoria and NSW, where broiler farms are mainly concentrated. Conventional shed-based and free range broiler farms would be selected to participate in the trial from within the local government areas listed in Table 1.

Proposed local government areas

| **New South Wales** | **Victoria** |
| --- | --- |
| Lake MacquarieCentral CoastHawkesburyPenrithLiverpoolCamdenWollondilly | Yarra RangesMornington PeninsulaSouth GippslandCardiniaCaseyGeelongColac OtwayGolden PlainsSurf CoastBulokeGannawarraLoddonCampaspeCentral GoldfieldsMount AlexanderMacedon RangesCity of Greater BendigoHindmarshWest WimmeraYarriambiack |

1. The first phase of the trial is expected to be restricted to specifically-selected farms in Victoria that do not currently vaccinate, allowing assessment of vaccine safety by comparison with unvaccinated control sheds on the same farm.
2. Further, trial locations would be decided by Bioproperties, in consultation with farm managers. Specific locations of participating farms would be notified to the OGTR before any dealings with the vaccine commence at that site.
	* + 1. Study design
3. Over the 5 year period, up to 2,000,000 broiler chickens would be vaccinated with the GM vaccine, representing approximately 40 farms or sheds, each holding approximately 50,000 chickens. The farm or shed would be the ‘experimental subject’. This number of subjects is necessary to detect a small to moderate difference in mortality rate where the incidence of natural field challenge with ILTV is low.
4. The safety and efficacy of the GM vaccine would be assessed on a farm or shed basis depending on how the vaccine is allocated. Where a farm has multiple sheds, each shed may be randomly assigned to receive one of the ILTV vaccines, either the GM vaccine or another APVMA-registered vaccine against ILTV as an active control. There may be a shed(s) not vaccinated against ILTV as a negative control. However, the use of an unvaccinated control shed would not be considered on a free range farm where maintaining physical separation would not be possible. Where a whole farm is vaccinated with the GM vaccine, a comparison would be made with other farms receiving the active or negative control treatment.
5. In the first phase of the trial, which is expected to last no longer than 12 months, a few farms (equivalent to up to 500,000 chickens) would be selected that do not currently vaccinate against ILTV and are more isolated from other poultry farms. These farms are ideal to assess the transmission and safety characteristics of the vaccine under field conditions. Approximately 40 broiler chickens not inoculated with the GM vaccine or another vaccine against ILTV (sentinel chickens) may be housed in the same shed containing chickens inoculated with the GM vaccine to study transmission characteristics. The sentinel chickens would be from the same batch of chickens inoculated with the GM vaccine. In the second phase of the trial, more farms would be selected in areas that have experienced ILTV outbreaks to assess the efficacy of the GM vaccine under field conditions.
	* + 1. Manufacture, supply and storage of the GMO
6. The GM vaccine would be manufactured at Bioproperties’ manufacturing facilities in Glenorie, NSW, which are APVMA-licenced and also certified by the Regulator. Manufacture would be done according to the *Australian Code of Good Manufacturing Practice (GMP) for Veterinary Chemical Products* (APVMA 2007).
7. The GM vaccine would be transported to Ringwood, Victoria for storage, then to a poultry company’s vaccine storage area before being distributed to the farms included in the field trials using couriers. The GM vaccine would be supplied as a freeze-dried pellet in a glass vial. These would be placed in trays, wrapped in plastic cling wrap, and placed into a Styrofoam box filled with dry ice and sealed with packaging tape. The Styrofoam box would be placed into a cardboard box. The primary container and external containers of the vaccine would be labelled to indicate the APVMA permit number, contents, purpose and storage requirements that have been approved by the APVMA. Transport and storage would be in accordance with the PC2 requirements of the Regulator’s *Guidelines for the Transport, Storage and Disposal of GMOs.*
8. Once the GM vaccine has arrived at the farm, the vaccine would be stored in a freezer (below ‑18°C) specifically used for veterinary medicines. The vaccine must be kept in a freezer or on ice until it is ready to be reconstituted.
	* + 1. Preparation and administration of the GMO
9. The GM vaccine would be used for inoculation of broiler chickens only. Long lived breeder or layer chickens would not be included in the trial. Broiler chickens would be inoculated between 7 to 14 days of age. The product leaflet approved by APVMA specifies how the vaccine must be used. Most of the broiler chickens in the shed would be vaccinated with the GM vaccine, while the sentinel chickens in the same shed would not be vaccinated against ILTV in order to study transmission of the GM vaccine to unvaccinated chickens. The sentinel chickens housed in the shed with GMO-inoculated chickens would be prevented from drinking the water containing the GM vaccine if administered by drinking water. The broiler chickens receiving the GM vaccine will only be vaccinated once and will not have received vaccination with any other ILTV vaccine. No other ILT vaccine would be given to the flock after vaccinating with the GM vaccine.
10. Chickens in the commercial industry are routinely vaccinated against bacterial and other viral pathogens. Details of all vaccinations would be recorded and maintained, as standard practice of the poultry company.
11. The GM vaccine needs to be reconstituted with sterile cold water prior to administration. Reconstitution of the vaccine would take place in a room or area adjacent to the shed where the water tanks or water pipe controls are located.
12. Each vial would contain approximately 108 plaque forming units (PFU) of live GM vaccine, representing 1000 vaccine doses. Each chicken would receive approximately 105 PFU. The APVMA permit allows vaccination by either eye drop or drinking water, but the most likely route would be via drinking water because this is a more efficient way to inoculate large numbers of birds.
13. Preparation and vaccination would be conducted by the farm manager with the aid of an assistant who may be supervised by a registered veterinarian or qualified, trained or experienced personnel.
14. For eye drops, the pellet must be reconstituted in 30 mL of water and 30 microlitres delivered to the eye using a dropper.
15. To prepare the vaccine for drinking, the volume of water to be consumed in 3 hours by all chickens in the shed must be calculated before reconstituting the vaccine. Sufficient doses of the vaccine for the whole flock would be reconstituted in a tank of water with skim milk as a stabiliser. The tank is connected to the drinking troughs or drinking nipples within the shed. No additional water would be provided until all of the vaccine-containing water has been consumed.
	* + 1. Sample collection
16. Vaccinated chickens and unvaccinated sentinel chickens included in the trial would be monitored throughout the rearing period. Data on mortality rates at the farm would be collected. The registered veterinarian may conduct post-mortem examination and collect samples at the farm to determine the cause of mortality.
17. Tracheal swabs from randomly selected vaccinated or unvaccinated sentinel chickens and faecal samples from the litter would be collected at various time points after vaccination.
18. The samples collected from vaccinated and unvaccinated sentinel chickens would be transported as biological specimens, and in accordance with the Regulator’s *Guidelines for the Transport, Storage and Disposal of GMOs.* Samples would be taken to facilities certified by the Regulator for testing and analysis.
19. Samples would be tested for the GMO or wild type ILTV DNA by quantitative polymerase chain reaction (qPCR). This would be performed until the likely extent of the persistence of the GMO has been determined, and would not be performed on all chickens.
20. Some of the live vaccinated and unvaccinated sentinel chickens would be transported from the farm to a certified facility in the University of Melbourne for research purposes. The tests, experiments and analyses undertaken at the University of Melbourne involving the live vaccinated and unvaccinated sentinel chickens and samples would be conducted under a Notifiable Low Risk Dealing (NLRD) authorised by the University of Melbourne’s institutional biosafety committee (IBC). The live vaccinated and unvaccinated sentinel chickens would be transported according to the IBC and animal ethics committee requirements.
	* + 1. Personal protective clothing
21. Commercial broiler farms may supply visitors and veterinarians with disposable overalls, hairnets and high rubber boots for use in poultry shed and free-range enclosures. Farm workers, including workers collecting chickens for transport, are required to wear clean, laundered work clothes each day or may wear disposable overalls.
22. Farm workers and manager preparing the vaccine would wear gloves in addition to eye protection. The product leaflet recommends wearing eye protection and masks when preparing the GM vaccine.
23. Veterinarians conducting post-mortem examination at the broiler farm routinely wear disposable gloves and overalls and dedicated footwear, and disinfect hands and footwear after examination.
	* + 1. Decontamination and disposal of the GMO
24. The poultry farm may have a shed or building used for entering and exiting the farm area. The shed or building contains a hand wash basin, hand sanitiser, a footbath, visitor log book, change room, and personal protective clothing, such as disposable overalls, rubber boots and hairnets. A typical shed for housing chickens has an anteroom that is routinely used for entering and exiting the shed. A footbath filled with fresh disinfectant and a hand wash basin or hand sanitiser are available in the anteroom. When entering and exiting the broiler shed, hands and boots would be disinfected against the GMO. The supplied disposable overalls would be disposed of via the normal farm waste bin when exiting the farm and the rubber boots remain on the farm.
25. Multiple sheds on any farm containing chickens, treated with the GM vaccine or any of the other active controls or no ILTV vaccine treatment, would be clearly identified. To minimise cross-contamination, measures would be implemented including housing and managing each treatment group separately, and decontaminating equipment or materials when entering and exiting each shed.
26. The APVMA permit states that water tanks, tubing or eye droppers used to deliver the product must be treated between flocks with an agent effective against the vaccine virus. The applicant proposes that after use, bottles, droppers, vials and other materials contaminated with the GM vaccine would be soaked in disinfectant solution such as Virkon (1% v/v), sodium hypochlorite (0.5%) or quaternary ammonium chloride (0.01% v/v). After soaking in disinfectant, the waste materials would be wrapped in paper and placed in regular waste bins.
27. After all the chickens have been removed from the shed and during full shed clean-out, water lines and tanks used for drinking water vaccination would be cleaned with commercial virucidal oxidising agents such as iodophore, chlorine dioxide, or stabilised hydrogen peroxide-based products. These would be added to the water tanks at an appropriate concentration and allowed to run through the water lines. The solution would be held within the water lines for the recommended contact time and then flushed using chlorinated water.
28. Due to the short growing period of broilers in the current industry, the re-use of litter for more than one flock is common practice in the industry. For the field trials, the applicant proposes to re-use litter only where the first flock is not vaccinated with ILT or where a vaccinated flock is followed by a vaccinated flock of the same vaccine. The applicant proposes that full cleanout of the shed and removal of litter would occur before an unvaccinated flock replaces a flock vaccinated with the GM vaccine. However, the APVMA permit states that the shed and litter are to be treated between flocks in a manner which is effective against the vaccine virus.
29. The applicant proposes to wash and disinfect vehicles, equipment, crates and bins after use with detergent and disinfectant solution.
30. For temporary storage of litter for disposal, dispersal would be restricted by covering the heaped litter with clean co-composting material and a tarpaulin. Chicken carcasses may be stored temporarily in a freezer prior to disposal by waste contractors or composting on farm land. For composting on farm land, the compost would be left for 3 to 6 months to ensure completion of the composting process.
31. The disposal of farm waste such as litter and chicken carcasses varies for each farm or poultry company. State legislation and local councils have requirements for disposal of waste generated in poultry farms, including free range farms (see Section 7).
	* + 1. Training of personnel
32. The entire field trial would be managed by a registered veterinarian consultant contracted by Bioproperties who manages the overarching protocol. The protocol for each site would be a version of the overarching protocol, modified to include the site location, names and contact details of the personnel participating at that site, and any minor changes required to accommodate farming practices at that site. A copy of the DIR licence would be attached to the protocol. The protocol would be prepared by Bioproperties Research and Development, and reviewed by Bioproperties Regulatory Affairs to ensure compliance with all regulatory conditions including the APVMA permit conditions and licence conditions that would be imposed by the Regulator. The Quality Assurance Manager would ensure that the protocol meets the Good Clinical Practice guidelines developed under the principles of International Cooperation on Harmonization of Technical Requirements for Registration of Veterinary Medicinal Products (VICH 2000). Each site would have a registered veterinarian who would ensure that the protocol is followed. The site veterinarian may be employed by the poultry company. All relevant personnel, their role and responsibilities, would be clearly indicated in the trial protocol.
33. The trial site protocol and the product leaflet also describe decontamination measures in response to spills of the GM vaccine. All farm workers and farm managers would be trained in decontaminating spills.
34. All workers responsible for handling the vaccine, inoculated chickens and contaminated equipment would be trained in handling the GMO. All farm workers would be trained in decontamination and disposal of the GMO in accordance with the trial protocol, any licence conditions imposed by the Regulator and the APVMA permit conditions.
	* + 1. Adverse events
35. The APVMA permit requires that Bioproperties must maintain a record of any adverse events, which includes any issues with the quality and safety of the product, and veterinary treatment must be sought as necessary.
36. Adverse events to vaccinated chickens would be reported to the Bioproperties Animal Care and Ethics Committee (ACEC) and the Regulatory Affairs Manager. The ACEC would provide advice on the steps to be taken to minimise any harm occurring to the chickens used in the field trials.
	* + 1. Record keeping
37. The APVMA requires that Bioproperties maintain a record of the trials performed under the permit. Specifically, details must include the date and location where the trials are conducted, commodities treated, rates and frequency of application, total amount of product used, and the names and addresses of persons conducting the trial, and any adverse events. These details must be maintained for a minimum period of two years from the date of expiry of the permit and must be made available to the APVMA upon request.
	* + 1. Fate of chickens after field trials
38. Vaccinated and unvaccinated sentinel chickens that have reached the appropriate age for harvesting, which could be as early as 28 days of age, would be transported and processed in the same way as for other commercial broiler chickens (see Section 7).
	1. Parent organism
39. The parent organism of the GMO is *infectious laryngotracheitis virus* (ILTV). The ILTV strain, CSW-1, from which the GMO was derived, was originally isolated from the Glenfield, NSW outbreak in 1959.
40. ILTV is a member of the *Iltovirus* genus of the subfamily *Alphaherpesvirinae,* family *Herpesviridae*. ILTV is also known as *Gallid herpesvirus 1* (Thureen & Keeler 2006).
41. ILTV infects the trachea and conjunctiva causing respiratory disease and conjunctivitis in chickens, although pheasants, peafowl and turkeys can also be infected with ILTV (Crawshaw & Boycott 1982; Portz et al. 2008). The virus can establish latent infection in the neurons innervating the trachea and be re-activated by stress (Garcia et al. 2013; Hidalgo 2004; Ou & Giambrone 2012). ILTV does not infect humans or other animals.
42. ILT disease has been known to affect poultry for decades. ILT disease was first reported in 1925 in Canada, followed by the USA in 1926, Australia and Great Britain in 1935, and Europe in 1940. By 1962, the disease had been described in at least 40 countries across North and South America, Middle East, Africa, Australia and Asia (Agnew-Crumpton et al. 2016; Blacker et al. 2011; Menendez et al. 2014; Moreno et al. 2010; Sellers et al. 2004; Volkova et al. 2012; Linares et al. 1994).
43. ILT disease is notifiable under NSW (*Stock Diseases Act 1923*) and Victorian (*Livestock Disease Control Act 1994*) legislation, creating a legal obligation to notify authorities if an animal is known or suspected of having ILT disease.
44. There is no treatment for ILT disease. Vaccines are currently used to protect chickens from ILT disease. Two of the currently available live attenuated vaccines in Australia, A20 and SA2, were derived from an Australian ILTV strain. The Serva vaccine strain, also used in Australia, was derived from a European strain. These strains are able to recombine with each other and with wild type ILTV resulting in virulent ILTV that have caused outbreaks in Australia.
	* 1. Basic Biology
45. ILTV has a linear double-stranded DNA genome approximately 155 kilo base pairs (kb) in length. The genome consists of unique long (UL) and short (US) segments, and inverted repeats – internal repeat and terminal repeat (IR and TR). The IR and TR flank the US region (see Figure 2) (Johnson et al. 1991). ILTV possesses three origins of viral DNA replication, one (OriL) located within the UL region, and two OriS within the US region (Fuchs et al. 2007).

**UL**

**US**

ORIL

ORIS

ORIS

**IR**

**TR**

Figure 2. Organisation of ILTV genome

Wild type ILTV genome organisation. IR: internal inverted repeat. ORI: origin of replication. UL: unique long. US: unique short. TR: terminal inverted repeat.

1. The virus is comprised of four distinct structural elements: envelope, tegument, capsid and core. The lipid envelope contains glycoprotein spikes which are responsible for stimulating humoral and cell-mediated immune responses. ILTV encodes ten structural glycoproteins (Thureen & Keeler 2006). Contained within the envelope is the capsid, which surrounds the core comprising the viral DNA packed at high density without internal proteins. The region between the envelope and capsid is the tegument which contains more than 20 viral proteins likely to have important roles in modulating virus-host interactions. The size of the particle varies between 200 and 350 nm, since ILTV incorporates large but variable amounts of tegument proteins (Fuchs et al. 2007; Davison & Clements 2009).
2. Infection is initiated by attachment of virus glycoprotein to the cell membrane receptor, heparan sulphate, followed by fusion of the envelope with the host cell plasma membrane. Within the cell, viral DNA is released from the capsid and migrates into the nucleus through nuclear pores.
3. Transcription and replication of viral DNA occur within the nucleus but the viral DNA does not integrate into the host genome. Transcription of ILTV DNA occurs in a highly regulated, sequentially ordered cascade similar to that of other alphaherpesviruses. The first ILTV peptides are detectable at 4 hours after *in vitro* infection of chicken kidney cells and ILTV DNA replication commences between 8 and 12 hours post-infection (Prideaux et al. 1992). Several of the virus-encoded proteins are enzymes and DNA-binding proteins that regulate viral DNA replication, but most are viral structural proteins. Viral DNA replication occurs by a rolling circle mechanism with the formation of concatemers which are cleaved into monomeric units and packaged into preformed capsids within the nucleus (Hidalgo 2004).
4. DNA-filled nucleocapsids acquire an envelope as they migrate through the nuclear membrane to the cytoplasm where the capsids associate with the tegument proteins. The nucleocapsids are re-enveloped in the Golgi region, followed by release of mature particles by exocytosis (Fuchs et al. 2007; Hidalgo 2004). ILTV growth was studied in primary chicken kidney cells *in vitro*. At 11 hours after infection of chicken kidney cells, little or no viable ILTV was produced. Between 11 hours and 24 hours after infection of chicken kidney cells, a logarithmic increase in ILTV titres was observed (Prideaux et al. 1992).
5. The natural route of entry for ILTV is through the upper respiratory tract and conjunctiva. Ingestion may be another mode of infection, although exposure of nasal epithelium following ingestion is necessary for infection (Garcia et al. 2013). ILTV infects the trachea and conjunctiva, but other tissues may be susceptible as well. The DNA of SA2 and A20 vaccine strains were detected by qPCR in the chicken’s Harderian gland, lungs and kidneys over a 28 day period after oral administration of these vaccines (Roy et al. 2015). DNA of various ILTV strains including an ILTV vaccine were detected in the conjunctiva, sinuses, trachea, cecal tonsils, thymus and cloaca, with peak genome copy numbers detected at 4-5 days post-inoculation (Oldoni et al. 2009).
	* 1. Host range
6. The chicken is the primary host and reservoir for ILTV.
7. Natural infection with ILTV has also been observed in pheasants and peafowl (Crawshaw & Boycott 1982). ILT disease and some mortality in peafowl and pheasants have been reported in a shed in Canada housing many bird species. ILT disease also occurred in Malay argus pheasants and chickens, but not in other species housed in the same shed including ocellated turkeys, Chinese painted quails, common guineafowls, fulvous tree ducks, African hornbills and several species of macaws and cockatoos (Crawshaw & Boycott 1982).
8. Turkeys are also naturally infected with ILTV, displaying clinical signs of nasal discharge, marked dyspnea, depression and tracheitis. Turkeys inoculated intratracheally with ILTV showed similar clinical signs of the disease (Portz et al. 2008).
9. Attempts to experimentally infect other birds such as ducks, starlings, sparrows, crows and pigeons have been largely unsuccessful (Beach 1931).
10. Ducks experimentally exposed to ILTV did not show signs of disease and appeared normal. However, neutralizing antibodies against the ILTV were detected 7 to 14 days after exposure. Growth of ILTV in embryonated duck eggs experimentally inoculated with ILTV was achieved (Yamada et al. 1980).
11. ILTV is not known to infect humans, other non-avian vertebrates or other organisms including invertebrates, plants, microorganisms and aquatic organisms.
	* 1. Clinical signs
12. ILT is a viral respiratory tract infection which produces severe production losses due to decreased weight gain, decreased egg production and mortality of infected chickens. Clinical signs generally appear between 6-12 days following natural exposure (Hidalgo 2004; Bagust et al. 2000). Experimental inoculation of a live attenuated ILTV vaccine or wild type strains onto the eye and into the nostril of chickens results in clinical signs to appear between 3-12 days post-inoculation (Oldoni et al. 2009). Generally, most chickens recover in 10-14 days after clinical signs begin to appear, but it can take as long as 3 weeks in some cases (Garcia et al. 2013).
13. Characteristic clinical signs include marked dyspnea, nasal discharge, moist rales, coughing, gasping, sneezing, depression, swelling of infraorbital sinuses and conjunctivitis. In severe forms of the disease, signs also include laboured breathing and expectoration of blood-stained mucus. Birds die from this disease due to suffocation, as the windpipe becomes completely blocked. Mortality rate varies between 5-70%, with most in the range of 10-20%, but can be as high as 90-100% in severe cases (Garcia et al. 2013; Hidalgo 2004; Ou & Giambrone 2012).
	* 1. Latency
14. Latency establishment after infection is the major biological survival mechanism of herpesviruses, enabling evasion of host immune surveillance. As is the case for other herpesviruses, ILTV establishes latent infections within their hosts. The latent infection is characterised by a shutdown of virus replicative functions and the inability to detect infectious virus. During latency, the double stranded DNA genomes are maintained as multiple copies of circular episomes within the nuclei of the cells in which they became latent (Avgousti & Weitzman 2015; Bloom et al. 2010; Wilson & Mohr 2012).
15. The trigeminal ganglion is the main site of latency for herpesviruses, including ILTV. The trigeminal ganglion provides the main sensory innervation to the tissues of the upper respiratory tract including the trachea. Chickens with latent infections that have recovered from ILT disease no longer showed signs of the disease. In clinically recovered chickens infected with ILTV, ILTV DNA was detected by PCR in the trigeminal ganglion on 31, 46 and 61 days post-infection (Williams et al. 1992).
16. The virus can remain latent until a stress stimulus causes some still unknown changes that can trigger transcription of the genes and ultimately lead to replication and production of virions, known as reactivation. The molecular mechanism of reactivation within a cell is dependent on many factors. Inducing gene expression from silenced promoters during reactivation from latency may be mechanistically distinct from activating gene expression during lytic infection (Avgousti & Weitzman 2015; Bloom et al. 2010; Wilson & Mohr 2012).
17. In a study involving chickens latently infected with CSW-1 ILTV, reactivation of the virus was found in six of 16 chickens over a period of three to 15 months following infection (Bagust 1986). In the same study, four of 9 chickens that had been vaccinated with the SA2 strain showed the reactivation of the vaccine strain over a period of two to 10 months. This study determined that wild and vaccine strains of ILT virus can establish long-term latent infections. However, the study could not demonstrate that immunosuppressive drugs induced reactivation of the virus. Similarly, no virus shedding was detected from the trachea of chickens after 6 weeks of receiving a different vaccine against ILTV and within a week after immunosuppressive drugs treatment (Hughes et al. 1991).
18. The effect of stress factors on virus shedding, including the onset of egg laying and rehousing with unfamiliar chickens, was studied in chickens latently infected with a wild type ILTV strain. About 6 weeks after ILTV inoculation and 3 weeks before the onset of lay, ILTV was isolated from the trachea of 2 out of 10 infected chickens and from only 2.5% of the tracheal swab samples. In the first week of the onset of lay, ILTV was isolated from 9 out 10 infected chickens and from 20% of tracheal samples, indicating a significant effect on viral shed from the stress of laying. Rehousing with unfamiliar chickens had a lesser effect, with ILTV being isolated from the trachea of only 1 out of 5 infected chickens. It appears that stress, in particular the onset of egg laying, may increase the ILTV shedding rate in a latently infected chicken (Hughes et al. 1989).
19. Most information on latency and reactivation comes from small animal models, principally mice and rabbits, which have been used to study latency and reactivation *in vivo* of herpes simplex virus-1 (HSV-1). Using one of the rodent *in vitro* culture systems of sympathetic neurons, it was suggested that there are two phases of gene transcription during reactivation induced by pharmacological agents. The first wave, termed phase I, occurs approximately 15-20 hours post-induction and leads to concurrent transcription of immediate-early, early and late genes. In phase I, no new viral protein synthesis or DNA replication is required, which begins at 25-30 hours post-induction. During phase II, viral DNA replication and synthesis of infectious viruses occur (Avgousti & Weitzman 2015; Bloom et al. 2010; Wilson & Mohr 2012).
20. To examine the influence of stress on the reactivation of latent HSV-1 in infected mice, the mice were exposed to heat stress. After 12 hours since being subjected to hyperthermia, immunolabelled cells in the trigeminal ganglion were found to be positive for HSV-1 antigens. Immunopositive cells increased at 24 hours and then decreased at 36 hours (Huang et al. 2011). In another study, mice latently infected with HSV-1 were subjected to heat stress and were sacrificed at 14, 24 and 48 hours after heat stress. Infectious virus was recovered from the trigeminal ganglion in 1 out of 10 mice at 14 hours after heat stress, reaching peak of 60% (6 out of 10) of mice at 24 hours. By 48 hours after heat stress, infectious virus was recovered from 20% of mice (Sawtell & Thompson 1992). However, it should be noted that there may be animal and virus species differences between HSV-1 and ILTV.
	* 1. Shedding
21. ILTV is shed from infected birds from various sites. Virus shed in the trachea can be released into the environment by aerosolisation or expectoration of tracheal exudates. As discussed above, the rate of virus shed following latent infection can be increased in situations of stress such as egg laying (Bagust et al. 2000; Hughes et al. 1989).
22. Chickens infected with various wild type ILTV strains shed the virus from the conjunctiva, sinuses and trachea between 2 and 9 days post-inoculation. Shedding at these sites was detected by PCR and viral isolation using chicken kidney cells. For most of the wild type ILTV strains, the peak of viral replication (102-106 genome copy numbers/reaction) detected by PCR occurred between 4-5 days post-inoculation, with only one wild type ILTV strain having a peak of viral replication (106 genome copy numbers/reaction) at 9 days post-inoculation. No infectious virus was isolated from the conjunctiva, sinus and trachea at 11 and 14 days post-inoculation (Oldoni et al. 2009).
23. Shedding of two live attenuated ILTV vaccine DNA have been detected by PCR at various sites in the chicken with DNA levels varying depending on the tissue and ILTV vaccine. DNA of ILTV vaccines were detected in the conjunctiva and trachea from 2-14 days after chickens were vaccinated. Peak genome copy numbers (106 genome copy numbers) were detected from 4-8 days post-vaccination in the conjunctiva and trachea, declining several-fold logarithmically to low genome copy numbers (102 genome copy numbers) at 14 days post-vaccination. ILTV genome copy numbers remained steady ranging from 101.94 to 102.7 in the cecal tonsils from 2 to 8 days and at 21 days post-vaccination. ILTV DNA was detected from the cloaca from 5 to 9 days post-vaccination, ranging from 101.44 to 102.6 genome copy numbers, and was undetectable from 10 days post-vaccination. Infectious virus was isolated from only two sites, the conjunctiva and trachea from 2-6 days post-vaccination, and no infectious virus was isolated from 7 to 28 days post-vaccination (Rodriguez-Avila et al. 2007).
24. The DNA levels of the SA2 and A20 vaccine strains have been detected in the Harderian gland, trachea, lung and kidneys up to 28 days after vaccination of chickens. At 6 days post-vaccination, higher DNA levels of the vaccine strains (105-109.8 copy numbers/mg tissue) were measured in the Harderian gland, trachea, kidneys and lung compared to 14 days post-vaccination, in which DNA levels declined by 2-6 logs lower. DNA of the vaccine strains were detected in the faeces from 2-28 days post-vaccination, with a peak (1010 copy numbers/g faeces) at 5 days post-vaccination, and declined thereafter by about 3-4 logs at 14 days post-vaccination (Roy et al. 2015). It is unclear if the vaccine DNA detected from these studies represents infectious virus particles.
	* 1. Transmission
25. Transmission of ILTV can occur via contact with ILTV shed in tracheal exudates, contaminated inanimate objects such as equipment and clothing, contaminated litter, manure and infected carcasses. Egg transmission of the virus has not been demonstrated. The virus may spread by aerosol movement or wind (Garcia et al. 2013; Hidalgo 2004; Bagust et al. 2000).
26. Risk factors have been identified that may have led to outbreaks in broiler farms in Mississippi, USA during 2002-2003. Based on the responses to a retrospective survey questionnaire, the report found that farm suppliers such as gas company representatives, who are likely to visit farms, and farm workers who visit other chicken farms, are likely vehicles of ILTV introduction onto broiler farms. Sharing of equipment used to remove broiler litter between subsequent flocks may also serve as an important vehicle of ILTV transmission. During the outbreak, shared litter removal equipment was associated with ILTV transmission despite a requirement being put in place for litter decontamination. Tunnel-ventilated broiler houses with inlets toward neighbouring poultry farms are more likely to get infected with ILTV. The report suggested risk mitigation measures including following biosecurity procedures, showering and changing footwear prior to entering broiler houses on their own farm, and that practices such as wearing plastic boots or changing boots may be more effective than footbaths in preventing ILTV transmission (Volkova et al. 2012).
27. The larvae and adult darkling beetles (*Alphitobius diaperinus*) are prevalent in poultry facilities. These beetles consume feed, water, poultry carcasses and faeces. The beetles live in compacted earth and litter, and can damage poultry house structures. The darkling beetles lay eggs in the bedding litter producing larvae that live in the litter, predominantly under feed pans. The larva then burrow into the earth floor of the broiler shed to pupate, and from the pupa the adult darkling beetle emerges. Earth floors of broiler houses are an important medium for pupation, but darkling beetles still occur, albeit in smaller numbers, in broiler sheds with higher density flooring materials such as concrete and bitumen (Poultry Hub 2017). Some strains of darkling beetles are resistant to certain insecticides with some adult strains being more resistant than larvae to certain insecticides (Hamm et al. 2006). Chickens may consume beetles rather than feed. ILTV DNA and virus were detected in adult beetles and larvae taken from the farms up to 42 days after an ILTV outbreak in commercial poultry farms in the USA. Ingestion of ILTV-positive beetles could lead to infection of chickens and therefore may serve as a source of ILTV transmission. The study did not show that beetles were infected with ILTV (Ou et al. 2012).
	* 1. ILTV vaccines
28. The vaccines now commonly used in commercial poultry flocks worldwide include attenuated live vaccines developed by consecutive passage of virulent virus in cell cultures (tissue culture origin [TCO]) or in embryonated hen eggs (chicken embryo origin [CEO]). Recombinant vaccines have also been produced using *herpesvirus of turkeys* (HVT) or *fowlpox virus* (FPV) modified to express ILTV glycoproteins that can elicit protective immune responses in vaccinated birds, and are now used commercially in some poultry-producing regions around the world (Coppo et al. 2013).
29. Currently in Australia, there are three APVMA registered vaccine strains against ILT: A20, SA2 and Serva vaccine strains. The three vaccines are live attenuated CEO vaccines.
30. The SA2 vaccine strain is an attenuated ILTV field strain of Australian origin. The A20 vaccine strain was produced by serial passages of the SA2 strain in primary chick embryo cell cultures and embryonated eggs in order to decrease its residual virulence. The Serva vaccine strain originated in Europe.
	* 1. ILTV classes and recombination between types
31. Based on the data obtained from whole genome sequence analysis of the A20, SA2 and Serva vaccine strains, SA2 and A20 genomes are divergent from the Serva genome with only 99.2% of the sequence identical to Serva genome (Lee et al. 2011a; Lee et al. 2011b).
32. The genome size of CSW-1 ILTV strain (151,671 bp) is smaller compared with the Serva strain (153,645 bp) due to large deletions within the UL region and in both the internal and terminal inverted repeats (Lee et al. 2013).
33. Using the BLAST online tool (National Center for Biotechnology Information (NCBI) 2017), the nucleotide sequence identity of the whole genome of CSW-1 strain[[2]](#footnote-3) was compared with other ILTV strains. The results showed that CSW-1 has 99.82% identity with the Serva[[3]](#footnote-4), 99.70% identity with the SA2[[4]](#footnote-5) and 99.69% identity with the A20 strain[[5]](#footnote-6).
34. Sequence analysis revealed that the UL21, 32, 34 and 43 genes of CSW-1, Serva and SA2 strains share 100% nucleotide and amino acid sequence identity (Lee et al. 2013). The ICP4, UL27, UL36, US5 and US8 genes showed the greatest nucleotide and amino acid sequence variability among the three ILTV strains. The phylogenetic relationships between CSW-1, SA2 and Serva strains vary depending on which gene was analysed.
35. ILTV strains can be categorised into different classes based on restriction fragment length polymorphism (RFLP) PCR of certain ILTV genes and genomic regions. ILTV strains with the same RFLP pattern were placed into one class. In Australia, this method has been used to identify ten different genotypes or classes of ILTV (Agnew-Crumpton et al. 2016; Blacker et al. 2011; Kirkpatrick et al. 2006).
36. The A20 and SA2 strains belong to class 1. ILTV classes 2, 3 and 5 comprise other strains isolated from outbreaks in commercial flocks in Australia and were found to be distinct from class 1. Class 4 comprises the CSW-1 strain (Blacker et al. 2011; Kirkpatrick et al. 2006).
37. Class 7 corresponds to the Serva vaccine strain. ILTV classes 8 and 9 are phylogenetically close to class 7, indicating a close genetic relationship between these classes (Blacker et al. 2011). Furthermore, a comparative sequence analysis has revealed that class 8 is also largely similar to both A20 and SA2, while class 9 was derived from recombination between A20 and Serva strains. The results suggest that recombination occurred between the co-circulating A20, SA2 and Serva strains giving rise to class 8 and 9. Furthermore, classes 8 and 9 are more virulent than their parent strains when studied *in vivo* in chickens (Lee et al. 2012).
38. Class 10 was isolated from Australian disease outbreaks in NSW in 2013. The samples used in this analysis were obtained mostly from commercial poultry flocks and a few backyard flocks. These flocks were vaccinated with one or a combination of the three available ILTV vaccines. Analyses of class 10 revealed a mosaic pattern, with some regions showing a high level of identity to specific field or vaccine strains of ILTV, while other regions were identical to a different field or vaccine strain. Class 10 shares genomic regions with classes 1, 7, 2 and 8 suggesting that class 10 may have emerged as a result of recombination events involving a previously recombined ILTV class (Agnew-Crumpton et al. 2016).
39. It is possible that recombination may have been facilitated by the conditions under which the ILTV vaccines were used, including the introduction of the European Serva strain into the Australian environment, using a combination of the three different ILTV vaccines on a single flock contrary to the APVMA approved directions for use[[6]](#footnote-7), inappropriate use of the vaccines and the mass delivery of multiple vaccines to large numbers of intensively housed birds. This finding highlights the risk associated with the use of multiple attenuated ILT vaccines under conditions imposing high selective pressures, which may foster recombination between co-circulating viruses and selection of more virulent or transmissible progeny (Agnew-Crumpton et al. 2016; Coppo et al. 2013).
	* 1. Recent outbreaks in Australia
40. ILTV outbreaks in chicken farms that commonly recur in Victoria and NSW have been caused by different classes of ILTV. From 2007 to 2009, ILTV class 2 was responsible for a large number of outbreaks mainly in Victoria. Class 8 was responsible for the majority of outbreaks in NSW in the same period. Class 4 (CSW-1) and 5 were not identified as causing outbreaks in NSW or Victoria during this period (Blacker et al. 2011).
41. Between 2009 and 2015, however, class 2 was very rarely detected and instead class 9 caused the largest number of ILTV outbreaks in Victoria. Since 2009, except for 2011, class 8 was replaced by class 9 as the predominant ILTV class to cause outbreaks in Australia, including NSW. Class 4 (CSW-1) was not identified as causing outbreaks in Australia between 2009 and 2015 (Agnew-Crumpton et al. 2016).
	* 1. Environmental stability and decontamination methods
42. Survival of ILTV in different conditions varies depending on the amount of virus initially present, the medium in which it occurs, pH, temperature and exposure to light. At 13-23°C, ILTV in tracheal exudates survived up to 110 days in the dark, but this was reduced to 7 hours in direct sunlight. At 4-10°C in the dark, ILTV in the trachea of chicken carcasses survived for 30 days. At 4°C, ILTV survived in desiccated tracheal exudate for at least 24 years (Jordan 1966).
43. ILTV is sensitive to heat, ether, chloroform, and other lipolytic solvents. The virus was destroyed in 1 minute by treatment with 3% cresol or a 1% lye solution. Storage media containing glycerol or sterile skim milk greatly increases survival (Ou & Giambrone 2012).
44. ILTV vaccine DNA has been detected at high levels in dust from laboratory chicken cages at 28 days after inoculation of chickens with either A20 or SA2 vaccine. Litter samples from these laboratory chicken cages also contained ILTV vaccine DNA which was shed from the vaccinated chickens (Roy et al. 2015). However, the infectivity of the vaccine strains detected from dust and litter was not investigated.
45. Biofilms in drinking water lines have been suspected of being a source of ILTV in the field. A common method of administering ILTV vaccine is through the drinking water. After running ILTV vaccine mixed with water into lines and flushing the lines with tap water three times, ILTV vaccine DNA was still detected from the lines for up to 21 days. Chickens drinking from this water line tested positive for ILTV DNA up to 21 days after flushing with water. After the vaccine application and flushing with water, a sanitising solution was held for 24 hours in the water lines and then flushed with tap water. A comparison of the different sanitising solutions revealed that ILTV vaccine was not detected in the water lines after sanitising with sodium bisulfate (0.31 mL/L) or hydrogen peroxide (30 mL/L) solution. However, ILTV vaccine was still detected after treatment with citric acid (3.05 mL/L) or sodium hypochlorite (0.19 mL/L). Chickens tested positive for ILTV DNA after drinking from the water lines treated with citric acid or sodium hypochlorite, while they tested negative after sodium bisulfate or hydrogen peroxide (Ou et al. 2011).
46. After an ILT outbreak in California affecting over 50 chicken farms, it was shown that ILTV was no longer isolated from chickens introduced into the farms that employed a thorough decontamination regime. This regime involved heating the farm shed to a minimum of 37°C for 100 hours, thorough cleaning and disinfection of the farm facilities and all equipment, heating again to a minimum of 37°C for 100 hours and downtime of 21 days of not letting flocks into the farm (Chin et al. 2009).
47. Litter containing ILTV heated at 38°C for 24 hours in an oven or in a room, or composting for 120 hours resulted in failure to detect ILTV by PCR. Similarly, ILTV was not detected after addition of commercial litter treatment chemicals (e.g. aluminium sulphate (Al+Clear®)) that reduces ammonia and pH in litter (Giambrone et al. 2008).
	1. The GMO – nature and effect of genetic modifications
		1. The genetic modification
48. The wild type parent strain was originally derived from the virulent strain isolated from a field outbreak of ILT in Glenfield, NSW in 1959 (NSW virulent G strain). The virulent G strain was later renamed CSW-1 after about 10 passages in chicken kidney cell culture. The CSW-1 strain underwent a further three passages in chicken embryo kidney (CEK), then one passage in leghorn chicken liver tumour (LMH) cell line, and another passage in CEK cell line before the genetic modifications were carried out as described below.
49. The GMO is a live attenuated virus with a deletion of the gene encoding glycoprotein G (gG). The gG gene was removed by a series of targeted homologous recombination steps (Figure 3). Initially the gG gene was replaced with the enhanced green fluorescent protein (eGFP) gene, resulting in ∆gG(eGFP) ILTV. This eGFP gene was then removed from the ∆gG(eGFP) ILTV genome to create the GMO (∆gG ILTV) (Devlin et al. 2006).





Figure 3. Construction of the GM virus

1. Wild type ILTV genome with gG gene flanked by upstream and downstream sequences.
2. gG was replaced with eGFP resulting in ∆gG(eGFP) ILTV.
3. eGFP was removed from ∆gG(eGFP) ILTV genome resulting in ∆gG ILTV (the GMO).

IR: internal inverted repeat. TR: terminal inverted repeat. gG: coding region of gG gene. Gu: US2, PK and UL47 genes upstream of gG. Gd: gJ, gD, gI, gE, and US9 genes downstream of gG. eGFP: enhanced green fluorescent protein.

1. The region of the GMO genome flanking the deletion was sequenced. The sequence data indicate that the gG transcription start and termination sequences are intact, and theoretically, transcription of an approximately 150 nucleotide mRNA could occur. As the translation initiation (ATG) codon remains in the transcript, this mRNA could theoretically result in the translation of a 27 amino acid non-functional protein corresponding to non-coding regions of the gG mRNA. The potential expression of the mRNA and protein has not been investigated.
2. Compared with the CSW-1 strain, the GMO also has a two base-pair deletion in the non-coding sequence four base pairs 5' to the initiation codon, and a single A to G transition in the non-coding sequence approximately 700 base pairs 5' to the initiation codon.
	* 1. Glycoprotein G
3. Glycoprotein G is conserved in most members of the *Alphaherpesvirinae* subfamily. Glycoprotein G is secreted or anchored on the plasma membrane of the infected cell (Bryant et al. 2003). The role of gG appears to vary in different alphaherpesviruses. Studies on *Equine herpesvirus* 1 and 4 (EHV-1 and 4) with gG deletion have shown that gG is not essential for virus growth *in vitro* (Huang et al. 2005)*.* *Bovine herpesvirus* 1 (BHV-1) with an inactivated gG exhibited defects in plaque formation and reduced *in vitro* growth (Nakamichi et al. 2001). Furthermore, gG in BHV-1 facilitates viral cell-to-cell spread by maintaining cell-to-cell junctions of infected cells (Nakamichi et al. 2002). For *herpes simplex virus* (HSV), both gG and gC are required for efficient infection of the apical surfaces of corneal epithelial cells *in vitro* (Tran et al. 2000). For some herpes viruses, such as *Feline herpesvirus* 1 (FeHV-1), EHV-1 and BHV-1, gG functions as a virus-encoded chemokine binding protein (vCKBP) that prevents chemokines interacting with their cellular receptors. As a result, an advantage may be conferred to the virus by inhibiting chemokine-mediated inflammatory reactions (Bryant et al. 2003; Costes et al. 2005).
	* 1. Characterisation of the GMO
			1. In vitro studies
4. Inoculation of LMH cells at a multiplicity of infection (MOI) of 0.002 showed no significant difference in the *in vitro* growth kinetics of the GMO and the CSW-1 parent strain, displaying peak titres of about 105 PFU/mL at 4 days post-inoculation. The removal of gG did not affect transcription of the upstream and downstream sequences immediately adjacent to gG. The ability of the virus to spread cell-to-cell as measured in plaque assays was similar to CSW-1 (Devlin et al. 2006).
	* + 1. Virulence
5. Attenuation of the GMO as a result of the gG gene deletion was demonstrated in a study by Devlin et al (2006). At 4 days post-inoculation, chickens inoculated with the GMO showed milder ILT disease symptoms and had greater weight gain compared to those inoculated with CSW-1 ILTV or with ILTV in which the gG gene was reinserted back (∆gG(R) ILTV) (Devlin et al. 2006).
6. At 4 days post-inoculation, chickens inoculated with the GMO had similar titres of the virus of about 104.95 genome equivalents/section isolated from the trachea as those inoculated with CSW-1 ILTV (104.8 genome equivalents/section) or ∆gG(R) (104.5 genome equivalents/section) ILTV, suggesting that the capacity for *in vivo* replication and shedding the virus from the trachea was not affected by the loss of gG (Devlin et al. 2006).
7. Chickens inoculated with the GMO had greater tracheal mucosal thickness than those inoculated with CSW-1 ILTV or ∆gG(R) ILTV. The increase in mucosal thickness is consistent with increased inflammatory cell infiltrate in the mucosa. This suggests that gG may play a role in influencing the inflammatory response at the site of ILTV infection (Devlin et al. 2006).
8. Mortality rates in chickens inoculated with the GMO were lower (2 out of 8) than for CSW-1 ILTV (5 out of 8) or for ∆gG(R) ILTV (6 out of 8) in this study (Devlin et al. 2006). In another study by the same group (Devlin et al. 2007), pathogenicity of the GMO was compared to that of the SA2 and A20 ILTV vaccine strains. In this study, mortality for chickens inoculated with the GMO (2 out of 8) was similar to that for the A20 strain (1 out of 8) at 21 days post-inoculation, and less than for the SA2 strain (7 out of 8 at day 8 post-inoculation, at which time this group was discontinued). Each chicken was inoculated with a similar dose of these strains. Based on these results, deletion of the gG gene results in a less virulent ILTV. It remains to be investigated whether gG functions as a virus-encoded chemokine binding protein.
	* + 1. Immunogenicity
9. A study by Devlin et al (2007) investigated the protective immune response generated by the GMO. Chickens were first vaccinated with the GMO, then 28 days post-vaccination, they were challenged with a CSW-1 ILTV strain. Chickens previously inoculated with the GMO had significantly less severe clinical ILT signs and tracheal histopathology compared to chickens without prior inoculation. After challenge with wild type ILTV, greater weight gains were observed in chickens previously inoculated with the GMO compared to unvaccinated chickens. Using PCR and plaque assay in LMH cells, no ILTV was detected in the trachea of previously vaccinated chickens four days after challenge with CSW-1 ILTV.
10. At 21 days after inoculation with the GMO without subsequent ILTV challenge, antibodies against ILTV were detected by ELISA in chickens inoculated with the GMO. Chickens inoculated with the A20 vaccine had significantly higher antibodies against ILTV compared to those inoculated with the GMO. However, the antibody titres were not considered by the authors to correlate with protection against ILT disease because local cell-mediated immune responses may be responsible for ILT disease protection (Devlin et al. 2007).
11. The protective immunity provided by the GMO was assessed in the laboratory by studying transmission of the CSW-1 ILTV strain to chickens previously vaccinated with the GMO (Devlin et al. 2011). Chickens were first inoculated with the GMO (3000 PFU) 3 weeks before one chicken infected with the CSW-1 strain was introduced into the same cage. After 6 days of exposure, only one out of 30 vaccinated chickens tested positive for CSW-1 ILTV DNA. This suggests that wild type ILTV would not spread among vaccinated chickens and may be cleared by the host immune system.
	* + 1. Efficacy
12. The efficacy of the GMO in protecting chickens after challenge with CSW-1 was compared with other live attenuated vaccine strains, A20, SA2 and Serva (Coppo et al. 2011). Each treatment group, comprising 20 or 21 chickens, was inoculated with the vaccine [dose at 103.48 PFU (GMO), 103.70 PFU (A20), 104.10 PFU (SA2), 102.50 median tissue culture infective dose (Serva)] 21 days prior to challenge with CSW-1 ILTV (103.65 PFU). Five days after challenge, chickens inoculated with the GMO displayed similar clinical ILT signs compared with the A20, SA2 and Serva group.
13. At 6 days after challenge, the chickens were sacrificed. Each chicken was weighed to calculate the weight gain, and tracheal histopathology was examined under the light microscope. The weight gain of chickens inoculated with A20 vaccine was the highest of all the groups, but the weight gains were similar between the SA2, Serva and the GMO. Tracheal histopathology was similar between the different vaccine groups (Coppo et al. 2011).
	* + 1. Transmission
14. To study transmission of the GMO, chickens that had been inoculated with either the GMO or CSW-1 ILTV (4500 PFU) 4 days earlier were introduced into cages with naïve chickens (3 replicates of 10 naïve chickens for each ILTV strain) for 6 days. Across the three replicates, 8 of the 30 in-contact chickens became infected with the GMO, while 4 in-contact chickens became infected with the CSW-1 ILTV strain. In one replicate in each experimental group no transmission was observed, and transmission of the GMO and CSW-1 were not found to be statistically significantly different (Devlin et al. 2011). Transmission of the GMO to susceptible bird species other than chickens has not been studied.
	* + 1. Phenotype stability
15. As a measure of the stability of the GMO’s attenuated phenotype, weight was measured in naïve chickens before they were housed in the same cages as the GMO-inoculated chickens and at the end of the transmission study (Devlin et al. 2011). Weight gain of naïve, in-contact chickens that became infected with the GMO was similar to those of naïve, in-contact chickens that did not become infected. This suggests that after one *in vivo* passage in chickens, the GMO remained attenuated.
	* + 1. In vivo stability
16. A study by Coppo et al (2011) took tracheal swabs from chickens inoculated with the GMO, A20, SA2 and Serva at 21 days post-inoculation to examine viral DNA presence using qPCR. GMO DNA was detected in 2 out of 21 chickens inoculated with the GMO at 21 days, with a mean of 2.17 log10 viral DNA copies in the trachea. Chickens inoculated with the A20 vaccine (9 out 19) had the lowest mean viral DNA copies (1.98 log10). Chickens inoculated with the SA2 vaccine (15 out 19) had the highest mean viral DNA copies (2.95 log10), followed by the Serva vaccine (2.46 log10 in 9 out of 20 chickens). The results show that the commercial vaccine strains persisted longer in the trachea than the GMO. However, it is unclear if the amount of ILTV DNA detected represents infectious virus particles.
	1. Receiving environment
17. The receiving environment forms part of the context for assessing risks associated with dealings with GMOs. It informs the consideration of potential exposure pathways, including the likelihood of the GMOs spreading or persisting outside the site of release. Relevant information about the receiving environment include state legislation and local council requirements relevant to poultry farming, the current commercial broiler farming and processing practices, biosecurity standards for poultry farms, waste management practices, site of release, the related viral species in the environment and potential hosts in the environment.
	* 1. Background on broiler farming
18. NSW and Victorian broiler farms, including free range farms, must comply with a range of legislation designed to protect people and the environment. Local councils and/or state government agencies must approve intensive agriculture developments including free range broiler farms. Local councils are generally the responsible authority for the administration or enforcement of planning schemes. This means that councils would assess and determine farm planning permit applications. Councils are also responsible for monitoring and enforcing the compliance of broiler farm operators with their planning permit conditions. In addition, the poultry company may require minimum distances between the broiler farm and other poultry farms or livestock farms owned or managed by them or by others.
19. Boundary setbacks may be required by councils and are defined as the distance between the nearest external edge of any new broiler chicken shed or litter stockpile or compost pile and the farm boundary. Boundary setbacks mitigate visual amenity issues and the immediate impact of odours, dust, aerosols and noise emissions from sheds, litter, or compost piles on the amenity of adjacent land and the surrounding area.
20. The separation distance is the distance from the nearest external edge of a broiler shed to the nearest external edge of a sensitive use (e.g. house or public building) on land beyond the broiler farm property. It excludes sensitive uses directly associated with the broiler farm operations – e.g. residential dwellings on the broiler farm property. Separation distances are used to reduce the effects of odour, dust, aerosols and noise. Separation distances usually extend across adjoining properties that are not owned by the farm owner. The greater the separation distance and the boundary setback, the lower the probability of offensive odour and dust adversely impacting the surrounding community.
21. A buffer is where the farm owner has legal control of the land needed to separate the poultry sheds from adjoining developments. A buffer may be open farmland, or a landscape area that hides views of the sheds or helps to disperse odours.
	* + 1. NSW requirements
22. The *Environment Planning and Assessment Act 1979* (NSW) (EP&A Act) is the major legislation governing land use and environmental assessments. The EP&A Act establishes a framework for local government zoning, assessment requirements, development control plans and development consent provisions. In addition, broiler farms in NSW that accommodate more than 250,000 chickens require a licence under the *Protection of the Environment Operations Act 1997* (NSW) (POEO Act).
23. The *Best Practice Management for Meat Chicken Production Manual* (the Manual) (NSW Department of Primary Industries 2012) provides guidance for the planning, design, construction and management of shed based broiler farms in NSW, but not for free range farms. The Manual recommends that new poultry farms be a minimum of 1000 metres from other intensive poultry farms (500 metres when there are extenuating circumstances such as farms with a common owner or farms supplying the same processor); 3000 metres to commercial duck farms; and 5000 metres to poultry breeder farms. In addition, the Manual recommends that new farms be away from waterways and wetlands (ideally 3000 metres) that are used extensively by waterfowl.
24. In relation to protecting ground water or watercourses, the Manual recommends locating the broiler farmland above the 1-in-100-year flood line; avoiding locating the farm near major potable water supply storages and watercourses within drinking water catchments; and protecting riparian zones with appropriate buffer zones and vegetative filter strips. The Sydney Catchment Authority specifically requires that broiler farms not be located within 100 metres of a major potable water supply or reservoir, or within 40 metres of a watercourse in the Sydney drinking water catchment.
25. However, as the Manual only relates to new poultry farms built since 2012, older broiler farms could be sited in close proximity to other poultry farms or residential areas.
	* + 1. Victorian requirements
26. In Victoria, broiler farms must be approved by local councils and the Victorian Department of Environment, Land, Water and Planning under the *Planning and Environment Act 1987*. In all Victorian planning schemes, a planning permit is required to use and develop land for a broiler farm including free range broiler farms. Broiler farms, including free range broiler farms, are prohibited in all urban zones, Rural Conservation Zone, the Green Wedge A Zone and Rural Living Zone.
27. Compliance with the *Victorian Code for Broiler Farms 2009* (the Broiler Code) (Victoria Department of Economic Development 2009) is mandatory for the establishment of all new broiler farms and expansion of the capacity of existing broiler farms, but does not apply to free range farms. Broiler farms that were lawfully established before the introduction of the Broiler Code may continue to operate in conformity with their previous lawful operations and the conditions of any valid planning permit that pertains to the broiler farm. Where the Broiler Code does not apply, it may still be a useful reference for identifying relevant issues and responses to inform the preparation and consideration of a proposal.
28. The Broiler Code details requirements including the location, siting, design, site access, waste management, farm operation and management. In addition, broiler farms must meet the requirements of relevant state and local government regulations.
29. The Broiler Code uses a mathematical formula to calculate the required minimum separation distance, based on the proposed farm capacity for housing chickens. For example, a farm with a capacity of 100,000 chickens requires a minimum separation distance of 325 metres, and a farm with a 400,000 capacity requires a minimum separation distance of 686 metres.
30. The boundary setback is specified in the Broiler Code as at least 100 metres. The Broiler Code has several other requirements for distances of the farm to certain other areas depending on the zoning of the land.
31. The Victorian Department of Economic Development, Jobs, Transport and Resources has provided some guidelines on biosecurity buffer distances for different types of farms. For example, the buffer distance for units in large farm complexes range from 200-500 m, and for new farms, the buffer distance is 1000 m (Victoria Department of Economic Development 2015). As noted above, older broiler farms built before 2009 could be sited in close proximity to other poultry farms or residential areas.
	* + 1. Corporate structures
32. The chicken meat industry is predominantly vertically integrated. This means that generally, individual companies own almost all aspects of production - breeding farms, multiplication farms, hatcheries, feed mills, some broiler farms, and processing plants. Two large integrated national companies supply more than 70% of Australia’s broiler chickens - Baiada and Inghams Enterprises. Inghams and Baiada are privately owned, with farming and processing operations in most states. The rest are medium-sized, privately owned companies, and a myriad of smaller processors.
33. Growing broiler chickens, from day old chicks to the day of processing, is generally contracted out by processing companies to contract growers. Approximately 800 growers produce about 80% of Australia’s broiler chickens under these contracts. Other broiler chickens are produced on large company farms, or on farms owned and managed by ‘intermediary’ companies which own a number of farms, each managed by a farm manager, and who enter into contracts with processing companies to grow out chickens on a larger scale.
34. Contract growers own the farm and provide the management, shedding, equipment, labour, bedding and other inputs to rear chickens. The processing company provides (and owns) the chickens and provides feed, medication and technical advice (Australian Chicken Meat Federation Inc. 2013b).
35. Farms proposed to be included in the trial would be those controlled, owned or contracted by the major commercial poultry processors. The applicant stated that each company operates according to quality management systems incorporating standards such as GMP and the Hazard Analysis of Critical Control Points (HACCP), and in accordance with strict state environmental codes. The companies are members of peak organisations such as the Australian Chicken Meat Federation (ACMF). The applicant stated that they demonstrate a commitment to implementing standards and guidelines such as the National Farm Biosecurity Manual for Chicken Growers (Australian Chicken Meat Federation Inc. 2010), which are generally included within the company quality assurance program.
	* + 1. Broiler chicken farm (including free range farm) routine management in Australia

Shed housing

1. Broiler chickens are farmed in large open poultry houses, usually referred to as ‘sheds’, ‘houses’, ‘barns’ or ‘units’. Shed sizes vary, but a typical shed is about 150 metres long and 15 metres wide and holds about 40,000 adult chickens. The larger sheds can contain up to 60,000 chickens. There are often three to ten sheds or more on one farm. A typical new farm would house approximately 320,000 chickens, with eight sheds holding approximately 40,000 chickens each.
2. Traditionally, broiler sheds have been ‘naturally ventilated’, with the sides of the shed open to fresh air. The amount of air circulating through the shed is changed by raising/lowering curtains running along the side of the shed, or by a vent opening at the top of the shed. Fans are sometimes used to encourage air flow, and water misting systems cool the chickens by evaporative cooling in very hot conditions.
3. An increasing number of chicken sheds have ‘tunnel ventilation’. Tunnel ventilation sheds have fans at one end of the shed which draw air into the shed through cooling pads in the walls, over the chickens and out the far end of the shed at high speed. Three or four temperature sensors in the shed allow automatic control of the fan, heating and cooling settings.
4. Feed lines and pans run the length of the shed and are supplied automatically from silos outside the shed via pipes. Feed silos are kept secure against all pests, and any spillage around silos is cleaned up immediately to prevent attraction of pests. Water lines run the length of the shed, with drinkers at regular intervals. Water and feed are placed so that chickens are never more than about 2 metres from food and water. The water and feed lines can be raised or lowered within the shed to allow feeding, or for pick-up (harvest) or shed clean-out (Australian Chicken Meat Federation Inc. 2013a).
5. Bedding used is a thick layer of litter, such as sawdust, wood shavings, rice hulls or other materials spread across the floor.
6. For shed floors, the Broiler Code has requirements stipulating: that the base of the broiler shed should be constructed from low-permeability materials such as concrete, compacted clay or another sealed surface; the finished floor level of the broiler shed should be above the natural surface level to prevent the entry of stormwater run-off, or alternatively, the shed should be bunded or a surface drainage system installed to prevent the entry of stormwater run-off; and a concrete stand area should be located at the entrance to each broiler shed.
7. A number of structures are regularly inspected and maintained including shed walls, roofs, ventilation, cooling systems, automated environmental controllers, sensors, water reticulation systems, silos and feed-lines. The surrounding area must be maintained to ensure they are clean and tidy (NSW Department of Primary Industries 2012).

Chicken rearing

1. Day-old chicks are transported from the hatchery to broiler farms, usually in ventilated chick boxes in specially designed, temperature controlled trucks. On arrival at the broiler farm, chicks are placed onto the floor of the shed, where they are initially confined to about a half or one-third of the total shed area (the ‘brooding area’) and given supplementary heating from gas heaters or heat lamps.
2. For the first two days of the flock’s life, the shed temperature is held at 31 - 32°C, the optimum temperature for chick comfort, health and survival. As the chickens grow, the shed temperature is gradually lowered by about 0.5°C each day, until it reaches 21 - 23°C at 21 days. The farmer aims to maintain shed temperatures within this range, although towards the end of grow-out period for large chickens, the temperature may be reduced.
3. As the chickens grow, the area available to them is increased until they have free run over the floor of the entire shed.
4. Generally, feed and clean water is available 24 hours a day, although some operators make feed available at specific 'meal times' only.
5. Farm workers regularly, at least once every day, monitor the flock’s health and progress, remove any dead chickens, and cull any sick or injured ones. Farm workers also check feeders and water systems. Careful management of ventilation and water system helps keep the litter clean and dry, as poor litter affects air quality and can affect bird health and performance.
6. Over the life of the broiler flock about 4% of chickens die as a result of natural causes or selective culling (Australian Chicken Meat Federation Inc. 2013a).

Pick-up or harvest

1. In Australia, a percentage of chickens are harvested from most flocks on several occasions. Harvesting, also known as ‘partial depopulation’, ‘thinning out’, or ‘multiple pick-up’, may be done up to four times until all chickens have been removed from the shed. Thinning out sheds allows more space for the remaining chickens and reduces the natural temperatures in the shed. The first harvest might occur as early as 28 days and the last at 55-60 days of age.
2. Immediately before pick-up the sheds are cleared of all dead chickens and any chickens not suitable for catching. The FSANZ Standard for Poultry Meat (Standard 4.2.2) requires that diseased poultry must not be sold or supplied for human consumption (FSANZ 2010). Feedlines are lifted not more than 3-6 hours before pick-up in accordance with the instructions given by the processor. Access to water is not removed until the pick-up crew arrives on the farm.
3. Chickens are often harvested at night as it is cooler and the chickens are more settled. They are generally picked up by specialised contracted pick-up crews under low lighting conditions so that they are calm and easy to handle. They are usually caught by hand and placed into plastic crates or aluminium modules designed for good ventilation and safety from bruising during transport. These crates or modules are handled by specialist forklift equipment and loaded onto trucks for transport to the processing plant. During pick-up, the farmer is available to help maintain all aspects of chicken welfare.
4. When all the chickens have been removed from the shed (after about 60 days), it is cleaned and prepared for the next batch of day-old chicks.

Shed clean-out

1. The next batch of chicks generally arrives in five days to two weeks, giving time to clean the shed and prepare for the next flock. The break also reduces the risk of common microorganisms being passed between batches as many pathogens die off. As each broiler flock spends 6 to 7 weeks in a shed and there is a two week break between batches, farmers run about 5.5 batches through a shed each year.
2. Some farms undertake a full cleanout after every batch. This includes removing bedding, brushing floors, scrubbing feed pans, cleaning out water lines, scrubbing fan blades and other equipment, and checking rodent stations. High pressure hoses clean the whole shed thoroughly at a standard rate of 6000 to 8000 litres of water per shed. Because low water volumes are used, there is little water run-off. The shed is disinfected, using low volumes of disinfectant which is sprayed throughout.
3. On other farms, a partial clean-up of the shed is done, including removing old litter and/or topping up with fresh litter and cleaning and sanitising all equipment. A full cleanout is done after every second or third batch of chickens (Australian Chicken Meat Federation Inc. 2013a).
4. The darkling beetles live in the litter, predominantly under feed pans. Earth floors of broiler houses are an important medium for pupation (Poultry Hub 2017). An insecticidal treatment may be applied in areas where shed insects such as beetles are a problem. However, some strains of darkling beetles are resistant to certain insecticides (Hamm et al. 2006).
5. Registered veterinarians or technicians may test the sheds after a full cleanout to confirm sheds have been adequately cleaned and potential disease agents removed.

Free range farms

1. Free range broiler chickens are produced using similar management, housing, rearing and feeding practices as conventional broiler chickens. Free range broiler chickens are harvested in the same timeframe as shed-based chickens. The major differences are that free range broiler chickens are allowed access to an outside run for part of each day (at least after the brooding period) and often have lower target stocking densities. Depending on the accreditation program adhered to, use of antibiotics to treat sick birds may preclude the meat from these chickens being sold as free range (Australian Chicken Meat Federation Inc. 2013a).
	* + 1. Water use
2. Broiler chicken farms must comply with the relevant state legislation to prevent contamination of surface and ground water, watercourses or bores and catching overland overflow.
3. Broiler chicken sheds operate as closed systems with little or no water escaping to the outside environment. Any water spilt inside the shed from drinking equipment or during cleaning would subsequently evaporate.
4. The risk of ground water contamination is primarily avoided via appropriate site selection and by engineered construction and compaction of the shed floor (NSW Department of Primary Industries 2012).
	* + 1. Environmental monitoring
5. Environmental monitoring and recording form part of farm management to ensure that the requirements of relevant state legislation are met. Growers are encouraged to develop, document and implement an Environmental Management Plan for the farm (NSW Department of Primary Industries 2012).
6. During high-risk activities (such as shed clean-out), a record is kept of management actions to minimise the risks. Records must be made available to relevant regulatory authorities.
	* + 1. Transport of live chickens to processing plant
7. Chicken farms are generally within 100 kilometres of the processing plant. Poultry processing plants are usually close to markets and labour sources, with many of the largest operations within 50 km of a capital city.
8. When chickens reach a suitable size for market (typically around 28 days of age), chickens are placed in crates for transport in an open truck and transported in accordance with the relevant state legislation. Crates, trucks, equipment and other materials used to transport the chickens from the shed to the processing plants are decontaminated with disinfectant after delivery of chickens. The Standard for Poultry Meat (Standard 4.2.2) requires that transportation vehicles and equipment be effectively cleaned, sanitised and in good repair to ensure poultry is not made unsafe or unsuitable for human consumption (FSANZ 2010).
9. In Victoria, live chickens taken from farms are transported in accordance with the *Australian Animal Welfare Land Transport of Livestock Standards and Guidelines*, referred to as the Land Transport Standards (LTS) (Animal Health Australia & Department of Agriculture 2012). The LTS has been adopted into the Victorian legislation under the *Livestock Management Act 2010* (Victoria).
10. In NSW, live chickens are transported in accordance with the *Prevention of Cruelty to Animals (Land Transport of Livestock) Standards 2013* which is required under the *Prevention of Cruelty to Animals Act 1979* (NSW). The provisions in the NSW Standards reflect those in the LTS.
11. The LTS has specific requirements for transporting poultry aimed at animal welfare, such as that container or crate openings must be 20 cm x 22 cm, containers must be stacked in a way that facilitates airflow, maintaining appropriate temperatures, preventing delays in transporting and unloading, and protecting poultry from various weather conditions.
	* + 1. Poultry processing and rendering plants
12. The processing plants slaughter, process and package chickens for wholesale or retail sale for human or animal consumption.
13. The processing plants are highly automated and adhere to high standards of cleanliness and hygiene. Meat processing plants must have documented procedures including those related to sanitation to ensure the safety of food. Meat processing plants are regulated by the state regulatory authorities.
14. In Victoria, some functions of PrimeSafe (the regulatory authority), as prescribed under the *Meat Industry Act 1993* (Victoria), include licensing meat processing facilities, reviewing the standards of meat produced for consumption or sale within the state, and reviewing the standards of the construction and hygiene of plant and equipment in a meat processing facility.
15. Likewise, meat processing plants in NSW must have a licence from the NSW Food Authority, which inspects the premises to ensure all buildings and equipment meet the relevant standards and requirements of the *Food Act 2003* (NSW). In addition, meat processing plants may require a licence from the NSW Environment Protection Authority (EPA).
16. As well as regular flush and ‘spot’ cleaning of the plant during a shift, a full daily cleaning also occurs at the processing plant. All sections of the processing plant, including the live bird area and the wastewater pits and pipes, are cleaned and flushed daily. All internal factory areas and contaminated external areas usually drain to wastewater pits and then to the effluent treatment and disposal system.
17. At some poultry processing plants, wastewater may be directed to a compact effluent treatment system such as a dissolved air floatation unit to remove grease and solids before it is discharged to the sewer in accordance with trade waste agreement. Solid waste removed from the effluent by the dissolved air floatation system may be, for example, transported daily to the local landfill to minimise odours.
18. Waste products from the processing of birds at the processing plant are collected from the processing line and separated into three distinct waste streams comprising blood, feathers and internal organs/heads/feet (offal) for transport to a rendering plant. Rendering plants that process substances for human consumption are required to apply for a licence from the state regulatory authorities (e.g. PrimeSafe, NSW Food Authority), meet relevant standards such as the *Australian Standard* 5008 *- Hygienic Rendering of Animal Products* (Standards Australia 2007), comply with legislation such as *Food Act 2003* (NSW), and be inspected by the state regulatory authorities.
	* 1. Biosecurity
			1. Biosecurity legislation
19. Each state and territory has their own biosecurity legislation. The *Biosecurity Act 2015* (NSW) provides tools and powers to manage animal and plant pests and diseases, weeds and contaminants that threaten the NSW economy, environment and community. The tools allow for practical responses proportionate to risk, and include: emergency powers in case of a significant biosecurity risks, as well as requiring people who deal with biosecurity, and who have knowledge of the biosecurity risks posed, to take reasonable steps to manage those risks. The Biosecurity Act includes strong enforcement tools, including significant penalty provisions especially for wilful or reckless acts. The *Livestock Disease Control Act 1994* (Victoria) provides for the prevention, monitoring and control of livestock diseases in Victoria, and also addresses issues related to licences, registrations and enforcement.
	* + 1. Poultry farm biosecurity standards
20. As part of current arrangements between the Victorian government and industry, poultry producers are expected to implement on-farm biosecurity programs and follow them on a daily basis to reduce the risk of transmission of disease onto and between poultry farms (Victoria Department of Economic Development 2015).
21. A number of documents provide guidelines including *National Farm Biosecurity Manual for Poultry Production* (Department of Agriculture 2009), *National Farm Biosecurity Manual for Chicken Growers* (Australian Chicken Meat Federation Inc. 2010) and *NSW Biosecurity Guidelines for Free Range Poultry Farms* (NSW Department of Primary Industries 2007). These outline biosecurity standards applicable to all poultry producers including free range farms.

Farm facilities – conventional and free range

1. The biosecurity standards applicable to both conventional and free range broiler farms are summarised below.
2. Each farm must keep a copy of the *National Farm Biosecurity Manual for Poultry Production* (Department of Agriculture 2009) or a more detailed document that encompasses the manual such that it is readily accessible to workers. All workers must be trained in the relevant parts of the manual and such training is to be recorded.
3. The production area (sheds or free range area, feed storage and handling area, and area immediately surrounding the sheds including pick-up areas) must have a perimeter fence or otherwise well-defined boundary (e.g. creek, vegetation) establishing a clearly defined biosecurity zone.
4. If livestock graze on the property then the production area must have a stock proof fence. Grazing near sheds (i.e. on part of the production area) is only permitted where the grazing area is separated by a stock proof barrier from the area used by poultry.
5. The main entrance to the production area must be capable of being closed to vehicle traffic (e.g. lockable gate which should be kept locked at all times) and must display appropriate signage including ‘Biosecure Area No Entry Unless Authorised’ or similar wording. In addition, signage including contact numbers must direct visitors to contact the producer before proceeding.
6. There must be a change area away from sheds with clean protective clothing and boots provided. Entry to sheds must only be made through entrances with a footbath containing a suitable disinfectant. There must be provision for scraping the soles of boots before dipping to ensure the disinfectant makes contact with the soles of the boots. An alternative system using separate production area- and shed-footwear may be used. Facilities for hand sanitation must also be placed at the entry to each shed.
7. Facilities should be available for the cleaning and disinfection of equipment before entry.
8. Feeding systems must wherever possible be closed to ensure that feed in silos and feed delivery systems are protected from access and contamination by wild birds and rodents. Feed spills should be cleaned up without delay to prevent the congregation of wild birds.
9. Drinking water should be accessed inside the shed; or, if watering stations are required outside, they should be of a type that cannot be easily accessed by wild birds (e.g. a nipple system). The watering system should be maintained, in order to prevent leakage and the creation of wet patches within or outside the shed. Water tanks should be checked regularly to ensure that they remain bird-proof.
10. Drinking water for poultry, as well as cooling water used in poultry sheds, must meet appropriate water standards. Water that does not meet the standard must be treated (e.g. chlorination, ultraviolet, iodine) to ensure that the standard is met. All surface water (dam, river etc.) must be treated before being used as drinking water for poultry. Treated water supply must be kept in a closed system from the point of treatment to the drinker.
11. All poultry housing must be designed and maintained so as to prevent the entry of wild birds and limit the access of vermin as far as is practical.
12. The production area should be adequately drained to prevent accumulation and stagnation of water likely to attract water fowl, especially in the areas around sheds.
13. Trees and shrubs should be selected to minimise wild bird attraction. The area around sheds must be kept free from debris and vegetation should be mown regularly. Vegetation buffers for environmental compliance should not be compromised.
14. An appropriate vermin control plan must be developed and implemented, including rodents, foxes, wild dogs and cats. A baiting program for rodents must be implemented where a risk assessment deems this necessary (e.g. live rodents, droppings, nests).
15. Beetle populations within shed litter should be controlled via an integrated pest management approach by using pesticides, composting and total shed and litter clean-out.
16. Only commercially produced avian species are to be kept in the production area and no other avian species (including aviary birds and pet birds) or pigs are to be kept on the property.
17. If more than one commercially produced avian species is kept in the production area, the species should be housed and managed separately, with suitable biosecurity arrangements for each species. Shared equipment should be cleaned and disinfected between uses.
18. Used litter and manure must not be stockpiled in the production area. Used litter and manure must be stored in an appropriately designed storage area away from the production area.
19. Dead bird disposal methods must conform with applicable environmental compliance requirements.

Free-range farms

1. The following biosecurity measures are specific for free-range farms.
2. Good fencing is required to prevent the entry of animals such as dogs, foxes and cats. In many situations, however, fencing alone is insufficient to stop such intrusions; therefore, some free range enterprises keep specially trained dogs with the chickens, as protection against other animals and against unauthorised human entry. Dogs must not enter sheds unless part of the flock security strategy. Guard dogs such as these are not regarded as a biosecurity risk but rather as a biosecurity tool.
3. Where footbaths are not appropriate for a free range paddock, a system should be documented and implemented to monitor and prevent any potential hazardous organic material or litter entering free range paddocks.
4. In free range farms, chickens may have some exposure to wild birds. Therefore, documented measures must be taken to minimise the congregation of waterfowl and the impact of wild birds. Wild bird attractiveness can be minimised by placing feeders and water inside the shed, rather than in the open range where wild birds would have easier access. Placement of bird netting in critical feeding areas may also reduce the risk.
5. In free range farms with sheds or other housing, manure deposits outside the hatch openings must be removed after each batch, and ramps used by chickens must be scraped and cleaned after each batch.
6. Grass on and around the farm must be kept cut to reduce rodent attraction.

Farm worker standards and visitors – conventional and free range

1. Production area personnel or any person residing on the property must not have contact with any other poultry, avian species or pigs unless they have a complete head-to-toe shower and change into new protective footwear and clothing prior to entering the production area.
2. Personnel must wear laundered clean clothes each day to work and ensure that they do not become contaminated by contact with avian species or pigs on their way to work. It is critical that boots worn in sheds are not worn or taken outside the production area.
3. Company service personnel visiting the production area must wear protective clothing and footwear, as approved by the production facility manager. Hands must be sanitised before entering sheds.
4. Contractors who have had contact with poultry or other birds that day or keep birds at their home must not enter sheds and/or ranges populated or ready to be populated with birds unless it is an emergency, and they have showered from head-to-toe, changed clothes and boots and wear hair covering. Tools taken into the production area must be cleaned before entry into sheds and must be free of dust and organic matter.
5. All persons must agree to comply with the entry conditions by signing the visitors’ log and such visits must be approved by the manager before visitors may enter sheds and ranges. This requirement also applies to vaccination crews.
6. Pick-up crews work from youngest to oldest or all young birds or all old birds on a shift basis in accordance with the processing company’s pick-up biosecurity procedures. Pick-up crews must not keep birds at their homes. Drivers must sanitise their hands and boots before and after each pick-up or delivery to a production area. Trucks carrying unused or used litter must be cleaned and disinfected between production areas.
7. A system for tracing movements of delivery personnel (e.g. through delivery dockets and feed company records) must be implemented.
	* + 1. High level biosecurity
8. In the event of an outbreak of disease, the *National Farm Biosecurity Manual for Poultry Production* (Department of Agriculture 2009) recommends the following measures:
* limiting visitors from entering the production area unless absolutely essential
* visitors who visit must have a head to toe shower before and after visit
* used clothing and personal protective equipment must remain on property
* any vehicle entering the property must be washed and disinfected before and after going onto the property
* poultry and litter must not be moved on or off property until disease status is clarified.
1. Farms require a contingency plan to cope with occurrences of high mortalities. An investigation must be conducted to ascertain the cause of death and the best option for the disposal of the dead birds. Where normal disposal methods are not feasible, the relevant regulatory authorities (e.g. the local council, the state EPA) may need to be contacted to help identify alternative options.
2. Subject to approval from local council, state EPA and other authorities, mass-death disposal options may include: rendering (if facilities are available), in-shed composting, external composting, disposal in a landfill site, or burial on-farm (see Section 7.3).
3. If the cause of the death is an Emergency Animal Disease, then the relevant Australian Veterinary Emergency Plan (Ausvetplan) would be activated and the appropriate authorities would be notified. Disposal of carcasses, used litter and feed, and decontamination of equipment, would be under the direct control of the state’s Chief Veterinary Officer.
4. The *Biosecurity Incident Management System* (Biosecurity Emergency Preparedness Working Group 2012) provides guidance for the management of biosecurity incident response in Australia and can be applied to all biosecurity sectors. Typically the states and territories have primary responsibility for preparing and responding to biosecurity incidents within their borders. The DAWR has a role in providing national leadership and coordination in preparing for and responding to biosecurity incidents.
	* 1. Waste management
5. In NSW, the handling of litter and waste on the farm must meet the requirements of waste management legislation such as the POEO Actand may require a permit from the NSW EPA. For farms in the Sydney drinking water catchment area, the Sydney Catchment Authority does not permit the disposal of chicken carcasses on site, except during an outbreak of exotic disease that results in a farm being quarantined.
6. In Victoria, the state government, catchment management authorities and local councils each have roles and responsibilities that relate, directly or indirectly, to farm waste management. Management of waste including used litter and dead chickens must be conducted in accordance with the conditions of the planning permit. For example, farms may not stockpile, compost or spread litter if the planning permit conditions require removal of all litter directly off-farm.
7. Incineration is not the preferred practice because it is expensive and must be conducted only in authorised incinerators built for purpose. Burning carcasses in open fires is not permitted.
8. The *Best Practice Management for Meat Chicken Production Manual (NSW Department of Primary Industries 2012)* and the *Victorian Code for Broiler Farms 2009* (Victoria Department of Economic Development 2009) stipulate the following relevant measures for disposal of litter and dead chickens:
* litter must be removed from the farm or operational area immediately as sheds are being cleaned out and transported from the farm in covered vehicles to avoid spillage and dust emissions
* chicken carcasses must be removed from the sheds daily and disposed of, or stored appropriately (e.g. in freezers), within 24 hours of death
* for collection of chicken carcasses by waste contractors, the collection point for carcasses must be as far as practical away from the farm site to prevent the collection vehicle from entering the site, collection point must be weather-proof and easily cleanable, and the collection vehicles and containment systems must be leak-proof and vermin-proof
* where birds need to be frozen before collection, secured freezers with adequate space must be provided. Carcass-storage containers and the collection area must also be regularly cleaned and disinfected to minimise the spread of disease
* any spillage in the collection areas must be immediately cleaned and disinfected
* records of collection (date and mass) must be maintained
* personnel disposing of carcasses should be instructed on maintaining personal hygiene and environmental protection measures
* carcasses (or bird bins) must not be left in public view and must be disposed of at licensed composting facilities, rendering plants or landfills.
	+ - 1. Composting
1. Composting facilities and those on farm land must comply with the state requirements. Under the EP&A Act (NSW), approval for composting sites and on farm land may be required from an appropriate authority (usually the local council) or state EPA. Composting facilities must comply with legislation including the *Environment Protection Act 1970* (Victoria).
2. The Environmental Guidelines on *Composting and Related Organics Processing Facilities* (NSW Department of Environment and Conservation 2004) focus on the appropriate environmental management of commercial composting facilities in NSW. Composting in Victoria must be conducted in accordance with *Designing, Constructing and Operating Composting Facilities* (Victoria Environment Protection Authority 2015). Both the NSW and Victorian composting guidelines refer to the AS 4454 – *Composts, Soil Conditioners and Mulches* (Standards Australia 2012).
3. The composting guidelines from each state cover topics related to distances of compost area from other sensitive areas including wildlife parks and water areas, vermin control, management of compost and dispersal of debris. Compost containing different types of organic matter are categorised depending on the risk of harm to human health and/or the environment. Animal carcasses are placed in the highest risk category. The requirements for these high risk organics are more stringent in order to minimise their impact on various environmental aspects. The guidelines recommend that high risk organics be composted in an enclosed environment with a high level of secondary odour controls.
4. The *Best Practice Management for Meat Chicken Production Manual (NSW Department of Primary Industries 2012)* and the *Victorian Code for Broiler Farms 2009* (Victoria Department of Economic Development 2009) stipulate the following relevant measures for composting on farm land:
* if litter is stored on farm (or composted) it must be managed to avoid contamination of surface waters, stormwater drains, waterways, catchments and ground waters, and avoid excessive fly breeding
* bunding may be required to prevent entry and contamination of stormwater run-off
* temporary litter stockpiles or compost piles must be separated by at least 100 metres from any broiler shed on the subject land, or sited and managed as otherwise stipulated by the processor to meet biosecurity requirements
* if re-using litter on farm, the litter application site must not be on land subject to flooding, steep slopes, rocky, slaking or highly erodible land or highly impermeable soils where there is any risk of nutrient run-off to waterways, groundwater and surrounding land
* nutrient-rich run-off from temporary litter stockpiles or compost piles must be collected in a sump or dam and may be re-used to add moisture to the pile.

Composting process

1. Composting is the microbiological transformation of organic materials under controlled aerobic conditions. There are two phases to the composting process which have different processing parameters. First is **pasteurisation,** which generates heat within the material to significantly reduce the number of viable pathogens and plant propagules. This is followed by **maturation,** which sees the decline in temperatures and moisture levels, slowing of microbial activity and an increase in biological stability of the organic material (Victoria Environment Protection Authority 2015).
2. Current composting practices use either purpose-built compost bins, which may be rotary, or composting bays or piles. Small volumes can be composted in bins. The size and number of compost rotary units required depend on the size of the operation and normal levels of bird mortality (3% to 5%). Rotary units require careful management to ensure that an aerobic environment is maintained in order to reduce the possibility of excessive odour generation.
3. Litter makes up the greatest part of the composted waste by volume. The litter and chicken carcasses from several sheds may be incorporated into a long windrow in open land, along with additional organic material such as sawdust/mill waste or green waste. The litter and carcasses must be covered with at least 300 mm of clean co-compost material to exclude flies or birds. The windrows may be 100 metres long, 3-4 metres in width and 2-3 metres high.
4. Intermediate volumes may be composted in piles or in bays formed from hay bales. Fencing is used to exclude stock that may disrupt the windrows or piles. Compost in large bays, piles or windrows need to be mixed or turned using earth moving equipment.
5. The temperature to effectively achieve pasteurisation ranges from 55°C to 75°C. The temperature reached by composting material influences the rate of decomposition, oxygen demand and microbial population. Other pasteurisation parameters include carbon to nitrogen ratio, total moisture levels, oxygen content, pH, porosity and bulk density.
6. The Victorian composting guidelines (Victoria Environment Protection Authority 2015) and the AS 4454 – *Composts, Soil Conditioners and Mulches* (Standards Australia 2012) require that for compost containing high risk materials (e.g. manure, carcasses), the core temperature must be maintained at 55˚C or higher for a period of 15 days or longer; and during this period of high temperature, the whole compost mass must be turned a minimum of five times.
7. After the completion of the composting process, the compost may later be sold and used in pastures to increase nutrients and add organic matter.
	* + 1. Burial on the farm
8. Burial was a traditional and economical option for disposal of carcasses. However, not all soil types or locations are suitable for on-site burial; for instance, areas may have a high risk of water table contamination or shallow soils. Disposal of chicken carcasses via burial is also unlikely to be suitable in more closely settled areas and on smaller properties, owing to the higher risk of odour or of predation by domestic animals.
9. On-site burial of dead chickens on the farm is currently undertaken only in an emergency situation or with the approval of the relevant authorities such as the state EPA or the Chief Veterinary Officer.
10. The state EPA recommends burial of carcasses on the farm to be undertaken at an area at least 100-300 m away from houses and water sources (e.g. ground water, surface water), that has good access to the site for earthmoving machinery and stock transport unless the stock are to be walked in for slaughter, has a pit base with at least 1-2 m above the level of the watertable, and covered with at least 2 m of heavy soil of low permeability and good stability (NSW Environment Protection Authority 2013; Victoria Department of Economic Development 2016).
	* + 1. Landfill
11. Contaminated farm waste including dead chickens may be disposed of at landfill sites.
12. Landfills may require a licence from the relevant authorities such as the state EPA to dispose of dead chickens from the poultry industry and must be done according to the requirements of the state EPA. The Victorian EPA *Landfill Licensing* guideline (Victoria Environment Protection Authority 2016) and the NSW EPA *Environmental Guidelines – Solid Waste Landfills* (NSW Environment Protection Authority 2016) have requirements in relation to managing waste in landfills to minimise environmental impact, such as segregating active landfill sites from surface water or groundwater, controlling debris, covering waste (such as carcasses) daily with soil or other material at least 0.15 - 0.30 metres thick, and controlling vermin in landfills.
	* + 1. Rendering plants
13. The *Best Practice Management for Meat Chicken Production Manual* recommends removal of chicken carcasses for rendering if a rendering plant is located close to the farm (typically within 100 kilometres).
14. Rendering plants may require a licence from the state EPA to dispose of dead birds from the poultry industry. The NSW EPA has recommendations for managing rendering plants such as cleaning spilled material in the premises, storage of rendering material, cleaning all equipment, machinery and bins, waste management and effluent treatment. In addition, the state EPA may impose licence requirements or conditions on the rendering plant e.g. treatment of the effluent.
15. Rendering is the process of separating the lipids or fats from animal tissue and water under the influence of heat and sometimes pressure. Variations in the process of rendering are employed by each rendering plant. Animal carcasses are usually processed as soon as they are delivered to the rendering plant. Generally, there are two principal methods of rendering.
16. In the wet rendering process, the tissue is ground to a small particle size of about 12 mm and preheated at around 95°C for between 5 and 60 minutes depending on the individual system. The heated slurry is then pressed or centrifugally separated into liquid and solid phases. The liquid which consists of lipids and water is then centrifugally separated into separate streams. The wet solids are dried then milled to a free-flowing meal.
17. In the dry rendering process, the tissue is ground to a particle size of about 30-40 mm then heated in a jacketed container, mechanical agitation is provided and the water evaporated either at atmospheric or increased pressure. The fat and solids are then separated over a screen. The fat is refined to remove any fine particles of solids remaining. The solids are pressed to remove excess fat then milled to a free-flowing meal.
18. In either case, continuous or batch processes may be utilised. Depending on the grade, the fat can be used for pharmaceuticals, food, soap or stock feed. The meal can be used in the pet food and fertiliser industries (Australian Renderers Association Inc. 2014).
19. Vapours from the condenser and those collected by hoods over the cookers and presses within the plant should be ducted to a treatment system such as a biofilter or afterburner. All wastewater, including washdown water and condensate not reused in boilers, is directed to an effluent treatment system (NSW Environment Protection Authority 2003).
	* 1. Site of release
20. The most likely route of administration of the GM vaccine would be via the drinking water system in commercial broiler chicken sheds as this is a more efficient way of vaccination than by eye drop. Even if the farm is free range, water would be provided inside the shed to minimise wild birds accessing water.
21. In the first phase of the field trial, only a few farms in Victoria that currently do not vaccinate against ILTV and are more isolated would be selected. In the second phase of the trial, more farms in NSW and Victoria would be selected in areas that have experienced previous ILTV outbreaks.
22. The main routes by which the GMO may enter the wider environment include spills of the GMO, and shedding of the GMO in tracheal exudates and faeces of vaccinated chickens.
	* 1. Related viral species in the receiving environment
23. The presence of related viral species may offer an opportunity for genetic recombination in the environment.
24. Three live attenuated ILTV vaccines are registered for use in chicken farms in Australia. From 2007-2015, ILTV outbreaks in NSW and Victoria have been caused by different classes of ILTV including those originally derived from the vaccine strains (see Section 5.9). ILT disease continues to be a problem in Australia with recent reports of ILT disease in both backyard and commercial poultry in NSW and Victoria (NSW Department of Primary Industries 2016; Victoria Department of Economic Development 2017). Information about the ILTV classes seen in Australian outbreaks is provided in Section 5.9.
25. Another virus belonging to the subfamily *Alphaherpesvirinae* that commonly infects poultry, including chickens is *Gallid herpesvirus* type 2 (Marek’s disease virus). Marek's disease affects both commercial and backyard poultry and is endemic in Australia.
26. *Psittacid herpesvirus 1* (PsHV-1) also belongs to the *Iltovirus* genus in the subfamily *Alphaherpesvirinae*. PsHV-1 causes Pacheco’s disease, an acute and potentially lethal respiratory infection in psittacine birds including macaws, parrots and cockatoos. Based on sequence analysis of PsHV-1 and ILTV, these viruses are relatively phylogenetically closely related. The similarity of their genomes suggests that they represent a class of avian alphaherpesviruses that diverged early from a common ancestor and are distinct from the Marek’s disease virus (Thureen & Keeler 2006). However, PsHV-1 and ILTV do not share the same host species.
27. There are also several other avian herpesviruses known but the herpesviruses tend to be host-specific. It is unlikely that recombination between different species of herpesviruses occur.
	* 1. Potential hosts in the environment
28. The potential for ILTV to infect other susceptible hosts that may be present at or near the proposed trial sites is taken into account in the risk assessment (Chapter 2). The primary host for ILTV is the chicken. ILT disease in turkeys, pheasants and peafowl are rarely reported (see Section 5.2). Throughout its long history since its initial reports in various parts of the world, ILTV outbreaks have occurred mostly in chicken farms.
29. In Australia, most chicken farms are usually separated by a large distance from other poultry farms and residential areas, and located in rural or semi-rural areas. However, some older chicken farms may be in close proximity to other poultry farms or residential areas. State and local council requirements set separation distances between sheds and houses external to the farm.
30. Chickens and other birds are likely to be kept in backyards outside the required separation distances imposed by local councils and/or state governments.
31. For biosecurity reasons, poultry farms only keep birds used for production. Some chicken farms may also rear, grow and sell turkeys, pheasants or game birds commercially. If more than one species of birds are produced on the farm, these must be housed and managed separately with suitable biosecurity arrangements for each species (Department of Agriculture 2009).
32. Other birds such as ducks may act as carriers of ILTV, but there is limited evidence of their role in spreading the virus and attempts to infect birds such as ducks were unsuccessful (see Section 5.2).
33. Production sheds on farms are designed to exclude wild birds and various biosecurity measures are in place to minimise wild birds accessing the production areas, including free range farms.
34. Australia has feral chickens, turkeys, pheasants and peafowls, from the family *Phasianidae*. Feral peafowls are declared as pests in Kangaroo Island, South Australia (South Australia Department of Environment 2017). Australia also has 3 native species in the *Phasianidae* family *(Coturnix pectoralis, Excalfactoria chinensis, and Coturnix ypsilohora)* (ABRS 2009), however their susceptibility to ILTV is unknown.
	1. Previous authorisations
		1. Australian authorisations
35. The APVMA has issued a permit for the use of the GM vaccine for research only. The GM vaccine has never been registered in Australia or elsewhere.
36. Work to develop the GMO in the laboratory including testing and preliminary experiments have been authorised under the Act as Notifiable Low Risk Dealings (NLRDs) conducted by the University of Melbourne and Royal Melbourne Institute of Technology (RMIT).
	* 1. International authorisations and experience
37. No application for the use or marketing of the GMO has been submitted to overseas regulatory authorities.
38. Risk Assessment
	1. Introduction
39. Risk assessment identifies and characterises risks to the health and safety of people or to the environment from dealings with GMOs (Figure 4). Risks are identified within the established risk context (see Chapter 1) and take into account current scientific and technical knowledge. Uncertainty and in particular, knowledge gaps, is considered throughout the risk assessment process.

**RISK ASSESSMENT PROCESS \***

**Risk**

**scenarios**

**Substantive Risks**

**Risk Evaluation**

*Consequence assessment*

*Likelihood assessment*

*Identification of substantive risks*

Negligible risks

RISK IDENTIFICATION

RISK CHARACTERISATION

**Risk context**

*Postulation of risk scenarios*

**\*** Risk assessment terms are defined in the *Risk Analysis Framework* 2013

Figure 4. The risk assessment process

1. Risk identification first considers a wide range of circumstances whereby the GMO, or the introduced genetic material, could come into contact with people or the environment. Consideration of these circumstances leads to postulating plausible causal or exposure pathways whereby dealings with a GMO (risk scenarios) may, in the short and long term, harm people or the environment.
2. Postulated risk scenarios are screened to identify substantive risks that warrant detailed characterisation. Substantive risks are further assessed when a risk scenario is considered to have some reasonable chance of causing harm. Pathways that do not lead to harm, or could not plausibly occur, do not advance in the risk assessment process.
3. Risk identification techniques used by the Regulator and evaluators at the OGTR include checklists, brainstorming, reported international experience and consultation. In conjunction with these techniques, risk scenarios postulated in RARMPs prepared previously for licence applications of the same and similar GMOs are also considered.
4. Substantive risks (i.e. those identified for further assessment) are characterised in terms of the potential seriousness of harm (Consequence assessment) and the likelihood of harm (Likelihood assessment). The level of risk is then estimated from a combination of the Consequence and Likelihood assessments. Risk evaluation then combines the Consequence and Likelihood assessments to determine level of risk and whether risk treatment measures are required. The potential for interactions between risks is also considered.
	1. Risk Identification
5. Postulated risk scenarios are comprised of three components (Figure 5):
6. Source of potential harm (risk source)
7. Plausible causal linkage to potential harm (causal pathway) and
8. Potential harm to an object of value (people or the environment).



Figure 5. Components of a risk scenario

1. In addition, the following factors are taken into account when postulating relevant risk scenarios for this licence application:
* the proposed dealings, which are conduct experiments with the GMO, transport and disposal of the GMO, and possession (including storage), supply and use in the course of any of these dealings
* restrictions placed on conduct of the experiments with the GMO, transport and disposal of GMO by other regulatory agencies and by the relevant States and local councils
* characteristics of the parent organism
* routes of exposure to the GMO
* potential for transmission
* potential exposure to the same gene from environmental sources
* the release environment
* practices during and after administration of the GM vaccine including broiler farming practices.
1. The APVMA has assessed the environmental safety and trade risks associated with the research trial use of the GM vaccine under APVMA permit in accordance with the AgVet Code and have determined they are satisfied that the risks associated with the use of the vaccine are acceptable when used in accordance with the conditions on the permit and in conjunction with OGTR approval. The APVMA has also considered the risk of recombination from the use of viral vaccines including the GM vaccine and requires that experiments be conducted to assess the ability of the GMO to recombine with other ILTV strains. The permit for the use of the GM vaccine addresses the following aspects:
* requirements for the directions for use, labelling, packaging, storage, and disposal of the GMO, contaminated materials and equipment, and chicken carcasses to ensure the safety of birds including non-target birds by limiting the spread and persistence of the GMO.
1. The current assessment focuses on risks posed to humans and the environment, including spread and persistence of the GMO beyond the field trials which may arise from inoculation of chickens, transport of live inoculated chickens, human and animal consumption of chickens, and transport and disposal of waste.
	* 1. Postulated risk scenarios
2. Six risk scenarios were postulated, as summarised in Table 2. These risk scenarios were evaluated considering both short and long term effects, restrictions imposed by APVMA, the current state and local council requirements, and in the context of practices proposed by the applicant. Detailed evaluations of these scenarios are provided later in this section. None of the risk scenarios were identified as a risk that could be greater than negligible and warranting further scrutiny.

Summary of risk scenarios from dealings with GMO

| **Risk scenario** | **Risk source** | **Causal pathway** | **Potential harm** | **Substantive risk?** | **Reason** |
| --- | --- | --- | --- | --- | --- |
| 1 | GM ILTV vaccine | Exposure of people handling the GMO or GMO-inoculated chickens during rearing, transport, processing or disposal  | Disease, toxicity or allergenicity | No | * ILTV has a very narrow host range, is not a human pathogen and is not expected to cause disease, toxicity or allergenicity in people
* ILTV is endemic in Australia but does not cause ill-health in people
* Other ILTV vaccines have a history of safe use with no adverse effects in people
* Vaccination would be conducted by trained workers supervised by a registered veterinarian or qualified personnel
* Only trained personnel would be allowed to handle the GMO
* Handling procedures in the poultry industry follow strict biosecurity measures
* Shedding of the GMO is expected to have ceased or declined to very low levels at the time of collection for processing
* Collection and transport of chickens to processing facilities minimise stress to protect animal welfare
* Processing facilities adhere to high standards of cleanliness and hygiene
 |
| 2 | GM ILTV vaccine | Exposure of people to the GMO when preparing or consuming meat from GMO-inoculated chickens | Disease, toxicity or allergenicity | No | * ILTV has a very narrow host range, is not a human pathogen and is not expected to cause disease, toxicity or allergenicity in people
* People are already exposed to meat that has come from chickens infected by ILTV strains or other ILTV vaccines, with no adverse effects
* Shedding of the GMO is expected to have ceased or declined to very low levels at the time of collection for processing
* Collection and transport of chickens to processing facilities minimise stress to protect animal welfare
* Processing plants employ hygiene and sanitation standards to ensure food safety as required by state authorities
* Chicken sold for human consumption lack the internal organs, gastrointestinal tract and head, which are the sites of infection of the GMO
* Cooking would destroy the GMO
* All food businesses in Australia are required to comply with the Food Safety Standards within the Food Standards Code, which includes cooking poultry thoroughly
 |
| 3 | GM ILTV vaccine | Exposure of susceptible wild birds to the GMO🡇Exposed birds become infected and develop ILT disease🡇ILT disease results in death🡇Decreased numbers of wild birds | Adverse impacts on desirable species | No | * ILTV has a very narrow host range
* Vaccination would be conducted by workers supervised by a registered veterinarian or qualified personnel
* Only trained personnel allowed to handle the GMO
* Vaccine spills would be disinfected
* Strict biosecurity measures are routinely exercised in chicken farms including free range farms
* Measures are in place to minimise wild birds accessing sheds, farms and water tanks where vaccination is conducted
* APVMA requires management of potential carriers
* Contaminated equipment, materials and vehicles would be disinfected after use
* Storage and disposal of used litter, carcasses and other contaminated farm waste would be done according to local council and state requirements
* Local council and state requirements impose conditions to prevent contamination of water sources
* Titres of infectious virus shed are expected to be low, so wild birds are unlikely to be challenged with a sufficient dose of GMO to develop ILT disease
* A large number of wild birds would have to be exposed to the GMO and develop severe ILT leading to death to impact wild species
 |
| 4 | GM ILTV vaccine | Exposure of susceptible wild birds to the GMO🡇Exposed birds become infected and develop a protective immune response🡇Reduced infection with virulent ILTV strains🡇Increase numbers of feral/pest birds | Adverse impacts on desirable species | No | * ILTV has a very narrow host range
* Chickens, pheasants and turkeys are not major pests in Australia
* Vaccination would be conducted by trained workers supervised by a registered veterinarian or qualified personnel
* Only trained personnel allowed to handle the GMO
* Vaccine spills would be disinfected
* Strict biosecurity measures are routinely exercised in chicken farms including free range farms
* Measures are in place to minimise wild birds accessing sheds, farms and water tanks where vaccination is conducted
* APVMA requires management of potential carriers
* Contaminated equipment, materials and vehicles would be disinfected after use
* Storage and disposal of used litter, carcasses and other contaminated farm waste would be done according to local council and state requirements
* Local council and state requirements impose conditions to prevent contamination of water sources
* Titres of infectious virus shed are expected to be low, so feral or pest birds are unlikely to be challenged with a sufficient dose of GMO to induce a protective immune response
* A large number of feral or pest birds would have to be exposed to the GMO and develop a protective immune response to impact other species
 |
| 5 | GM ILTV vaccine | Chickens are co-infected with the GMO and another ILTV strain🡇Recombination between GMO and other strain🡇Generation of a new virulent ILTV strain🡇Infection of chickens and/or other susceptible avian species | Increased disease burden in chickens and other susceptible avian species | No | * APVMA requirements include not using any other ILTV vaccine in a flock inoculated with the GMO, not vaccinating unhealthy birds and isolating treated flocks from susceptible populations of chickens not included in the trials
* Flocks vaccinated with other registered live attenuated vaccines as active controls would be housed in separate sheds and managed separately
* After the treated chickens have been removed from the shed and before a new batch of chickens is introduced, the shed would be fully cleaned
* State requirements to separate broiler farms from other poultry farms and sensitive uses
* Strict high level biosecurity measures and notification requirements for ILT disease would limit the spread of a virulent strain to susceptible flocks
 |
| 6 | GM ILTV vaccine | Establishment of the GMO outside the trial limits🡇Exposure of people, animals and susceptible birds to the GMO leading to risk scenarios 1-5 | As per Risk scenarios 1-5 | No | * Vaccine administration would take place in sheds
* All contaminated materials, equipment, sheds and vehicles would be disinfected after use
* APVMA requires disinfection of drinking system, cleaning of spills and management of potential carriers
* Strict biosecurity measures are routinely exercised in chicken farms including free range farms
* Collection and transport of chickens to processing facilities minimise stress to protect animal welfare
* Titres of infectious virus shed are expected to be low, so birds are unlikely to be challenged with a sufficient dose of GMO to induce disease or a protective immune response
* Temporary storage of used litter and carcasses and disposal of contaminated farm waste would be done according to local council and state requirements
* Local council and state requirements that impose conditions to prevent contamination of water sources
* State requirements to separate broiler farms from other poultry farms and sensitive uses
* Survival of the GMO is low at ambient conditions and in sunlight
 |

* + - 1. ***Risk scenario 1 – Exposure of people to the GMO***

|  |  |
| --- | --- |
| *Risk source* | GM ILTV vaccine |
| *Causal pathway* | Exposure of people handling the GMO or GMO-inoculated chickens during rearing, transport, processing or disposal |
| *Potential harm* | Disease, toxicity or allergenicity |

**Risk source**

1. The source of potential harm for this postulated risk scenario is the GM ILTV vaccine strain.

**Causal Pathway**

1. There are a number of ways that people may be exposed to the GMO while undertaking the dealings as part of this field trial or during subsequent processing of the chickens in commercial facilities.
2. People may be exposed directly to the GM vaccine during transportation of the GM vaccine to the farms. The GM vaccine is supplied as a freeze-dried pellet in a glass vial which makes it unlikely to leak. The applicant proposes to double-contain the GM vaccine, and this is also a requirement of the APVMA permit. Transport and storage of the GM vaccine would be in accordance requirements of the Regulator’s *Guidelines for the Transport, Storage and Disposal of GMOs*. These measures would reduce the likelihood of exposure of people to the GMO during transport.
3. The extent of exposure to people would be mainly limited to specific farms located in NSW and Victoria.
4. Exposure to people involved with the field trials may occur via inhalation of aerosols or splash during preparation of the GMO for administration. The freeze-dried vaccine needs to be reconstituted in water and, if the vaccine is to be administered via drinking water, the reconstituted vaccine would be diluted in a mixture of water and skim milk powder. The APVMA permit recommends the operator to wear eye protection and a mask while preparing and administering the GM vaccine, however, the applicant proposes to wear gloves and eye protection while preparing and administering the GM vaccine. In the event of a spill of the prepared GMO, the APVMA permit also requires that the spill area be treated using disinfectant. The applicant proposes that workers cleaning spills would wear gloves.
5. Prior to inoculation, workers would be trained in handling, preparing and administering the GM vaccine. Preparation of the GM vaccine and inoculation would be conducted under the supervision of a registered veterinarian or qualified personnel, and the workers would follow the vaccine label, APVMA permit conditions and trial protocol(s). After administration of the GMO, the APVMA permit requires any unused vaccine to be rendered non-viable. All contaminated vials, eye droppers, bottles and other materials would be soaked in disinfectant prior to disposal in the normal waste bin. These measures would minimise the likelihood of exposure to the GMO.
6. Once the GM vaccine has been administered to the chickens, shedding of GMO into the environment is likely to occur for a limited time while the GMO is actively replicating. As discussed in Chapter 1, Sections 5.5 and 6.3.7, the GMO was detected in the trachea of only a small fraction of chickens (2/21 chickens) 21 days post-inoculation, with the highest levels at 4 days post-inoculation (Coppo et al. 2011; Devlin et al. 2006). Studies have shown that chickens inoculated with wild type or other live attenuated ILTV vaccine strains shed the virus from 2 to 28 days post-inoculation. Peak shedding occurs from 4-9 days post-inoculation, and then shedding declines several logs by 14 days post-inoculation. Some wild type ILTV strains were no longer isolated from infected tissues in chickens at 11 and 14 days post-inoculation (Oldoni et al. 2009; Rodriguez-Avila et al. 2007; Roy et al. 2015). It is likely that the GMO would be shed for a similar period, with GMO levels declining to low levels or ceasing by 14 days post-inoculation. Stress may re-activate the virus and lead to further shedding, however re-activation takes some time. If the chickens were stressed during collection or transport, the earliest period in which replication of the virus would be expected to resume following latency is between 12-24 hours post-induction of the stress (Huang et al. 2011; Sawtell & Thompson 1992). Any exposure to the GMO from treated chickens shedding is likely to be much lower than the dose intentionally administered to chickens.
7. Shedding of the GMO from treated chickens means that people working at or visiting the trial sites (sheds and free range fields) may be exposed to the GMO. This could occur when handling the GMO-inoculated chickens (alive or dead carcasses) or cleaning sheds, vehicles and equipment used on site, and moving litter or waste. Strict biosecurity measures are followed at broiler farms including free range farms (refer to Chapter 1, Section 7 for more information). All workers and visitors must wear overalls and high rubber boots before entering sheds. Hands and boots are disinfected before and after entering the sheds. Veterinarians conducting post-mortem examination wear gloves and overalls, and disinfect hands after examination. Disposal of farm waste such as litter and chicken carcasses must be done in accordance with the state and/or local council requirements. These measures would reduce exposure to the GMO, but some exposure is still expected.
8. People could be exposed to the GMO shed from treated chickens when collecting them from the farm, transporting them to processing facilities and when handling chickens at these facilities. Chickens are proposed to be inoculated with the GM vaccine from 7-14 days old. Harvesting of broiler chickens usually commence from the age of 28 days. By the time the broiler chickens would be collected, shedding of the GMO is expected to have ceased or declined to low levels 14 days after inoculation and by 28 days of age.
9. After the chickens have recovered from ILT disease, the GMO may become latent, such that it can be reactivated by stressful situations and resume shedding. However, chickens that are not fit for transport are removed before pick-up, so only healthy chickens are transported to processing facilities, and collection is usually conducted at night under dim light to minimise stress on chickens. Transport of live chickens to the processing facilities would be in accordance with the state legislation to protect their welfare and minimise stress during transport.
10. Disposal of chicken carcasses at rendering facilities is conducted in accordance with state or local council requirements, ensuring high standards of cleanliness and hygiene. Processing and rendering facilities, where slaughter of chickens is conducted, are highly automated with minimal direct manual contact with chickens, minimising worker exposure to any microbiological contaminants. The rendering process would destroy any GMOs that may still be present.
11. As discussed in Chapter 1, Section 5.10, ILTV has been shown to survive only 7 hours in direct sunlight at ambient temperatures but would survive much longer in the dark and at low temperatures. Stability of the GMO when shed by chickens has not been studied but is expected to be similar to wild type ILTV strains, and therefore would deteriorate over time in field conditions.

**Potential harm**

1. ILTV has a very narrow host range and is not a human pathogen. ILTV occurs naturally in the environment, and live attenuated ILTV vaccines are widely used in poultry, so people working in the poultry industry are currently exposed with no reports of disease, infection (clinical or subclinical), toxicity or allergic reactions.
2. The GMO does not contain any new genetic material and the sequence is highly similar to the parent ILTV strain, with one gene deleted. As discussed in Chapter 1, Section 6.1, sequencing of the region of the genome from which the gG gene was deleted indicates that there is a theoretical potential for a novel 150 nucleotide mRNA transcript from across the deletion site, encoding a 27 amino acid protein. The expression of these has not been investigated. There may be potential that the expression of this 27 amino acid protein could produce adverse effects in a host such as toxicity. However, as toxic proteins have specific enzymatic properties, structural properties and recognise specific molecular targets, such a random sequence will unlikely to have any toxic properties (Hammond et al. 2013). As the genome sequences are not novel and exposure to the GMO is expected to be minimal, even if these products are expressed they are not expected to lead to any toxic or allergenic reactions. The small size of the potential protein makes it extremely unlikely to act as an allergen, as to elicit an allergic reaction a protein must contain at least two antibody binding sites, each 15 amino acids long, to facilitate cross-linking of antibodies. This gives a theoretical minimum size of approximately 30 amino acids (Huby et al. 2000).

**Conclusion**

1. Risk scenario 1 is not identified as a substantive risk because exposure is limited by standard industry handling and decontamination practices, including state-legislated practices, and the GMO is not expected to cause disease or other harms in people who are exposed. Therefore, this risk could not be considered greater than negligible and does not warrant further detailed assessment.
	* + 1. ***Risk scenario 2 – People consuming GM-vaccinated chickens***

|  |  |
| --- | --- |
| *Risk source* | GM ILTV vaccine |
| *Causal pathway* | Exposure of people to the GMO when preparing or consuming meat from GMO-inoculated chickens |
| *Potential harm* | Disease, toxicity or allergenicity |

**Risk source**

1. The source of potential harm for this postulated risk scenario is the GM ILTV vaccine strain.

**Causal Pathway**

1. It is proposed that broiler chickens treated with the GMO would enter the human food supply. Therefore, people may be exposed to the GMO, or to material from the GMO, when preparing or consuming meat from GMO-inoculated chickens.
2. As discussed in Risk Scenario 1, shedding of the GMO is expected to have ceased or declined to very low levels by the time harvest of broiler chickens normally commences at 28 days of age. Shedding may resume in stressed chickens, however re-activation of shedding takes considerable time, and stress is minimised by only harvesting healthy chickens, usually collecting chickens at night under dim light, and by transport in accordance with the state legislation designed to protect animal welfare.
3. As discussed in Chapter 1, Section 5.1, tissues in which ILTV may be found include the trachea, conjunctiva and Harderian gland in the eye, sinuses, thymus, lungs, kidneys, cecal tonsils and cloaca. ILTV is not known to replicate in chicken skeletal muscle. The GMO is expected to be found in these same tissues as other ILTV strains during active infection.
4. Also as discussed in Chapter 1, Section 5.10, ILTV in the trachea of chicken carcasses has been documented to survive at low temperatures for 30 days (Jordan 1966). However, processing facilities adhere to high standards of cleanliness and hygiene to ensure food safety. Any faecal contamination is reduced by following the Primary Production and Processing (PPP) Standard for Poultry Meat (Standard 4.2.2) (FSANZ 2010) that requires the processor to ensure that food is not made unsuitable or unsafe for human consumption. Furthermore, chicken meat sold for human consumption lack the internal organs, gastrointestinal tract and the head where the GMO may be present. Overall, the practices employed from collection of live chickens at the farm, transport and processing would minimise any residual GMO in the chicken meat or products derived from treated chickens. Any trace amount of GMO present would not survive cooking. All food businesses in Australia are required to comply with the Food Safety Standards within the Food Standards Code, which specifies what steps food businesses must take to ensure food is handled safely, including cooking poultry thoroughly (FSANZ 2010). Therefore, exposure to any GMO while preparing or consuming GMO-inoculated chickens is highly unlikely.

**Potential harm**

1. As described in Risk Scenario 1, ILTV is not a human pathogen, and the GMO is not expected to cause disease, toxicity or allergenicity in people.

**Conclusion**

1. Risk scenario 2 is not identified as a substantive risk because minimal, if any, exposure of people to the GMO from consumption of GMO-inoculated chickens is expected, and the GMO is not expected to cause disease or other harms in people who are exposed. Therefore, this risk could not be considered greater than negligible and does not warrant further detailed assessment.
	* + 1. ***Risk scenario 3 – Exposure of susceptible wild bird species to the GMO***

|  |  |
| --- | --- |
| *Risk source* | GM ILTV vaccine |
| *Causal pathway* | Exposure of susceptible wild birds🡇Exposed birds become infected and develop ILT disease🡇ILT disease result in death🡇Decreased numbers of wild birds |
| *Potential harm* | Adverse impacts on desirable species in the environment |

**Risk source**

1. The source of potential harm for this postulated risk scenario is the GM ILTV vaccine strain.

**Causal Pathway**

1. As discussed in Chapter 1, Section 5.2, ILTV has a very narrow host range, with the chicken being the primary host and reservoir, and the only other bird species observed to be naturally infected by ILTV are pheasants, peafowl and turkeys, all members of the family *Phasianidae* (Crawshaw & Boycott 1982; Portz et al. 2008).
2. There are several potential pathways by which susceptible wild bird species could come into contact with the GMO. Birds could be exposed to the GMO through various ways such as drinking water containing the GMO provided for treatment of chickens, through contact with spilt GMO or contaminated materials during disposal at the farm, direct contact with GMO-inoculated chickens, exposure to excreted fluids or faeces from vaccinated chickens, contact with pests potentially carrying the GMO, accessing areas used by treated chickens, drinking from surrounding waterways contaminated with the GMO or aerosols carrying the GMO.
3. As indicated in Chapter 1, Section 5.6, ILTV can be spread between nearby commercial poultry sheds or free range fields by air movement, particularly when tunnel ventilation is used. The capacity for transmission of the GMO has not been assessed in the field, however aerosols may lead to infection of susceptible wild birds close to sheds or free range fields containing GMO-inoculated chickens. Transmission to wild birds is expected to be less likely than transmission to a nearby shed employing tunnel ventilation because susceptible wild birds are not expected to be present in high numbers or densities, and would not remain in one place such that prolonged exposure may occur, particularly during the day when chickens in the shed or fields would be active and therefore shedding would be highest. There are also no reports of wild chickens, turkeys, peafowls or pheasants being infected or suffering from ILT disease.
4. Vaccination of broiler chickens with the GM vaccine would be performed according to the trial protocol(s), APVMA permit conditions and label instructions. The GM vaccine would be administered inside a shed (this includes free range farms), most likely by drinking water. Sheds are designed to limit access by wild birds. When administered in drinking water, the GM vaccine would be provided in an amount of water calculated to be consumed within 3-4 hours, and no additional water would be supplied until all of the GMO-containing water had been consumed. Even if the GM vaccine was prepared in an open bucket outside the shed prior to administration via drinking water, it would be unlikely that susceptible wild birds would come into contact with the GM vaccine in the bucket because there are measures in place to discourage wild birds accessing the farm, and the number of susceptible wild birds near broiler farms would be expected to be low. After administration of the GM vaccine, the APVMA permit requires any unused vaccine to be rendered non-viable. The APVMA also requires that in the event of a spill, the area must be treated with disinfectant. All contaminated bottles, vials, eye droppers, buckets and other materials would be soaked in disinfectant prior to reuse or disposal in the normal waste bin at the farm. These measures would greatly minimise the potential for exposure of susceptible wild birds to the GMO prepared for treatment.
5. As discussed in Chapter 1, Section 6.3, chickens treated with the GMO can transmit the GMO to other chickens in close physical contact. However, the minimum infective dose of the GMO has not been determined. The proposed field trials would study transmission of the GMO in the field.
6. Once the GM vaccine has been administered to the chickens, shedding of GMO into the environment via faeces or tracheal exudate is likely to occur for a limited time while the GMO is actively replicating. It is possible that susceptible wild birds could have access to production areas, particularly free range fields containing GMO-inoculated chickens. However, as discussed in Chapter 1, Section 5.10, ILTV does not survive well at ambient temperatures or when exposed to sunlight. For example, ILTV in tracheal exudates survived for 7 hours in direct sunlight, but particular environmental conditions such as low temperatures and darkness could potentially prolong virus survival (Jordan 1966). As discussed in Chapter 1, the GMO was detected in the trachea in a small percentage (9.5%) of chickens inoculated with the GMO up to 21 days, with high levels at 4 days post-inoculation (Coppo et al. 2011; Devlin et al. 2006). Studies have shown that chickens inoculated with wild type or other live attenuated ILTV vaccine strains shed the virus from 2 to 28 days post-inoculation. Peak shedding occurs from 4-9 days post-inoculation, then declining several logs by 14 days post-inoculation, and some virus strains were no longer isolated at 11 days post-inoculation (Oldoni et al. 2009; Rodriguez-Avila et al. 2007; Roy et al. 2015). At 14 days post-inoculation, it is expected that the amount of GMO shed would have ceased or declined significantly. Pick-up and transport of GMO-inoculated chickens could be done as early 28 days of age (14 days after inoculation of chickens), at which time, viral shedding, if any, would have ceased or be at low levels. Therefore, exposure of wild birds to viable GM virus would be low, and only likely when treated chickens are present and during the short period of peak viral shedding.
7. The susceptible wild birds could feed on pests potentially carrying the GMO, such as darkling beetles, living in broiler sheds. The APVMA permit requires that populations of wild birds and potential carriers, such as rodents and beetles, to be managed. Furthermore, current biosecurity measures require all broiler farms to control and manage vermin or pests at the farm, and to restrict access of wild birds to the production area including sheds or housing, water and feed. Sheds would be cleaned and decontaminated after removal of GMO-inoculated chickens. These measures would reduce the opportunity for exposure of susceptible wild birds to the GMO.
8. After the chickens have recovered from ILT disease, the GMO may become latent, such that it can be reactivated by stressful situations, such as during pick-up and transport, and resume shedding. Susceptible wild birds could be exposed to the GMO in the plume of dust, aerosols and debris from the GMO-inoculated chickens emitted from an open truck travelling to the processing plant. However, chickens that are not fit for transport are removed before pick-up, so only healthy chickens are transported to processing facilities, and collection is usually conducted at night under dim light to minimise stress on chickens. Transport of live chickens to the processing facilities would be in accordance with the state legislation to protect their welfare and minimise stress during transport. If the chickens were stressed during collection or transport, the earliest period in which replication of the virus would be expected to resume from latency may be between 12-24 hours post-induction of stress (Huang et al. 2011; Sawtell & Thompson 1992). By this time, the chickens would have arrived at the processing plant for slaughtering.
9. Susceptible wild birds could be exposed to the GMO at disposal sites containing contaminated farm waste at the farm or off-site. Used litter and chicken carcasses may be temporarily stored at the farm prior to disposal. Transport of litter and carcasses would be covered in a truck to prevent dispersal. Temporary storage of used litter prior to disposal would be covered with clean co-compost material and a tarpaulin, and chicken carcasses would be temporarily stored in a freezer. There are state and local council requirements for the temporary storage of chicken carcasses and used litter to minimise the spread of pathogens.
10. Disposal of contaminated farm waste, including litter, manure and chicken carcasses are conducted according to state and local council requirements (see Chapter 1, Section 7 for more information). Such waste may be disposed of at the farm by composting, burial, or taken off-site to a landfill or a commercial composting facility or, for carcasses, to a rendering facility. Compost in piles or open windrows is covered with at least 300 mm thickness of clean co-compost material to exclude birds accessing the litter and carcasses. Studies have shown that composting for 120 hours or heating litter to 38°C for 24 hours renders ILTV undetectable by the highly sensitive PCR method (Giambrone et al. 2008). Burial at the farm is uncommon but if conducted, waste would be covered by at least 2 m of soil. Landfills are required to cover the carcasses daily with at least 150 - 300 mm of soil or other material. There are a number of state requirements to minimise access of vermin including birds at these disposal sites. These measures would reduce exposure of susceptible wild birds to the GMO at these waste storage or disposal sites.
11. Waste or stormwater run-off from sheds, outdoor areas used by GMO-inoculated chickens or waste disposal areas may be contaminated with the GMO, leading to contamination of various water sources used by susceptible wild birds. However, as discussed in Chapter 1, local councils and state authorities have a number of regulations to ensure that water sources and catchment areas are not contaminated with run-off from the waste facilities, disposal sites and poultry farms.

**Potential harm**

1. As discussed in Chapter 1, Section 6.3, the GMO has been shown to cause milder ILT disease in chickens and a lower death rate than the parent ILTV strain (Devlin et al. 2006). Thus, even if a susceptible wild bird becomes infected with the GMO, the disease is not expected to be any worse than those caused by circulating ILTV strains. The host immune response would likely clear the GMO that would limit its spread in the body. There are also no reports of wild susceptible birds suffering or dying from ILT disease.
2. For this scenario to lead to harm to the environment, a large number of wild birds would have to become infected with the GMO, and ILTV would need to be an important factor limiting the wild bird population. However, as discussed above, ILTV has a very narrow host range limited to chickens, pheasants, peafowls and turkeys. These wild species of birds are not found in large numbers near commercial broiler farms.

**Conclusion**

1. Risk scenario 3 is not identified as a substantive risk because unintentional exposure of susceptible wild birds to the GMO is expected to be low and the potential harm to the environment from unintentional exposure is minimal. Therefore, this risk could not be considered greater than negligible and does not warrant further detailed assessment.
	* + 1. ***Risk scenario 4 – Immune response in susceptible feral/pest bird species to the GMO***

|  |  |
| --- | --- |
| *Risk source* | GM ILTV vaccine |
| *Causal pathway* | Exposure of susceptible wild birds🡇Exposed birds become infected and develop a protective immune response🡇Reduced infection with virulent ILTV strains🡇Increase numbers of feral/pest birds |
| *Potential harm* | Adverse impacts on desirable species in the environment |

**Risk source**

1. The source of potential harm for this postulated risk scenario is the GM ILTV vaccine strain.

**Causal Pathway**

1. As discussed in Risk Scenario 3, the only susceptible birds are chickens, turkeys, pheasants and peafowls and the number of these birds present in the wild is expected to be low. Susceptible wild birds could be exposed to the GMO through various routes, but the proposed measures and current requirements by the APVMA, state and local councils would minimise the likelihood of exposure to the GMO.

**Potential harm**

1. As discussed in Chapter 1, Section 6.3, the GMO has been shown to cause milder ILT disease in chickens and a lower death rate than the parent ILTV strain (Devlin et al. 2006). Furthermore, chickens inoculated with the GMO did not become infected with the CSW-1 ILTV strain when later exposed (Devlin et al. 2011). Susceptible birds that were unintentionally infected with the GMO may similarly be protected from other virulent ILTV strains present in the field. This may lead to increased survival of pest birds that have been exposed to the GMO, particularly during an ILTV outbreak. Increased numbers of pest birds in the environment could adversely impact other, desirable, species.
2. For this scenario to lead to harm to the environment, a large number of feral or pest birds would have to become infected with the GMO, and ILTV would need to be an important factor limiting the pest bird population. However, as discussed above, ILTV has a very narrow host range limited to chickens, pheasants, peafowls and turkeys, which are not significant pests in mainland Australia and not found in large numbers near commercial broiler farms.

**Conclusion**

1. Risk scenario 4 is not identified as a substantive risk because unintentional exposure of susceptible feral or pest birds to the GMO is expected to be low and the potential harm to the environment from unintentional exposure is minimal as susceptible species are not significant pests. Therefore, this risk could not be considered greater than negligible and does not warrant further detailed assessment.
	* + 1. ***Risk scenario 5 – Recombination between GMO and viruses***

|  |  |
| --- | --- |
| *Risk source* | GM ILTV vaccine |
| *Causal pathway* | Chickens are co-infected with the GMO and another ILTV strain🡇Recombination between the GMO and other strain🡇Generation of a new virulent ILTV strain🡇Infection of chickens and/or other susceptible avian species |
| *Potential harm* | Increased disease burden in chickens and other susceptible avian species |

**Risk source**

1. The source of potential harm for this postulated risk scenario is the GM ILTV vaccine strain.

**Causal Pathway**

1. The probability of recombination occurring in viruses is dependent on co-circulation of different viruses in the same geographical area, genetic similarity between the viruses, rate of co-infection of a host with both viruses and viral population size within the infected host.
2. As discussed in Chapter 1, Section 5.8, ILTV is endemic in Australia with different classes of ILTV circulating in NSW and Victoria, and there are three registered live attenuated ILTV vaccine strains (ie. SA2, A20 and Serva) currently used. New classes of ILTV that have caused outbreaks in Australian in recent years may have resulted from recombination between wild type ILTV and ILTV vaccine strains (Agnew-Crumpton et al. 2016; Blacker et al. 2011; Lee et al. 2012). Therefore, co-infection of a host cell by different ILTV strains does occur, however recombination may have been facilitated by using a combination of the three ILTV vaccines on a single flock (Agnew-Crumpton et al. 2016; Coppo et al. 2013).
3. At the trial sites, GM-vaccinated broiler chickens may be inadvertently exposed to circulating ILTV. However, the APVMA permit only allows the GMO to be given to healthy chickens, vaccination of the same flock with another type of ILTV vaccine is not permitted, and treated flocks must be isolated from susceptible populations of chickens that are not involved in the trial. When studying transmission of the GMO, unvaccinated sentinel chickens may be housed in a shed containing GMO-inoculated chickens.
4. Flocks treated with the GMO or sentinel chickens exposed to GMO-inoculated chickens are expected to develop protective immunity to other ILTV strains. In addition, biosecurity measures would also minimise the likelihood of exposure of GMO-inoculated chickens to other ILTV strains circulating in the environment during the trial. This would reduce the likelihood of co-infection and recombination with other ILTV strains.
5. During the trials, where a broiler farm has multiple sheds, flocks in different sheds may be given the GMO or another ILTV vaccine, or remain unvaccinated. Free range farms would not have an unvaccinated shed, which reduces the potential for cross-contamination. The same poultry farm may also be growing other susceptible bird species, such as turkeys, commercially. Each shed would be clearly identified and managed separately, as a biosecurity measure. To minimise cross-contamination, people entering/exiting sheds must disinfect hands and boots, movement of people and vehicles would be controlled and all contaminated equipment must be disinfected after use. The measures would reduce the likelihood of co-infection of broiler chickens and other susceptible bird species grown at the trial farms with different vaccine strains.
6. Chickens in neighbouring poultry farms or backyards adjacent to the trial areas may be exposed to the GMO via aerosols in the wind. Aerosol transmission would depend on the direction of the wind and weather conditions. As discussed in Risk Scenarios 1-4, the controls proposed by the applicant, the APVMA permit conditions, local council and state requirements and the strict biosecurity measures employed at poultry farms would minimise the likelihood of unintentional exposure of chickens in neighbouring poultry farms or backyards adjacent to the trial area to the GMO. Furthermore, state authorities require new broiler farms including free range farms to have separation distances, buffer zones and boundary setbacks, which may reduce the impact of aerosol transmission. These would reduce the potential for co-infection of chickens, including those in other poultry farms or backyards, with the GMO and other ILTV strains, thereby minimising the potential for recombination. However, older farms may be clustered together in close proximity and to rural residential areas. Poultry that were vaccinated against ILTV may recombine with the GMO if it was unintentionally spread to these areas. The spread of the GMO to adjacent poultry farms or backyards clustered together would be minimised by administering the GMO to chickens in sheds and, for non-free range farms, keeping the chickens in sheds.
7. After one flock of GMO-inoculated chickens has been removed from a shed and before a new flock is introduced, the shed would be fully cleaned and disinfected to avoid unintentionally infecting the new flock with the GMO. As discussed in Chapter 1, darkling beetles are a common pest in broiler sheds that live on the shed floors and ceilings. These insects may be resistant to certain insecticides. Subsequent batches of chickens in the shed used to house GMO-inoculated chickens could feed on insect potentially carrying the GMO, such as darkling beetles, living in broiler sheds. The APVMA permit requires that populations of wild birds and potential carriers, such as rodents and beetles, to be managed. Furthermore, current biosecurity measures require all broiler farms to control and manage vermin or pests at the farm. These measures would minimise the spread of the GMO to subsequent batches of chickens in the shed that may be vaccinated with other ILTV vaccine strains, thereby minimising the potential for recombination.
8. Live transport of inoculated chickens in open trucks passing by susceptible birds in farms or backyards may potentially disperse and transmit the GMO via aerosols, dust, or debris. These susceptible birds in farms or backyards may be inoculated with the available ILTV vaccines or be infected with the circulating ILTV strains. As discussed above, shedding of the GMO would have declined several logs or ceased at 14 days of inoculation. Harvesting and transport procedures in accordance with the state legislation to protect animal welfare would minimise stress in chickens. If the chickens were stressed during collection or transport, the earliest period in which replication of the virus would be expected to resume from latency is between 12-24 hours post-induction of stress (Huang et al. 2011; Sawtell & Thompson 1992). By this time, the chickens would have arrived at the processing plant for slaughtering. Therefore, it is unlikely that susceptible birds in farms or backyards would be inadvertently exposed to the GMO and become co-infected with the GMO and other circulating ILTV strains or vaccine strains.

**Potential harm**

1. Recombination between the GMO and another ILTV strain could result in viral progeny having any permutation of genomic segments of the two parent strains. Recombination could produce a less, similar or more virulent phenotype than either parent strain.
2. The CSW-1 strain is derived from an Australian field isolate from 1959, so is not novel to Australia. The class of ILTV to which CSW-1 belongs has not been identified in recent outbreaks, so it is not clear if it is currently circulating. However, it has been prevalent in the past and is likely to have contributed to the genetic makeup of current strains through past recombinations, so it is not expected to add a significant level of genetic variation to the current pool of circulating viruses. Comparison of the full genomic sequences of CSW-1 to the other live attenuated ILTV vaccine strains reveals that they are between 99.69% and 99.82% identical. The GMO does not contain any novel sequences or genes, with the only modification being deletion of the gG gene. Furthermore, the gene deletion would have increased genetic and phenotypic stability in the GM viral genome compared to a point mutation, which is more capable of reversion to the wild-type sequence, especially if the virus has a high mutation rate (Bull 2015; Hanley 2011). Any recombinant carrying the gG gene deletion is expected to retain the associated attenuated phenotype of the GMO. Nevertheless, despite the seemingly low likelihood of recovery of a viral gene from deletion, viruses originally containing deletions, but which have now regained the gene, have been isolated in laboratory experiments (Bull 2015; Jimenez-Guardeno et al. 2015). These factors make the likely outcome of any recombination between the GMO and another ILTV strain to be a virus of similar or lower virulence than the other strain involved.
3. The APVMA requires that experiments be conducted to assess the ability of the GM vaccine to recombine with other ILTV strains. This would be taken into consideration by the APVMA in assessing the risks of using the GM vaccine.
4. In the unlikely event of a novel virulent ILTV strain arising from recombination between the GMO and another ILTV strain within a farm participating in the trial, the opportunity for it to spread to other susceptible birds would be restricted by higher level of biosecurity measures and notification requirements for ILT disease.

**Conclusion**

1. Risk Scenario 5 is not identified as a substantive risk as the opportunity for recombination is restricted by the biosecurity measures employed and recombination between the GMO and another ILTV strain is expected to result in a virus of less or similar virulence than the current circulating ILTV strains. Therefore, this risk could not be considered greater than negligible and does not warrant further detailed assessment.
	* + 1. ***Risk scenario 6 – Spread and Persistence of the GMO***

|  |  |
| --- | --- |
| *Risk source* | GM ILTV vaccine |
| *Causal pathway* | Establishment of the GMO outside the trial limits 🡇Exposure of people, animals and susceptible birds to the GMO leading to risk scenarios 1-5 |
| *Potential harm* | As per risk scenarios 1-5 |

**Risk source**

1. The source of potential harm for this postulated risk scenario is the GM ILTV vaccine strain.

**Causal Pathway**

1. The GMO could be dispersed in the wider environment via pathways discussed in Risk Scenarios 1-5, and may persist in the environment if it is able to establish an infection cycle in susceptible wild birds or farmed poultry.
2. As discussed for these scenarios, the limits and controls proposed by the applicant, APVMA permit conditions, biosecurity measures, and local council and state requirements would limit the dispersal of the GMO in the environment. For the GMO to establish it must be able to be transmitted efficiently between flocks. The capacity of the GMO to be transmitted between birds in close contact has been demonstrated in the laboratory, however transmission has not been assessed in the field.
3. Establishment of the GMO in the wider environment may lead to ongoing exposure of susceptible birds in the wild, commercial or backyard poultry, other animals and people beyond the trial limits.

**Potential harm**

1. If the GMO were to establish outside of the trial limits, ongoing exposure is not expected to cause harm to people or the environment. The GMO is less virulent than circulating ILTV strains, so would cause less-severe disease in exposed susceptible birds than these strains. Other potential harms are not expected to be significant for the reasons discussed in risk scenarios 1-5.

**Conclusion**

1. Risk scenario 6 is not identified as a substantive risk because spread of the GMO outside the trial limits would be minimised by the limits and controls proposed by the applicant, State, local council and APVMA requirements, and no significant adverse effects from the GMO have been identified. Therefore, this risk could not be considered greater than negligible and does not warrant further detailed assessment.
	1. Uncertainty
2. Uncertainty is an intrinsic part of risk and is present in all aspects of risk analysis[[7]](#footnote-8).
3. There are several types of uncertainty in risk analysis (Bammer & Smithson 2008; Clark & Brinkley 2001; Hayes 2004). These include:
* uncertainty about facts:
* knowledge – data gaps, errors, small sample size, use of surrogate data
* variability – inherent fluctuations or differences over time, space or group, associated with diversity and heterogeneity
* uncertainty about ideas:
* description – expression of ideas with symbols, language or models can be subject to vagueness, ambiguity, context dependence, indeterminacy or under-specificity
* perception – processing and interpreting risk is shaped by our mental processes and social/cultural circumstances, which vary between individuals and over time.
1. Uncertainty is addressed by approaches such as balance of evidence, conservative assumptions, and applying risk management measures that reduce the potential for risk scenarios involving uncertainty to lead to harm. If there is residual uncertainty that is important to estimating the level of risk, the Regulator will take this uncertainty into account in making decisions.
2. As field trials of GMOs are designed to gather data, there are generally data gaps when assessing the risks of a field trial application. However, field trial applications are required to be limited and controlled. Even if there is uncertainty about the characteristics of a GMO, limits and controls restrict exposure to the GMO, and thus decrease the likelihood of harm.
3. For DIR 154, uncertainty is noted particularly in relation to:
* the degree of attenuation of the GMO under field conditions
* the level of shedding of infectious GMO from inoculated chickens
* the ability of the GMO to be transmitted from inoculated chickens to other chickens or other birds in the natural environment
* the ability of the GMO to establish an infection cycle and persist in the environment.
1. These areas of uncertainty have been accommodated in the risk assessment by assuming that shedding, transmission and persistence may be equal to other ILTV strains which are able to spread and persist in the environment. Accommodating this uncertainty resulted in an estimate of risk of negligible.
2. Additional data, including information to address these uncertainties, may be required to assess possible future applications with reduced limits and controls, such as a commercial release of the GMO.
3. Chapter 3, Section 4, discusses information that may be required for future release.
	1. Risk evaluation
4. Risk is evaluated against the objective of protecting the health and safety of people and the environment to determine the level of concern and, subsequently, the need for controls to mitigate or reduce risk. Risk evaluation may also aid consideration of whether the proposed dealings should be authorised, need further assessment, or require collection of additional information.
5. Factors used to determine which risks need treatment may include:
* risk criteria
* level of risk
* uncertainty associated with risk characterisation
* interactions between substantive risks.
1. Six risk scenarios were postulated whereby the proposed dealings might give rise to harm to people or the environment. In the context of the current release sites, limits and controls proposed by the applicant, the APVMA permit conditions, biosecurity measures, local council and state requirements, and considering both the short and long term consequences, none of these scenarios were identified as substantive risks. The principal reasons for these conclusions are summarised in Table 2 and include:
* attenuated phenotype of the GMO
* ILTV’s limited host range
* APVMA permit conditions for the use of GM vaccine
* local council and state requirements for broiler farms, processing and rendering facilities, and waste disposal
* suitability of the controls proposed by the applicant.
1. Therefore, risks to the health and safety of people, or the environment, from the proposed release of the GMO into the environment are considered to be negligible. The *Risk Analysis Framework* (OGTR 2013), which guides the risk assessment and risk management process, defines negligible risks as risks of no discernible concern with no present need to invoke actions for mitigation. Therefore, no controls are required to treat these negligible risks. Hence, the Regulator considers that the dealings involved in this proposed release do not pose a significant risk to either people or the environment.
2. Risk management plan
	1. Background
3. Risk management is used to protect the health and safety of people and to protect the environment by controlling or mitigating risk. The risk management plan addresses risks evaluated as requiring treatment and considers limits and controls proposed by the applicant, as well as general risk management measures. The risk management plan informs the Regulator’s decision-making process and is given effect through licence conditions.
4. Under Section 56 of the Act, the Regulator must not issue a licence unless satisfied that any risks posed by the dealings proposed to be authorised by the licence are able to be managed in a way that protects the health and safety of people and the environment.
5. All licences are subject to three conditions prescribed in the Act. Section 63 of the Act requires that each licence holder inform relevant people of their obligations under the licence. The other statutory conditions allow the Regulator to maintain oversight of licensed dealings: Section 64 requires the licence holder to provide access to premises to OGTR inspectors and Section 65 requires the licence holder to report any information about risks or unintended effects of the dealing to the Regulator on becoming aware of them. Matters related to the ongoing suitability of the licence holder are also required to be reported to the Regulator.
6. The licence is also subject to any conditions imposed by the Regulator. Examples of the matters to which conditions may relate are listed in Section 62 of the Act. Licence conditions can be imposed to limit and control the scope of the dealings. In addition, the Regulator has extensive powers to monitor compliance with licence conditions under Section 152 of the Act.
	1. Risk treatment measures for substantive risks
7. The risk assessment of risk scenarios listed in Chapter 2 concluded that there are negligible risks to people and the environment from the proposed trial of the GM vaccine. These risk scenarios were considered in the context of the scale of the proposed release and the proposed containment measures (which include standard industry practice, APVMA permit conditions, and state and local requirements), and considering both the short and the long term. The risk evaluation concluded that no specific risk treatment measures are required to treat these negligible risks. Limits and controls proposed by the applicant and other general risk management measures are discussed below.
	1. General risk management
8. The limits and controls proposed in the application were important in establishing the context for the risk assessment and in reaching the conclusion that the risks posed to people and the environment are negligible. Therefore, to maintain the risk context, licence conditions have been imposed to limit the release to the proposed size, location and duration, and to restrict the spread and persistence of the GMO and its genetic material in the environment. The conditions are discussed and summarised in this Chapter and listed in full in the licence.
	* 1. Licence conditions to limit and control the release
			1. Consideration of limits and controls proposed by Bioproperties
9. Chapter 1 provides details of the limits and controls proposed by Bioproperties in their application. Many of these are discussed in the six risk scenarios characterised for the proposed release in Chapter 2. The appropriateness of these limits and controls is considered further below.
10. The applicant proposes that a maximum of 40 broiler chicken farms would be included in the field trials in NSW and Victoria and the duration of the field trials would be limited to five years. The applicant proposes inoculating up to 2 million broiler chickens, and would exclude layers and breeders. Access to the farms participating in the trial would be controlled. People visiting the Trial areas would be aware of the presence of the GMO, as signs are required to be placed at all entrances to the Trial areas. These measures would limit the potential exposure of humans and other organisms to the GMO (Risk scenarios 1-6), and are included in the licence.
11. The APVMA permit for the GM vaccine recommends the operator to wear eye protection and a mask while preparing and administering the GM vaccine, which would reduce the likelihood of the GM vaccine entering a person’s eyes and mouth as well as being inhaled. The applicant also proposes that workers preparing the GM vaccine would wear gloves, in addition to eye protection (Risk scenario 1). Wearing personal protective equipment including eye protection and gloves would be sufficient to reduce exposure of workers to the GMO and are included in the licence. Additionally, vaccination would be conducted by trained farm workers under the supervision of a registered veterinarian or qualified personnel, ensuring that workers handle the GM vaccine appropriately. These measures have been included in the licence.
12. The applicant has stated that the administration age of the chickens being inoculated will be from 7 to 14 days old. As discussed in Chapter 1, live attenuated ILT vaccines and wild type ILTV are shed at peak levels from chickens from 4-9 days post-inoculation, and decline to low levels by the time chickens would start to be harvested at 28 days of age. To minimise the exposure of people outside the trial areas to the GMO (Risk scenarios 1 and 2) and the potential for dispersal of the GMO (Risk scenarios 3-6), conditions have been imposed in the licence which prohibits the harvesting and transport of GMO-treated chickens to the processing plants within 14 days of inoculation or if the chickens display clinical signs or symptoms of ILT disease. These would ensure minimal shedding of the GMO at harvest and minimise the likelihood of dispersing the GMO during transport of the live chickens.
13. The APVMA permit only allows the GM vaccine to be given to healthy chickens, vaccination with any other ILTV vaccine is not permitted and requires GMO-inoculated chickens to be isolated (Risk scenario 5). Unvaccinated sentinel chickens may be housed in the same shed as GMO-inoculated chickens to study transmission of the GMO. A licence condition has been imposed requiring these unvaccinated sentinel chickens to be treated in the same way as GMO-inoculated chickens. If a farm has multiple sheds and each shed may be vaccinated with different ILTV vaccines or remain unvaccinated, the applicant proposes that each shed would be managed separately, people must disinfect their hands and boots when entering/exiting the shed, movement of people would be controlled, and contaminated equipment disinfected after use. These measures would reduce the potential for recombination. The licence conditions include segregating GMO-inoculated chickens from all other poultry, decontaminating hands and footwear when exiting a Shed or Range, and decontaminating equipment. The licence also requires that a Compliance Management Plan, addressing cross-contamination and segregation of treated flocks, be provided to the Regulator.
14. The applicant proposes that GMO-inoculated chickens would be confined to sheds, and if relevant, to free range fields until the time of harvest. Effective containment of live GMO-inoculated chickens as well as employing biosecurity measures at the trial areas would reduce exposure of susceptible birds to the GMO and dispersal of the GMO in the environment (Risk scenarios 3-6). Conditions are included in the licence requiring that experimentation with the GMO, GMO-inoculated chickens or samples may only be undertaken within a trial area unless conducted in a certified PC2 facility under an NLRD authorisation, and a contingency plan addressing any escape or predation of GMO-inoculated chickens be provided to the Regulator. However, the GMO-inoculated chickens shed the virus, with peak shedding occurring 4-9 days post-inoculation, then declining or ceasing by 14 days post-inoculation. Licence conditions have been imposed requiring administration of the GMO inside a shed, and confinement of chickens in the shed for the first 14 days post-inoculation or while displaying signs or symptoms of infectious laryngotracheitis. These measures would minimise exposure of GMO-treated chickens to wild birds or other animals entering the trial farms.
15. Strict biosecurity and waste management measures as well as other effective broiler farm management are routinely applied in broiler farms to minimise pathogen occurrence and spread, and to protect people and the environment. Many measures are required under State legislation, local council regulations or the APVMA permit relevant to this trial. These measures combined minimise exposure to, and dispersal of, the GMO and are discussed in detail below.
16. State guidelines, legislation and/or local council requirements stipulate separation distances or biosecurity buffer distances for new broiler farms from other commercial poultry farms and sensitive areas. To minimise the likelihood of dispersal of the GMO via aerosols or particulates, a licence condition has been imposed requiring that the boundary of the Trial area on a Participating farm be at least 1000 metres from the poultry located on other poultry farms that are not included in the field trials. This requirement is consistent with current recommendations for separation distances for poultry farms in NSW and Victoria (as discussed in Chapter 1, Sections 7.1.1 and 7.1.2). Furthermore, state and local council requirements stipulate various proximity distances of broiler farms or disposal sites to water sources (Risk scenarios 3-6). Therefore, licence conditions require the trial areas to be at least 50 m from waterways (notwithstanding requirements of state and council regulations). A licence condition has also been included requiring immediate notification of any extreme weather conditions affecting the trial area to allow assessment and management of any risks.
17. Decontamination measures for people, sheds and equipment are proposed by the applicant. For people, this includes supplying and wearing overalls and high rubber boots to all shed visitors and workers, and disinfecting hands and boots when entering and exiting the shed. All equipment and materials contaminated with the GMO such as bottles, vials, droppers and other GMO-contaminated materials would be disinfected after use. Vehicles and equipment used during transport would be disinfected after use. Litter and dead chickens would be disposed of by composting, burial, rendering or landfill following State/Territory and/or local council requirements. These measures would limit spread and persistence of the GMO (Risk scenarios 3-6). The licence requires that a Compliance Management Plan, addressing decontamination measures and entry/exit procedures, be provided to the Regulator.
18. The APVMA permit states ‘the shed and litter are to be treated between flocks in a manner which is effective against the vaccine virus’. To minimise spread and persistence of the GMO, conditions have been imposed in the licence requiring decontaminating any areas used to temporarily store carcasses or litter which are potentially contaminated with the GMO, decontaminating sheds between batches of chickens and not permitting the re-use of contaminated waste, including litter.
19. The APVMA permit has a requirement to manage pests that are potential carriers of the GMO (e.g. dogs, cats, rodents, wild birds and darkling beetles). This would minimise the transmission of the GMO to susceptible wild or feral/pest birds, transmission to the next batch of chickens introduced in the shed that housed GMO-inoculated chickens, and the spread of the GMO in the environment (Risk Scenarios 3-6). The licence requires that a Compliance Management Plan, addressing pest, vermin and wild bird management, be provided to the Regulator.
20. As discussed in Chapters 1 and 2, there is the potential for the GMO to persist at a site following completion of the trial. This may potentially occur in the sheds even after thorough decontamination (such as via the presence of darkling beetles), or out on a range if there are areas that are protected from heat or sunlight. Therefore, additional licence conditions have been imposed requiring monitoring for clinical signs and symptoms of infectious laryngotracheitis in the Flock and testing for the unintended presence of the GMO in chickens displaying these signs or symptoms after they are introduced in the decontaminated shed. The licence holder is required to keep records of such monitoring and testing data and to notify the Regulator of any positive findings. This measure will confirm that the GMO has not persisted at a site after the completion of the field trials.
21. Live GMO-inoculated chickens would be transported to processing facilities as well as to laboratories for analysis. Transport to processing facilities would be in accordance with the relevant state legislation, which includes use of ventilated crates and transport in an open truck. As discussed in Chapters 1 and 2, peak shedding of the GMO is likely to occur 4-9 days post-inoculation or when displaying symptoms of ILT disease. Therefore, a licence condition has been imposed prohibiting the transport of chickens to processing facilities within 14 days post-inoculation or if displaying signs or symptoms of ILT disease. Live GMO-treated chickens may also be transported to research facilities for further study. If live GMO-treated chickens are required to be transported to the PC2 facilities within 14 days post-inoculation or if displaying signs or symptoms of ILT disease, a licence condition has been imposed that requires measures to be implemented to minimise dispersal of the GMO during transport. The licence requires that a Compliance Management Plan, addressing collection and transport of live GMO-inoculated chickens, be provided to the Regulator. The licence also requires that transport routes be provided to the Regulator.
22. The GM vaccine and samples containing the GMO (excluding live GMO-inoculated chickens) would be transported and stored according to the Regulator’s *Guidelines for the Transport, Storage and Disposal of GMOs*. These are standard protocols for the handling of GMOs to minimise exposure to the GMOs (Risk Scenarios 1 and 3), and dispersal into the environment (Risk Scenario 6). The licence requires that transport and storage of the GM vaccine and samples be in accordance with the Guidelines.
23. The applicant proposes to destroy any GMO-inoculated chickens not required for experimentation or transported to processing facilities. This would minimise the spread and persistence of the GMO (Risk scenario 6), and has been included in the licence.
24. Bioproperties would be required to submit a Compliance Management Plan to the Regulator before inoculating chickens with the GMO at each farm. This plan would detail how the licence holder intends to comply with the licence conditions, including decontamination processes and compliance with State, local council and industry requirements/guidelines.
	* + 1. Summary of licence conditions to be implemented to limit and control the release
25. A number of licence conditions have been imposed to limit and control the proposed release, based on the above considerations. These include requirements to:
* limit the field trials to a maximum of 40 farms in NSW and Victoria, from the date of licence issue to August 2022
* limit to a maximum of 2,000,000 broiler chickens that may be inoculated with the GMO
* locate the trial sites at least 1000 m from poultry located on other poultry farms
* locate the trial sites at least 50 m away from waterways
* confine GMO-inoculated chickens in the sheds for the first 14 days after administration of the GMO or if displaying symptoms of ILT disease
* prohibit the harvesting of GMO-inoculated chickens within 14 days of inoculation or if displaying symptoms of ILT disease for transport to Processing facilities
* manage pests, vermin and other animals
* decontaminate all sheds and equipment that have been contaminated with the GMO and peoples’ hands and footwear upon exit of shed or range
* not allow the use of other vaccines against ILTV in GMO-inoculated chickens
* transport and store the GMO and samples from GMO-inoculated chickens in accordance with the Regulator’s *Guidelines for the Transport, Storage and Disposal of GMOs*, in force at the time
* demonstrate compliance with a range of relevant State and local requirements and guidelines
* destroy all GMOs and GMO-inoculated chickens not required for further analysis or transported to processing facilities.
	+ 1. Other risk management considerations
1. All DIR licences issued by the Regulator contain a number of conditions that relate to general risk management. These include conditions relating to:
* applicant suitability
* contingency plans
* identification of the persons or classes of persons covered by the licence
* reporting requirements
* access for the purpose of monitoring for compliance.
	+ - 1. Applicant suitability
1. In making a decision whether or not to issue a licence, the Regulator must have regard to the suitability of the applicant to hold a licence. Under Section 58 of the Act, matters that the Regulator must take into account include:
* any relevant convictions of the applicant
* any revocation or suspension of a relevant licence or permit held by the applicant under a law of the Commonwealth, a State or a foreign country
* the capacity of the applicant to meet the conditions of the licence.
1. On the basis of the information submitted by the applicant and records held by the OGTR, the Regulator considers Bioproperties suitable to hold a licence. The licence includes a requirement for the licence holder to inform the Regulator of any information that would affect their suitability.
2. In addition, the applicant organisation must have access to a properly constituted Institutional Biosafety Committee and be an accredited organisation under the Act.
	* + 1. Contingency Plan
3. Bioproperties is required to submit a contingency plan to the Regulator before inoculating chickens with the GMO at each Participating farm. This plan would detail measures to be undertaken in the event of unintentional release of the GMO (e.g. a spill), loss of the GMO stock, severe weather conditions at the trial areas, transmission of the GMO to poultry other than the GMO-inoculated chickens, detection of recombination between the GMO and another ILTV strain, an outbreak of ILT disease that may potentially be linked to GMO exposure, and escape, loss or predation of GMO-inoculated chickens.
4. Bioproperties is required to provide the Regulator with a method to reliably detect the GMO or the presence of the genetic modification in a recipient organism, and a written procedure for monitoring of clinical signs or symptoms of infectious laryngotracheitis in GMO-naïve chickens and testing for the unintended presence of the GMO in GMO-naïve chickens. These methods and procedures are required before dealing with the GMO.
	* + 1. Identification of the persons or classes of persons covered by the licence
5. The persons covered by the licence would be the licence holder and employees, agents or contractors of the licence holder and other persons who are, or have been, engaged or otherwise authorised by the licence holder to undertake any activity in connection with the dealings authorised by the licence. Prior to dealing with the GMO, Bioproperties is required to provide a list of people and organisations that would be covered by the licence, or the function or position where names are not known at the time.
	* + 1. Reporting requirements
6. The licence requires the licence holder to immediately report any of the following to the Regulator:
* any additional information regarding risks to the health and safety of people or the environment associated with the dealings
* any contraventions of the licence by persons covered by the licence
* any unintended effects of the release.
1. A number of written notices are also required under the licence regarding dealings with the GMO at each farm, and inoculation of each batch of broiler chickens with the GMO at each farm to assist the Regulator in designing and implementing a monitoring program for all licensed dealings. The notices would include:
* local government area, address of the farm, GPS coordinates of farm
* whether the farm is free-range
* brief description/diagram/map of the farm and any associated sheds, ranges, houses or buildings (if relevant) in the trial area and what each is used for
* location of areas used for any composting or burying of farm waste
* expected date of inoculation with the GMO
* number and age of broiler chickens to be inoculated with the GMO
* intended method of GMO administration
* identification of the particular shed/range where the GMO-inoculated chickens will be kept
* proposed processing facilities for the GMO-inoculated chickens
* expected concurrent presence of other poultry
* expected dates of harvesting the GMO-inoculated chickens for transport to the processing facilities
* expected date of decontamination of sheds that have housed the GMO-inoculated chickens.
	+ - 1. Monitoring for Compliance
1. The Act stipulates, as a condition of every licence, that a person who is authorised by the licence to deal with a GMO, and who is required to comply with a condition of the licence, must allow inspectors and other persons authorised by the Regulator to enter premises where a dealing is being undertaken for the purpose of monitoring or auditing the dealing. Post-release monitoring continues until the Regulator is satisfied that all the GMOs resulting from the authorised dealings have been removed from the release sites.
2. If monitoring activities identify changes in the risks associated with the authorised dealings, the Regulator may also vary licence conditions, or if necessary, suspend or cancel the licence.
3. In cases of non-compliance with licence conditions, the Regulator may instigate an investigation to determine the nature and extent of non-compliance. The Act provides for criminal sanctions of large fines and/or imprisonment for failing to abide by the legislation, conditions of the licence or directions from the Regulator, especially where significant damage to the health and safety of people or the environment could result.
	1. Issues to be addressed for future releases
4. Additional information has been identified that may be required to assess an application for a commercial release of the GMO, or to justify a reduction in limits and controls. This includes:
* the degree of attenuation of the GMO under field conditions
* the ability of the GMO to establish an infection cycle and persist in the environment.
	1. Conclusions of the RARMP
1. The RARMP concludes that the proposed limited and controlled release of the GMO poses negligible risks to the health and safety of people or the environment as a result of gene technology. These negligible risks do not require specific risk treatment measures.
2. However, conditions have been imposed to limit the release to the proposed scale, location and duration, and to restrict the spread and persistence of the GMO and its genetic material in the environment, as these were important considerations in establishing the context for assessing the risks.

References

ABRS (Accessed:26-4-2017) Australian Biological Resources Study Australian Faunal Directory. [ABRS](http://www.environment.gov.au/biodiversity/abrs/online-resources/fauna/afd/index.html).

Agnew-Crumpton, R., Vaz, P.K., Devlin, J.M., O'Rourke, D., Blacker-Smith, H.P., Konsak-Ilievski, B. et al. (2016) Spread of the newly emerging infectious laryngotracheitis viruses in Australia. *Infect Genet Evol* **43**: 67-73.

Animal Health Australia, Department of Agriculture, F.a.F.D. (2012) *The Australian Animal Welfare Standards and Guidelines for the Land Transport of Livestock.*

Australian Chicken Meat Federation Inc. National Farm Biosecurity Manual for Chicken Growers. [ACMF a](http://www.chicken.org.au/page.php?id=175#4).

Australian Chicken Meat Federation Inc. (Accessed:5-4-2017a) Growing Meat Chickens. [ACMF b](http://www.chicken.org.au/page.php?id=6).

Australian Chicken Meat Federation Inc. (Accessed:5-4-2017b) Structure and Ownership. [ACMF c](http://www.chicken.org.au/page.php?id=2).

Australian Renderers Association Inc. (2014) *A pocket guide to Australian rendered products.*

Avgousti, D.C., Weitzman, M.D. (2015) Stress Flips a Chromatin Switch to Wake Up Latent Virus. *Cell Host Microbe* **18**: 639-641.

Bagust, T.J. (1986) Laryngotracheitis (Gallid-1) herpesvirus infection in the chicken. 4. Latency establishment by wild and vaccine strains of ILT virus. *Avian Pathol* **15**: 581-595.

Bagust, T. J., Jones, R. C., and Guy, J. S. (2000) Avian infectious laryngotracheitis. *Revue scientifique et technique (International Office of Epizootics)* **19**: 483-492.

Bammer, G., Smithson, M. (2008) *Uncertainty and risk: Multidisciplinary perspectives.* Bammer, G., Smithson, M., eds. Earthscan, London.

Beach, J.R. (1931) A BACTERIOLOGICAL STUDY OF INFECTIOUS LARYNGOTRACHEITIS OF CHICKENS. *Journal of Experimental Medicine* **54**: 801-808.

Biosecurity Emergency Preparedness Working Group (2012) *Biosecurity Incident Management System.*

Blacker, H.P., Kirkpatrick, N.C., Rubite, A., O'Rourke, D., Noormohammadi, A.H. (2011) Epidemiology of recent outbreaks of infectious laryngotracheitis in poultry in Australia. *Australian Veterinary Journal* **89**: 89-94.

Bloom, D.C., Giordani, N.V., Kwiatkowski, D.L. (2010) Epigenetic regulation of latent HSV-1 gene expression. *Biochimica et Biophysica Acta* **1799**: 246-256.

Bryant, N.A., Davis-Poynter, N., Vanderplasschen, A., Alcami, A. (2003) Glycoprotein G isoforms from some alphaherpesviruses function as broad-spectrum chemokine binding proteins. *EMBO Journal* **22**: 833-846.

Bull, J.J. (2015) Evolutionary reversion of live viral vaccines: Can genetic engineering subdue it? *Virus Evol* **1**:

Chin, R.P., Garcia, M., Corsiglia, C., Riblet, S., Crespo, R., Shivaprasad, H.L. et al. (2009) Intervention strategies for laryngotracheitis: impact of extended downtime and enhanced biosecurity auditing. *Avian Diseases* **53**: 574-577.

Clark, A.J. and Brinkley, T. (2001) Risk management: for climate, agriculture and policy. Commonwealth of Australia, Canberra.

Coppo, M.J., Noormohammadi, A.H., Hartley, C.A., Gilkerson, J.R., Browning, G.F., Devlin, J.M. (2011) Comparative in vivo safety and efficacy of a glycoprotein G-deficient candidate vaccine strain of infectious laryngotracheitis virus delivered via eye drop. *Avian Pathol* **40**: 411-417.

Coppo, M. J. C., Noormohammadi, A. H., Browning, G. F., and Devlin, J. M. (2013) Challenges and recent advancements in infectious laryngotracheitis virus vaccines. *Avian Pathology* **42**: 195-205.

Costes, B., Ruiz-Arguello, M.B., Bryant, N.A., Alcami, A., Vanderplasschen, A. (2005) Both soluble and membrane-anchored forms of Felid herpesvirus 1 glycoprotein G function as a broad-spectrum chemokine-binding protein. *Journal of General Virology* **86**: 3209-3214.

Crawshaw, G. J. and Boycott, B. R. (1982) Infectious Laryngotracheitis in Peafowl and Pheasants. *Avian Diseases* **26**: 397-401.

Davison, A.J., Clements, B. (2009) Chapter 25: Herpesviruses: general properties. In: *Topley & Wilson's Microbiology & Microbial Infections - Virology*, 10 Edition, Volume 1, Mahy, B., ter Meulen V., eds . John Wiley & Sons. 488-505.

Department of Agriculture, F.a.F.D. (2009) *National Farm Biosecurity Manual Poultry Production.*

Devlin, J.M., Browning, G.F., Hartley, C.A., Gilkerson, J.R. (2007) Glycoprotein G deficient infectious laryngotracheitis virus is a candidate attenuated vaccine. *Vaccine* **25**: 3561-3566.

Devlin, J.M., Browning, G.F., Hartley, C.A., Kirkpatrick, N.C., Mahmoudian, A., Noormohammadi, A.H. et al. (2006) Glycoprotein G is a virulence factor in infectious laryngotracheitis virus. *Journal of General Virology* **87**: 2839-2847.

Devlin, J.M., Hartley, C.A., Gilkerson, J.R., Coppo, M.J., Vaz, P., Noormohammadi, A.H. et al. (2011) Horizontal transmission dynamics of a glycoprotein G deficient candidate vaccine strain of infectious laryngotracheitis virus and the effect of vaccination on transmission of virulent virus. *Vaccine* **29**: 5699-5704.

FSANZ (2010) Final Assessment Report - Proposal P282 - Primary Production & Processing Standard for Poultry Meat.

Fuchs, W., Veits, J., Helferich, D., Granzow, H., Teifke, J.P., Mettenleiter, T.C. (2007) Molecular biology of avian infectious laryngotracheitis virus. *Vet Res* **38**: 261-279.

Garcia, M., Spatz, S., Guy, J.S. (2013) Chapter 5: Infectious laryngotracheitis. In: *Diseases of Poultry 13th edition*, Swayne, D.E., Glisson J.R., McDougald L.R., Nolan L.K., Suarez D.L., Nair V.L., eds . Wiley Blackwell.

Giambrone, J. J., Fagbohun, O., and Macklin, K. S. (2008) Management Practices to Reduce Infectious Laryngotracheitis Virus in Poultry Litter. *Journal of Applied Poultry Research* **17**: 64-68.

Hamm, R.L., Kaufman, P.E., Reasor, C.A., Rutz, D.A., Scott, J.G. (2006) Resistance to cyfluthrin and tetrachlorvinphos in the lesser mealworm, Alphitobius diaperinus, collected from the eastern United States. *Pest Manag Sci* **62**: 673-677.

Hammond, B., Kough, J., Herouet-Guicheney, C., Jez, J.M. (2013) Toxicological evaluation of proteins introduced into food crops. *Crit Rev Toxicol* **43 Suppl 2**: 25-42.

Hanley, K.A. (2011) The double-edged sword: How evolution can make or break a live-attenuated virus vaccine. *Evolution (N Y )* **4**: 635-643.

Hayes, K.R. (2004) Ecological implications of GMOs: robust methodologies for ecological risk assessment. Best practice and current practice in ecological risk assessment for genetically modified organisms. CSIRO Division of Marine Research, Tasmania.

Hidalgo, H. (2004) Infectious Laryngotracheitis: A Review. *Brazilian Journal of Poultry Science* **5**: 157-168.

Huang, J., Hartley, C.A., Ficorilli, N.P., Crabb, B.S., Studdert, M.J. (2005) Glycoprotein G deletion mutants of equine herpesvirus 1 (EHV1; equine abortion virus) and EHV4 (equine rhinopneumonitis virus). *Archives of Virology* **150**: 2583-2592.

Huang, W., Xie, P., Mingming, X., Li, P., and Zao, G. (2011) The Influence of Stress Factors on the Reactivation of Latent Herpes Simplex Virus Type 1 in Infected Mice. *Cell Biochemistry and Biophysics* **61**: 115-122.

Huby, R.D., Dearman, R.J., Kimber, I. (2000) Why are some proteins allergens? *Toxicological Sciences* **55**: 235-246.

Hughes, C.S., Gaskell, R.M., Jones, R.C., Bradbury, J.M., Jordan, F.T. (1989) Effects of certain stress factors on the re-excretion of infectious laryngotracheitis virus from latently infected carrier birds. *Res Vet Sci* **46**: 274-276.

Hughes, C.S., Williams, R.A., Gaskell, R.M., Jordan, F.T., Bradbury, J.M., Bennett, M. et al. (1991) Latency and reactivation of infectious laryngotracheitis vaccine virus. *Archives of Virology* **121**: 213-218.

Jimenez-Guardeno, J.M., Regla-Nava, J.A., Nieto-Torres, J.L., DeDiego, M.L., Castano-Rodriguez, C., Fernandez-Delgado, R. et al. (2015) Identification of the Mechanisms Causing Reversion to Virulence in an Attenuated SARS-CoV for the Design of a Genetically Stable Vaccine. *PLoS Pathog* **11**: e1005215.

Johnson, M.A., Prideaux, C.T., Kongsuwan, K., Sheppard, M., Fahey, K.J. (1991) Gallid herpesvirus 1 (infectious laryngotracheitis virus): cloning and physical maps of the SA-2 strain. *Archives of Virology* **119**: 181-198.

Jordan, F. T. W. (1966) A Review of the Literature on Infectious Laryngotracheitis (ILT). *Avian Diseases* **10**: 1-26.

Kirkpatrick, N.C., Mahmoudian, A., O'Rourke, D., Noormohammadi, A.H. (2006) Differentiation of infectious laryngotracheitis virus isolates by restriction fragment length polymorphic analysis of polymerase chain reaction products amplified from multiple genes. *Avian Diseases* **50**: 28-34.

Lee, S.-W., Markham, P.F., Coppo, M.J.C., Legione, A.R., Markham, J.F., Noormohammadi, A.H. et al. (2012) Attenuated Vaccines Can Recombine to Form Virulent Field Viruses. *Science* **337**: 188.

Lee, S.W., Devlin, J.M., Markham, J.F., Noormohammadi, A.H., Browning, G.F., Ficorilli, N.P. et al. (2011a) Comparative analysis of the complete genome sequences of two Australian origin live attenuated vaccines of infectious laryngotracheitis virus. *Vaccine* **29**: 9583-9587.

Lee, S.W., Devlin, J.M., Markham, J.F., Noormohammadi, A.H., Browning, G.F., Ficorilli, N.P. et al. (2013) Phylogenetic and molecular epidemiological studies reveal evidence of multiple past recombination events between infectious laryngotracheitis viruses. *PLoS One* **8**: e55121.

Lee, S.W., Markham, P.F., Markham, J.F., Petermann, I., Noormohammadi, A.H., Browning, G.F. et al. (2011b) First complete genome sequence of infectious laryngotracheitis virus. *BMC Genomics* **12**: 197.

Linares, J.A., Bickford, A.A., Cooper, G.L., Charlton, B.R., Woolcock, P.R. (1994) An outbreak of infectious laryngotracheitis in California broilers. *Avian Diseases* **38**: 188-192.

Menendez, K. R., García, M., Spatz, S., and Tablante, N. L. (2014) Molecular epidemiology of infectious laryngotracheitis: a review. *Avian Pathology* **43**: 108-117.

Moreno, A., Piccirillo, A., Mondin, A., Morandini, E., Gavazzi, L., Cordioli, P. (2010) Epidemic of infectious laryngotracheitis in Italy: characterization of virus isolates by PCR-restriction fragment length polymorphism and sequence analysis. *Avian Diseases* **54**: 1172-1177.

Nakamichi, K., Kuroki, D., Matsumoto, Y., Otsuka, H. (2001) Bovine herpesvirus 1 glycoprotein G is required for prevention of apoptosis and efficient viral growth in rabbit kidney cells. *Virology* **279**: 488-498.

Nakamichi, K., Matsumoto, Y., Otsuka, H. (2002) Bovine herpesvirus 1 glycoprotein G is necessary for maintaining cell-to-cell junctional adherence among infected cells. *Virology* **294**: 22-30.

National Center for Biotechnology Information (NCBI) BLAST - Basic Local Alignment Search Tool. https://blast.ncbi.nlm.nih.gov/.

NSW Department of Environment and Conservation (2004) *Environmental Guidelines Composting and Related Organics Facilities.*

NSW Department of Primary Industries (2007) *NSW biosecurity guidelines for free range poultry farms.*

NSW Department of Primary Industries (2012) *Best Practice Management for Meat Chicken Production in NSW - Manual 1 and 2.*

NSW Department of Primary Industries Animal Health Surveillance January-March 2016. [NSW DPI](http://www.dpi.nsw.gov.au/__data/assets/pdf_file/0005/656465/ahs-q1-2016.pdf).

NSW Environment Protection Authority (2003) *Industry Sector: Livestock Processing Industries (Animal Slaughter and Rendering) Compliance Performance Report, September 2003.*

NSW Environment Protection Authority Dead stock disposal. [NSW EPA](http://www.epa.nsw.gov.au/mao/deadstockdisposal.htm).

NSW Environment Protection Authority (2016) *Environmental Guidelines Solid waste landfills.*

OGTR (2013) Risk Analysis Framework. Report No: Version 4, Document produced by the Australian Government Office of the Gene Technology Regulator.

Oldoni, I., Rodriguez-Avila, A., Riblet, S.M., Zavala, G., Garcia, M. (2009) Pathogenicity and growth characteristics of selected infectious laryngotracheitis virus strains from the United States. *Avian Pathol* **38**: 47-53.

Ou, S., Giambrone, J. J., and Macklin, K. S. (2011) Infectious laryngotracheitis vaccine virus detection in water lines and effectiveness of sanitizers for inactivating the virus. *Journal of Applied Poultry Research* **20**: 223-230.

Ou, S.C., Giambrone, J.J. (2012) Infectious laryngotracheitis virus in chickens. *World J Virol* **1**: 142-149.

Ou, S. C., Giambrone, J. J., and Macklin, K. S. (2012) Detection of infectious laryngotracheitis virus from darkling beetles and their immature stage (lesser mealworms) by quantitative polymerase chain reaction and virus isolation. *Journal of Applied Poultry Research* **21**: 33-38.

Portz, C., Beltrao, N., Furian, T.Q., Junior, A.B., Macagnan, M., Griebeler, J. et al. (2008) Natural infection of turkeys by infectious laryngotracheitis virus. *Veterinary Microbiology* **131**: 57-64.

Poultry Hub Darkling beetles. [Poultry Hub](http://www.poultryhub.org/production/husbandry-management/housing-environment/pest-management/darkling-beetles/).

Prideaux, C.T., Kongsuwan, K., Johnson, M.A., Sheppard, M., Fahey, K.J. (1992) Infectious laryngotracheitis virus growth, DNA replication, and protein synthesis. *Archives of Virology* **123**: 181-192.

Rodriguez-Avila, A., Oldoni, I., Riblet, S., Garcia, M. (2007) Replication and transmission of live attenuated infectious laryngotracheitis virus (ILTV) vaccines. *Avian Diseases* **51**: 905-911.

Roy, P., Fakhrul Islam, A.F., Burgess, S.K., Hunt, P.W., McNally, J., Walkden-Brown, S.W. (2015) Real-time PCR quantification of infectious laryngotracheitis virus in chicken tissues, faeces, isolator-dust and bedding material over 28 days following infection reveals high levels in faeces and dust. *Journal of General Virology* **96**: 3338-3347.

Sawtell, N.M., Thompson, R.L. (1992) Rapid in vivo reactivation of herpes simplex virus in latently infected murine ganglionic neurons after transient hyperthermia. *Journal of Virology* **66**: 2150-2156.

Sellers, H.S., Garcia, M., Glisson, J.R., Brown, T.P., Sander, J.S., Guy, J.S. (2004) Mild infectious laryngotracheitis in broilers in the southeast. *Avian Diseases* **48**: 430-436.

South Australia Department of Environment, Water and Natural Resources Pest Animals. [SA DEWNR](http://www.naturalresources.sa.gov.au/kangarooisland/plants-and-animals/pest-animals).

Standards Australia (2007) *Hygienic Rendering of Animal Products AS 5008:2007.*

Standards Australia (2012) *Composts, soil conditioners and mulches AS 4454:2012.*

Thureen, D. R. and Keeler, C. L. (2006) Psittacid Herpesvirus 1 and Infectious Laryngotracheitis Virus: Comparative Genome Sequence Analysis of Two Avian Alphaherpesviruses. *Journal of Virology* **80**: 7863-7872.

Tran, L.C., Kissner, J.M., Westerman, L.E., Sears, A.E. (2000) A herpes simplex virus 1 recombinant lacking the glycoprotein G coding sequences is defective in entry through apical surfaces of polarized epithelial cells in culture and in vivo. *Proc Natl Acad Sci U S A* **97**: 1818-1822.

VICH (2000) Good Clinical Practice.

Victoria Department of Economic Development, J.T.a.R.D. (2009) *Victorian Code for Broiler Farms 2009.*

Victoria Department of Economic Development, Jobs Transport and Resources DEDJTR Biosecurity guidelines for poultry producers. [Agriculture Victoria a](http://agriculture.vic.gov.au/agriculture/pests-diseases-and-weeds/animal-diseases/poultry/biosecurity-guidelines-for-poultry-producers).

Victoria Department of Economic Development, Jobs Transport and Resources DEDJTR Disposing of carcasses after bushfire, flood or drought. [Agriculture Victoria b](http://agriculture.vic.gov.au/agriculture/emergencies/recovery/livestock-after-an-emergency/disposing-of-carcasses-after-bushfire-flood-or-drought).

Victoria Department of Economic Development, Jobs Transport and Resources DEDJTR VetWatch. [Agriculture Victoria c](http://agriculture.vic.gov.au/agriculture/pests-diseases-and-weeds/animal-diseases/vetsource-information-for-vets/vetwatch).

Victoria Environment Protection Authority (2015) *Designing, constructing and operating composting facilities.*

Victoria Environment Protection Authority (2016) *Landfill licensing.*

Volkova, V., Thornton, D., Hubbard, S.A., Magee, D., Cummings, T., Luna, L. et al. (2012) Factors associated with introduction of infectious laryngotracheitis virus on broiler farms during a localized outbreak. *Avian Diseases* **56**: 521-528.

Williams, R.A., Bennett, M., Bradbury, J.M., Gaskell, R.M., Jones, R.C., Jordan, F.T. (1992) Demonstration of sites of latency of infectious laryngotracheitis virus using the polymerase chain reaction. *Journal of General Virology* **73 ( Pt 9)**: 2415-2420.

Wilson, A.C., Mohr, I. (2012) A cultured affair: HSV latency and reactivation in neurons. *Trends in Microbiology* **20**: 604-611.

Yamada, S., Matsuo, K., Fukuda, T., Uchinuno, Y. (1980) Susceptibility of ducks to the virus of infectious laryngotracheitis. *Avian Diseases* **24**: 930-938.

**Appendix A**

Advice received by the Regulator from prescribed experts, agencies and authorities on the consultation RARMP is summarised below. All issues raised in submissions that related to risks to the health and safety of people and the environment were considered in the context of the currently available scientific evidence and were used in finalising the RARMP that formed the basis of the Regulator’s decision to issue the licence.

**Abbreviations: Ch:** chapter; **CMP:** Compliance Management Plan; **gG:** glycoprotein G; **GMO:** genetically modified organism; **ILT:** infectious laryngotracheitis; **ILTV:** infectious laryngotracheitis virus; **m**: metres; **OGTR**: Office of the Gene Technology Regulator; **PPP**: Primary Production and Processing **RARMP:** Risk Assessment and Risk Management Plan; **Sec:** Section; **Sub. No.**: submission number**.**

| **Sub. No.** | **Summary of issues raised** | **Comment** |
| --- | --- | --- |
| 1 | Agrees with the overall conclusions of the RARMP. | Noted. |
| The Regulator should further consider potential consequences of infection in other birds. | A new Risk Scenario (#3) has been added that considers this as a potential harm to wild birds (Ch.2 Sec. 2.1.3). |
| The Regulator should further consider trial locations where birds are able to be appropriately contained during the virus shedding period. | Licence conditions have been added that requires GMO-inoculated chickens to be kept inside the shed while they display clinical signs or symptoms of ILT disease and they must not be harvested until the disease signs or symptoms disappear. An additional licence condition has also been imposed requiring the Trial Areas to be at least 1000 m from poultry located on other poultry farms. |
| 3 | Does not have specialist scientific expert to make an assessment. No comment provided. | Noted. |
| 2 | The risk assessment conclusions and risk management plan are appropriate to mitigate any risk to public health and safety from the food supply. | Noted. |
| 4 | Does not have specialist scientific expert on the matter. Would like to be kept informed of the location of any trial sites within our local government area and would like further information in regard to any site specific controls for these trial sites. | The licence specifies the controls for the trial. The locations of each trial site will be made available on the [OGTR website](http://www.ogtr.gov.au/internet/ogtr/publishing.nsf/Content/home-1). |
| 5 | Agrees with the overall conclusions of the consultation RARMP that given the limits and control measures proposed, the risks to the environment are negligible. | Noted. |
| The consultation RARMP notes that the ILTV has a narrow host range, being restricted to the family *Phasianidae*, and there are three native Australian species in this family. The RARMP would benefit from giving scientific names of these three species. | The scientific names for the three Australian species have been added in Ch.1 Sec.7.6. |
| The GMO contains an intact translation start codon that could probably be translated to a small non-functional protein. The RARMP would benefit from a brief assessment of the possibility that this novel protein could produce any adverse effects such as toxicity in a host. | The suggested reason for why the translated non-functional protein is unlikely to have a toxic effect has been added in Risk Scenario 1 (Ch.2 Sec. 2.1.1). |
| It was suggested that deletions in the genome are more genetically and phenotypically stable than point mutations, and that resultant viruses containing the deletion would be non-virulent. Nevertheless, viruses, which have regained the previously deleted gene, have been isolated in laboratory experiments. | Risk scenario 5 discusses the effects of recombination between the GMO and another ILTV strain. The suggested information has been added in Risk Scenario 5 (Ch.2 Sec. 2.1.5). |
| 6 | No substantial objection to the licence application at this stage. | Noted. |
| However, it is noted that the consultation RARMP highlights the risk associated with the use of multiple attenuated ILT vaccinations which may trigger recombination of ILTV and result in a more virulent or transmissible progeny. Specific points of clarification was requested:* the disease causing capacity of the parental strain of GMO
* the consequence of restoring the gG gene function into the GMO
* effect of the deletion on the replication capacity of GMO
* any experiments to assess the risk of this GM vaccine undergoing genetic recombination with live ILTV vaccines currently used in Australia and circulating strains
* shedding of the GMO by inoculated birds and/or transmit it to other chickens or avian species.
 | ILTV’s ability to undergo recombination to create new classes of ILTV in the field more virulent than the wild type ILTV, including the recent study by Lee et al 2012 referred to in the submission, has been discussed in Ch.1 Sec.5.8 and 5.9.Responses to the questions raised are clarified below:* The parental strain used in the GM vaccine is CSW-1 which was originally isolated from a NSW outbreak in 1959. Recent outbreaks in Australia have not been caused by CSW-1 strain and appear to be no longer circulating (see Ch.1, Sec 5.9). The CSW-1 strain has since been studied in several laboratory experiments. Compared to the GM vaccine, the CSW-1 strain is more virulent, causing higher mortality rates than the GM vaccine (see Ch.1 Sec.6.3).
* It is expected that restoration of the gG gene in a virus would restore the virulence/pathogenicity similar or less than the parent strains (discussed in Ch.2 Sec. 2.1.5, Risk Scenario 5).
* Replication and growth of the GMO *in vitro* and *in vivo* have been characterised and shown to be similar to CSW-1 (discussed in Ch.1 Sec. 6.3).
* No experiments have been conducted to study the ability of the GMO to undergo recombination with another ILTV wild strain or vaccine strain. Recombination would be studied in the proposed field trials.
* Shedding and transmission of the GMO have been studied and shown to be transmissible to naïve chickens. Peak shedding of ILTV strains generally occurs 4-9 days post-inoculation, and GMO DNA can be detected from tracheal swabs up to 21 days in some chickens (discussed in Ch.1 Sec. 5 and 6).
 |
| 7 | Has some concerns and considerations related to biosecurity.RARMP suggests limiting administration to 2 million birds over a 5 year period in a restricted area will provide containment of the GMO. Concerned that the level of delivery may lead to establishment in the environment. | The risk of persistence and establishment of the GMO in the environment is considered negligible in Risk Scenario 6. Additional licences conditions have been included in the final licence to minimise spread and persistence, including confinement to sheds immediately following inoculation or if showing disease symptoms, isolating trial areas at least 1 km from poultry located on other poultry farms and testing for the GMO in chickens at the sites following completion of the trials. |
| Concerned that there may not be appropriate controls over movements of live birds and transport routes to processing plants given that this has been a suspected mechanism for spreading virulent virus. These viruses including current vaccines have been shown to recombine and generate virulent strains. Is the OGTR confident that these risks are appropriately managed? | The risk of spreading the GMO to susceptible birds that may lead to recombination with circulating ILTV strains after transport to processing plants is discussed in Risk Scenario 5 (Ch.2 Sec. 2.1.5). The reasons that the risk of recombination between the GMO and another ILTV strain is considered to be negligible include the GMO contains a gene deletion and resultant viruses containing the gene deletion would not be more virulent than the parent strain. Additional licence conditions have been imposed prohibiting the transport of chickens to processing plants immediately following vaccination or if displaying disease symptoms. |
| Infectious laryngotracheitis is a notifiable disease under state legislation, and there are legislative requirements under its use in research and development. The OGTR could consider ways to inform animal health authorities to ensure requirements are being met. It is important to note that dead poultry is considered a Restricted Animal Material and a number of state regulations apply to their disposal. | The state legislation and requirements including the notifiable disease requirements for ILT disease are discussed in the RARMP. Part of the CMP includes disposal of carcasses and licence conditions are imposed to notify the Regulator in the event of an ILT outbreak. |
| 8 | Support the OGTR’s conclusion that DIR 154 poses negligible risk of harm to human health and safety and the environment. | Noted. |
| It was not clear how the OGTR will ensure that the applicant demonstrates compliance with a range of state and local requirements and guidelines (discussed in Ch.1 Sec.7). | The state agencies, local councils, the licence holder, poultry farmer and poultry company all have a responsibility to ensure compliance with the relevant legislation and requirements.OGTR regularly monitors field trial sites to ensure compliance with licence conditions. Penalties may apply to non-compliance with licence conditions. |
| 9 | Whether the Standard 1.5.2 of Australia New Zealand Food Standards Code – Food produced using gene technology – has been taken into consideration in the assessment. | According to Standard 1.5.2, food produced using gene technology means a food which has been derived or developed from an organism which has been modified by gene technology. The broiler chickens appear to not be captured by Standard 1.5.2 because the chickens have not been modified by gene technology, but instead have been inoculated with a GM vaccine. The GM vaccine does not genetically modify any cells in the chicken. |
| It is noted that chicken meat processing practices would minimise any residual GMO in the chicken meat or product derived from treated chickens, and that any trace amount would not survive cooking (Ch2. Sec.2.1.2). It is unclear whether any novel DNA will remain detectable either in or on the meat or other type of derived product. | As discussed in Ch.1 Sec.6.1 and Ch.2 Risk Scenario 1, the GMO does not contain any new genetic material and the sequence is highly similar to the parent ILTV strain, with one gene deleted. In the unlikely event that DNA from the GMO were present in or on raw meat products, the risk to health and safety of people would be negligibleFurthermore, as noted, the GMO, including its DNA and proteins, is expected to not survive the cooking process. |
| Concern about whether consideration has been given to issues beyond food safety to include those which fall within the broader remit of the Australia New Zealand Food Standards Code. | In the context of the proposed field trials and the how the chickens would be consumed after slaughter, the RARMP has considered, as the most relevant, the requirements of Standard 4.2.2 – Primary Production and Processing (PPP) Standard for Poultry Meat which is aimed at ensuring the safety of poultry meat for human consumption. |

1. APVMA permit number PER81178, in force from 11 March 2016 to 30 June 2021. [↑](#footnote-ref-2)
2. (Genbank accession number: JX646899.1) [↑](#footnote-ref-3)
3. (Genbank accession number: HQ630064) [↑](#footnote-ref-4)
4. (Genbank accession number: JN596962.1) [↑](#footnote-ref-5)
5. (Genbank accession number: JN596963.1) [↑](#footnote-ref-6)
6. The APVMA takes into consideration the risk of recombination from the use of the viral vaccines. The product leaflets approved by the APVMA for the use of A20 and SA2 vaccine strains do not recommend concurrently using ILTV originating from genetically distinct strains in a flock or on a site. [↑](#footnote-ref-7)
7. A more detailed discussion is contained in the Regulator’s *Risk Analysis Framework* available from the [OGTR website](http://www.ogtr.gov.au/internet/ogtr/publishing.nsf/Content/riskassessments-1) or via Free call 1800 181 030. [↑](#footnote-ref-8)