



Permit to Import Quarantine Material

Permit: IP15002681 Valid From: 14 Jan 2015 Valid To: 14 Jan 2017 Page 1 of 24

Importer	Exporter
Health Support Queensland Pathology Queensland Central Laboratory Level 4, Block 7 Royal Brisbane & Women's Hospitals Campus, Herston Road Herston QLD 4006 Attn: Matt Ford	Various Suppliers Exporters Various Addresses In All countries

You are authorised to import the following material under the listed conditions

Note: This permit covers the Department of Agriculture quarantine requirement only.

All imports may be subject to quarantine inspection on arrival to determine compliance with the listed permit conditions and freedom from contamination. Imports not in compliance or not appropriately identified or packaged and labelled in accordance with the import conditions they represent may be subject to seizure, treatment, re-export or destruction at the importer's expense.

Additionally, all foods imported into Australia must comply with the provisions of the *Imported Food Control Act 1992*, and may be inspected and/or analysed against the requirements of the Australia New Zealand Food Standards Code.

All imports containing or derived from Genetically Modified material must comply with the *Gene Technology Act 2000*.

It is the importer's responsibility to identify, and to ensure it has complied with, all requirements of any other regulatory organisations and advisory bodies prior to and after importation including The Australian Customs and Border Protection Service, The Department of Health and Ageing, Therapeutic Goods Administration, Australian Pesticides and Veterinary Medicines Authority, Department of Sustainability, Environment, Water, Population and Communities, Food Standards Australia New Zealand and any state agencies such as Departments of Agriculture and Health and Environmental Protection authorities. Importers should note that this list is not exhaustive.

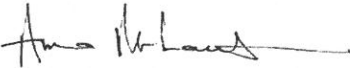

This permit is granted for the purposes of the *Quarantine Act 1908* and *Quarantine Proclamation 1998* of the Commonwealth of Australia. The laws of Australian States and Territories may also impose restrictions on the import of animals, plants and other goods into those States and Territories. This import permit does not prevent the application of those State and Territory laws. The importer should seek its own advice on any restrictions that may apply in any State or Territory into which it is proposed to import the animals, plants or other goods to which this permit relates.

Import conditions are subject to change at the discretion of the Director of Quarantine. This permit may be revoked without notice.

Notification of the import must be provided to the Department of Agriculture for all imported goods other than goods imported as accompanied baggage or goods imported via the mail and not prescribed under the *Customs Act 1901*. Notification must be consistent with *Quarantine Regulations 2000* (examples include a Quarantine Entry or a Quarantine declaration).

Commodity Name	Condition Number(s)	Country	End Use
Antibodies - Purified & raised against microorganisms and viruses as listed in PCT1104	PC0701 AND PC0992 AND PCT1104 AND PC0017	All countries	In-vitro use or in-vivo use in laboratory organisms only

This permit is granted subject to the condition that fees determined under Section 86E are paid

 Delegate of Director of Quarantine Printed Name Anna Moorhouse	Stamp: 
Date 14 Jan 2015	

Commodity Name	Condition Number(s)	Country	End Use
Antibodies - Purified & raised against microorganisms, including viruses (as listed in PC6800)	PC0017 AND PC6800 AND PC6848 AND PC0992	All countries	In-vitro use or in-vivo use in laboratory organisms only
Antigens - Purified & derived against microorganisms, including viruses (as listed in PC6800)	PC0017 AND PC6849 AND PC6800 AND PC0992	All countries	In-vitro use or in-vivo use in laboratory organisms only
Antigens - Purified & derived from microorganisms & viruses as listed in PCT1104	PC0701 AND PC0992 AND PCT1104	All countries	In-vitro use or in-vivo use in laboratory organisms only
Microorganisms, including viruses (as listed in PC6800)	PC0017 AND PC6800 AND PC6760 AND PC0691	All countries	In-vitro use or in-vivo use in laboratory organisms only
Microorganisms, including viruses (As listed in PC0600)	PC0017 AND PC0600 AND PC0691	All countries	In-vitro use or in-vivo use in laboratory organisms only
Animal fluids and tissues (excluding reproductive material) - sourced from low risk species	PC0017 AND PC6798 AND PC0992	All countries	In-vitro use or in-vivo use in laboratory organisms only
Diagnostic and Research Only Kits - not testing for microorganisms, viruses or prions - use in lab	PC0017 AND PC6282 AND PC0766	All countries	In-vitro
Human Diagnostic Tests and Human Diagnostic kits (excluding those testing for high risk human pathogens and diseases)	PC0017 AND PC6792 AND PC6369 AND PC6806	All countries	In-vitro
Cell lines and/or supernatant fluid - derived from laboratory animals & humans	PC0017 AND PC5887 AND PC6021	All countries	In-vitro use or in-vivo use in laboratory organisms only
Human fluids & tissues - free from listed diseases	PC4206	All countries	In-vitro use or in-vivo use in laboratory organisms only
Genetic material and vectors - low risk	PC0017 AND PC6797 AND PC5887	All countries	In-vitro use or in-vivo use in laboratory organisms only
Blood collection tubes (empty) - all biological additives	PC0017 AND PC0766	All countries	In-vitro
Purified / refined laboratory reagents - low risk laboratory material	PC0017 AND PC6799 AND PC0992 AND PC0701	All countries	In-vitro use or in-vivo use in laboratory organisms only

Commodity Name	Condition Number(s)	Country	End Use
Laboratory reagent (rabbit complement (greater than 20ml))	PC0017 AND PC0992 AND PC6866 AND PC0675	All countries	In-vitro use or in-vivo use in laboratory organisms only
Human Fluids and Tissues - not known to be infected (Including samples on microscope slides and samples dried onto filter paper, dip sticks or swabs)	PC0017 AND PC6743	All countries	In-vitro use or in-vivo use in laboratory organisms only

Condition	Condition Text
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PC0017 **Biological Imports Program (BIP) - Administrative conditions**

1. This import permit (or number) and all required documentation must accompany each consignment and must be valid at the time the cargo is landed.
2. In order to facilitate clearance of mail shipments, the import permit (or number) and all documentation should be securely attached to the outside of the package and marked 'Attention Quarantine'.
3. The importer must meet all costs associated with the import of this product.
4. The importer (or agent) must lodge a quarantine entry for each consignment.
5. Documents must be provided with each consignment which:
 - a) identify the consignment e.g. entry number; and
 - b) identify all goods being imported as part of this consignment e.g. invoice or waybill or importers manifest; and
 - c) describe the goods being imported (where not clear) Example 1: Product XRab = Purified protein derived from rabbits. Example 2: Product AX = Synthetic antibiotic. Example 3: Comte = Cheese.

Note: It is the importer's responsibility to provide any additional information which is requested in order to demonstrate that the import permit covers all goods being imported.
6. Consignments that do not meet the import conditions will remain under the Department's control pending export or destruction at the importers expense.
7. For further information please contact:

Regional - Clearance assistance: <http://www.daff.gov.au/biosecurity/about/contact/regional>

Canberra - Biological Import Program - Administrative assistance:
bioadmin@agriculture.gov.au

Canberra - Biological Import Program - Technical assistance: biologicals@agriculture.gov.au

PC0600 Nocardia spp. (excluding N. Nocardia crassostreae, kampachi, seriolae and salmonicida and vaccinii)
 Burkholderia cepacia complex

Condition	Condition Text
PC0675	Manufacturer's declaration

Certification

Each consignment must be accompanied by a manufacturer's declaration.
The declaration must meet the format described by this permit and state:

- a) The sera was only derived from animals with no history of or clinical signs of infectious disease, and;
- b) The sera was only derived from SPF animals maintained under veterinary supervision.

The declaration must be provided by:

The manufacturer

PC0691 **Packaging Requirements**

1. Cultures must be pure cultures (unless otherwise specified by this Import Permit) and labelled with the scientific name of the organism as it appears on this Import Permit.

Post Entry Requirements

- 2. This Import Permit allows for the importation of goods for in vitro laboratory studies (or in vivo use in laboratory organisms only), unless approved by the Department of Agriculture for specific in vivo use in non-laboratory organisms.
- 3. Laboratory organisms are those defined in the following list and must be contained under laboratory or animal house conditions: guinea pigs, hamsters, mice, rabbits, rats or micro-organisms. Work in all other animals and plants is not permitted.
- 4. For in vivo use in non-laboratory organisms (e.g. chickens, sheep, cattle, etc.) or plants a separate application for in vivo use must be lodged with, and approved by the Department of Agriculture. This also applies if the product is to be used in veterinary vaccine or veterinary therapeutic manufacture.
- 5. It is the end user's responsibility to ensure that all laboratory products are used in accordance with the current AS/NZS 2243 Safety in Laboratory standards. This includes handling and disposal procedures.
- 6. It is the importer's responsibility to ensure compliance with all international (e.g. IATA) and domestic requirements concerning the safe handling, transport and labelling of biological material.
- 7. It is the end user's responsibility to ensure that all laboratory products are used in accordance with the Office of the Gene Technology Regulator (OGTR) requirements.