MINISTRY FOR PRIMARY INDUSTRIES (MPI) STANDARD TREATMENT SUPPLIER PROGRAMME

Requirements for the supplier of official treatments

<table>
<thead>
<tr>
<th>REVIEW</th>
<th>This MPI standard is subject to periodic review.</th>
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<tbody>
<tr>
<td>ENDORSEMENT</td>
<td>This MPI standard is hereby endorsed.</td>
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<tr>
<td>Director Plant, Food, and Environment MPI</td>
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AMENDMENT RECORD and IMPLEMENTATION SCHEDULE

Amendments to this Standard originally issued on 1 June 2006 will be given a consecutive number and will be dated.

Please ensure that all amendments are inserted, obsolete pages removed, and the record below is completed.

<table>
<thead>
<tr>
<th>Amendment No:</th>
<th>Date:</th>
<th>Specification:</th>
<th>Implementation Date:</th>
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<tr>
<td>1</td>
<td>01/07/13</td>
<td>Update of Ministry of Agriculture and Forestry (MAF) to Ministry for Primary Industries (MPI)</td>
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<td>6</td>
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</tbody>
</table>
REVIEW

This MPI standard is subject to review and amendment at any time to ensure that it continues to meet current needs. The current version of the Standard will be available on the MPI web site at:


Last Review: May 2006

CONTACT

Ken Glassey
Senior Adviser
Biosecurity & Environment
Ministry for Primary Industries

Implementation Schedule

See transition schedule
1.0 INTRODUCTION

1.1 Background

This standard sets out requirements that treatment suppliers seeking approval to operate under a quality management system and a programme of surveillance without direct IVA supervision for all treatments must follow.

Treatment suppliers are required to demonstrate their ability to consistently provide product and services that meet applicable regulatory requirements by operating a quality management system. Treatment suppliers shall develop and implement a quality management system in compliance with this standard, and apply to an IVA for audit and to MPI for approval.

The requirements for a treatment supplier’s quality management system set out are based upon the principles of ISO 9000 series of quality management system standards, but recognise that small businesses may require less formal structures and systems.

When an IVA is satisfied that all the requirements of this standard are met, the IVA will recommend to MPI that the treatment supplier be approved to undertake those services within the scope of the treatment supplier’s system. MPI will consider that recommendation, and if appropriate will accept it and approve the treatment supplier. Once approved, whenever a treatment is carried out, the approved treatment supplier can carry out treatments without direct supervision by an IVA, and may issue its own treatment certificates.

When the treatment suppliers system has been submitted to their chosen IVA and passed a desk audit they shall operate under a heightened audit regime while going through the process of gaining approval.

All treatment suppliers are subject to legislation administered by other government departments; this standard does not cover those responsibilities eg HSNO and Health and safety.

1.2 Scope

This standard specifies the requirements to be met by a treatment supplier to become approved to carry out treatments:

i. As directed by MPI for imported risk goods;

ii. As required to meet export certification requirements (for the inclusion as an additional declaration on phytosanitary certificates);

iii. As required to treat and apply the ISPM 15 mark to wood packaging material that is in accordance with the international ISPM 15 standard. Reference should be made to MPI export certification standard: Technical Requirements: Registered Certification Mark (ISPM 15) to comply with this standard.
Treatment Suppliers may elect to operate under supervision, or seek approval under this standard.

1.3 References

a) Biosecurity Act 1993 (as amended by the Biosecurity Amendment Act (1997)
b) Hazardous Substances and New Organisms Act 1996
c) Health Act 1956
e) AS/NZS ISO 9000: 2000 Quality management systems – Fundamentals and vocabulary
g) ISO 17020: 1998: General criteria for the operation of various types of bodies performing inspection.
h) ISO 19011:2003 Guidelines for quality and/or environmental management systems auditing
i) Ministry for Primary Industries New Zealand Export Certification Standard: Requirements to be met by an Independent Verification Agency (IVA) to gain authorisation to undertake import and plant export certification activities
j) Ministry for Primary Industries New Zealand Standard: Approved Biosecurity Treatments
k) Technical Requirements: Registered Certification Mark (ISPM 15)
l) International Plant Protection Convention 1992
m) Export specifications ICPR

1.4 Definitions

For the purpose of this MPI standard the definitions found in Appendix 1 of the treatment supplier overview and general requirements document shall apply.

2.0 SYSTEM REQUIREMENTS

2.1 General

2.1.1 The treatment supplier shall establish, document where appropriate, implement and maintain a quality management system and continually improve its effectiveness in accordance with the requirements of this standard.

2.1.2 The treatment supplier shall:
   i. Ensure it has an IVA appointed at all times;
   ii. Identify the processes to be followed;
   iii. Ensure that both the operation and controls of the processes are effective;
   iv. Monitor, maintain and analyse the results of these processes;
   v. Maintain adequate records as evidence that the processes are effective.
2.1.3 The treatment supplier shall
   i. Appoint one of its staff members to have overall responsibility for establishment, implementation and maintenance of the treatment supplier’s quality management system.
   ii. Clearly define the responsibility and authority for this role in a written job description or similar document

Note 1 In a small business the Owner/Manager will usually fill this role.

2.1.4 Management shall review the quality management system annually to ensure its suitability and effectiveness to continue to meet the requirements of this standard. Records of the management review are to be maintained. The review shall include:
   i. Results of internal and external audits;
   ii. Communication from external parties, including complaints;
   iii. Changes in this standard, regulation or legislation and their impact on treatments;
   iv. Status of corrective and preventive actions;
   v. Recommendations for improvements.

2.1.5 Where a treatment supplier is operating over multiple locations they shall conduct at least an annual internal audit to verify that practices comply with planned activities. Where non compliances are identified, the root cause of the non compliance shall be determined, and appropriate actions taken to ensure that the cause(s) of problems are corrected [corrective action] and that potential problems are averted [preventive action]. All possible consequences that may have arisen from the root cause of a problem shall be identified, and if needed corrected.

   Records of internal audit and corrective and preventive actions shall be maintained.

2.2 Control of documents and records

2.2.1 Quality management system documentation shall include:
   i. A quality manual. The quality manual shall include procedures to address the following:
      a) Segregation of treated and untreated material;
      b) Traceability of treated material from the treatment stage through to storage and despatch;
      c) Recording of transfers or sales to other approved facilities such as manufacturers of wood packaging materials;
      d) Application of any relevant certification marks (if applicable);
      e) Maintenance of records;
      f) The training provided to staff members responsible for quality control or involved in the treatment of material to ensure understanding of the requirements for approval to treat material and/or apply any relevant certification marks;
...g) Procedures for administering the required treatment that assures that the minimum requirements are achieved;

h) Procedures for monitoring the treatment;

i) Mechanisms to detect treatment failure and the appropriate corrective actions that may be applied;

j) Calibration of monitoring or measuring equipment; and

k) Procedures for issuing treatment certificates;

l) A site plan of the facility (if permanent); and

m) An organisational structure clearly identifying the person(s) responsible for quality control activities;

n) Response to corrective action requests.

ii. Documents needed by the treatment supplier to ensure complete control of its processes;

iii. Records required by this standard.

Note 2 The amount of documentation required can vary depending on the complexity of the treatment.

Note 3 Documentation may be electronic, hard copy or any other form of medium.

A copy or copies of your quality manual or relevant procedures/work instructions shall be available for use by all employees that have a role or perform a function in the approved system.

Any alterations, amendments or corrections to the quality system or quality manual that may affect compliance with the requirements of approval shall be submitted in writing to your IVA for approval prior to their implementation. A record of approval shall be maintained by the approved facility.

2.2.2 The treatment supplier shall have a system for uniquely identifying and controlling all its documents (required under 2.2.1) to ensure that only the current editions are in use and that no unauthorised changes are made.

2.2.3 The treatment supplier shall ensure that current editions of documents (required under 2.2.1) are available to everyone who needs them.

2.2.4 Records shall be sufficient to demonstrate that all required processes have been carried out, that all treatments have been undertaken in compliance with the quality management system and the requirements of this standard. They shall be legible, stored in a manner that allows their retrieval, and protected from deterioration.

2.2.5 Records shall be maintained for a minimum of two years.

2.3 Resources

Management shall ensure that sufficient appropriate human, physical and financial resources are available to ensure that each treatment can be performed effectively.
2.4 **Contract review**

Prior to accepting an order to undertake a treatment, a treatment supplier shall review the order to ensure that
i. The correct treatment to be applied is identified; and
ii. The treatment supplier holds current approval for that treatment; and
iii. Staff who are qualified to undertake that treatment are available to perform or supervise the treatment; and
iv. The treatment supplier has the continuing capability to meet the treatment requirements
v. If there is any likelihood of damage that the owner of the goods is informed, and is given the option of continuing with the treatment or not.

2.5 **Staff competence, awareness and training**

i. The treatment supplier shall ensure that its staff are fully trained and competent for the work they do, and that treatment technicians at all times meet the personnel qualification criteria set out in Appendix III of this standard.
ii. An up to date register of treatment technicians showing the treatments that they are competent to perform shall be maintained;
iii. Where necessary staff shall be provided with written work instructions setting out how the treatment supplier requires critical jobs or tasks to be carried out;
iv. In house training of staff shall be carried out by competent treatment technicians;
v. Training by other providers may be recognised as adequate for the purposes of this standard. Recognised training courses include:
   a) Grow Safe Programme. (Application of chemicals);
   b) Individual Pest Control. Open Polytechnic of New Zealand Unit Standards;
   c) HSNO approved handler course.
vi. Records of training shall be kept and staff competence shall be regularly reviewed by management to determine whether training is required.

2.6 **Treatment materials**

i. The treatment supplier shall establish and maintain documented specifications for all treatment materials required;
ii. When purchasing materials, specifications of requirements for purchase shall be clearly stated on purchase documents;
iii. On delivery of materials, treatment suppliers shall verify that the material delivered meets purchase specifications;
iv. Safety information, handling and application procedures shall be obtained from suppliers, and manufacturer’s stated requirements regarding storage, safety and use shall be followed. If manufacturer’s stated requirements differ from MPI specifications, MPI should be notified and no treatment shall occur until the reason for the difference has been resolved by MPI;
v. Records of raw material specifications and manufacturer’s requirements shall be maintained.
2.7 Measuring and monitoring equipment

i. All measuring, test and inspection equipment used for measuring or monitoring treatment activities shall be routinely calibrated at least annually and/or verified to levels of accuracy appropriate to its use;

ii. Measuring and monitoring equipment shall be stored, transported and used in a manner that protects it from loss of accuracy;

iii. All equipment and products used in treatment processes shall be inspected before use to ensure that they meet the specification for the treatment to be applied, and are suitable for use;

iv. Should equipment calibration and/or verification show that equipment is not capable of the required accuracy of measurement, the treatment supplier shall review all measurements that have been made using that equipment since the last calibration and/or verification, and to determine what actions should be taken as a consequence of this inaccuracy. Appropriate actions, including re-treatment if required, shall be taken. Records of this review shall be kept for two years;

v. Equipment calibration records shall be maintained.

2.8 Process control

i. Where the absence of documented instructions creates a risk to effective performance of a treatment, documented procedures and instructions shall be provided;

ii. The treatment supplier shall document how the necessary checks on equipment, application and product handling will be done to ensure the treatment is successful.

2.9 Emergency response procedures

Treatment suppliers shall develop and document emergency response procedures to be followed in case of an emergency event. Staff shall be trained in those procedures and shall be aware of what actions they should take in case of an emergency event at all times. Procedures shall cover

i. What to do if personnel, auditors or the public are at risk;

ii. Actions to be taken to protect the environment;

iii. Actions to be taken to protect the product being treated;

iv. Means of notifying emergency services and, if needed, MPI.

3.0 SUBCONTRACTING

Treatment suppliers shall normally perform the treatment services for which they hold approval. Where any part of their treatment service is sub-contracted the treatment supplier shall ensure, and be able to demonstrate, that their sub-contractor’s documented procedures are either a part of their own approved system/procedures, or are independently approved by MPI.
3.1 Reporting

The Treatment supplier shall provide to their IVA the following reports, which shall contain information as stated within Table 1.

Table 1. Reporting requirements

<table>
<thead>
<tr>
<th>Topics</th>
<th>Time frame</th>
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<tbody>
<tr>
<td>Changes to their register of competent staff.</td>
<td>Within 5 working days of change</td>
</tr>
<tr>
<td>Critical non-compliance detections and corrective actions to be applied.</td>
<td>Within 5 working days after detection</td>
</tr>
<tr>
<td>Interceptions of MPI phytosanitary certified produce by importing countries or on cleared risk goods.</td>
<td>Within 5 working days of interception</td>
</tr>
<tr>
<td>Issues raised by importers or other offshore organisations identifying risks relating to export phytosanitary certification or clearance of goods.</td>
<td>Within 5 working days of receipt of issue</td>
</tr>
<tr>
<td>Importing countries requirements obtained from sources other than MPI.</td>
<td>Prior to undertaking export treatments</td>
</tr>
<tr>
<td>Number of official treatments carried out.</td>
<td>Monthly</td>
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4.0 TECHNICAL REQUIREMENTS

4.1 Treatment specifications

4.1.1 For export certification

Post harvest treatments of produce for export shall be carried out to the importing country’s specifications (ICPRs) and/or international standards such as ISPM 15.

Treatments being applied to export plants, forest and plant products or Regulated Articles will be those specified by the importing country as a condition of entry.

Only a treatment supplier or wood packaging manufacturer who is listed on the MPI register for certification mark, can apply the certification mark, as per the Technical Requirements: Registered Certification Mark (ISPM 15).
4.1.2 **For import clearance**

Treatments being applied to imported plants, forest and plant products or regulated articles to obtain biosecurity clearance must be a MPI approved treatment as outlined in the appropriate MPI Standard.

If the proposed treatment differs from a MPI supplied specification it must be approved by the Director before the goods are given Biosecurity clearance.

4.1.3 **Access to information**

Treatment suppliers shall document how they access up-to-date specifications for each type of treatment to be applied, and these specifications shall be readily available to treatment technicians and IVA at time of deciding to apply specific treatments.

4.2 **Treatment plan**

For each treatment to be provided the treatment supplier shall develop a treatment plan setting out the critical control points for effective control of the treatment. Each treatment type shall have a monitoring procedure (see clause 4.6).

4.3 **Treatment facilities**

Facilities used for undertaking treatments of consignments shall be capable of delivering the specified treatment to the required specifications. Facilities shall allow the treatment technician to ascertain that the entire treatment achieved the required outcomes eg: nil leaks if fumigating, heat is maintained, spillages and risk goods are contained etc.

Treatment of imported risk goods shall take place at an approved transitional facility or a place where an inspector has authorised those goods to proceed.

Treatments shall take place at the closest appropriate and practicable point of entry or departure where possible or a satisfactory method of segregation and transportation as per 4.5 maintained.

4.4 **Product identification**

i. The treatment supplier is to uniquely identify all products being treated by suitable means throughout the treatment process. This may be by individual item or batch identification;

ii. The processes used for product identification shall be documented;

iii. The product’s status with respect to the stage of treatment (for example untreated, treated but not released, or treated and approved for release) shall be clearly identified by label, location or other suitable means;

iv. Product shall be traceable from at least producer or point of entry into New Zealand to point of export certification or biosecurity clearance;
v. ISPM 15 treated product will be identified as per MPI export certification standard: Technical Requirements: Registered Certification Mark (ISPM 15).

4.5 Product segregation and post treatment security

4.5.1 Product segregation

The treatment supplier shall document their method(s) used to ensure product segregation while the products are under their control to ensure it is:

i. Segregated from other untreated products so as to avoid possible release of unwanted Organisms or cross contamination;

ii. Protected from possible release of unwanted organisms (imports) or contamination during storage and transport (exports);

iii. Protected from possible product substitution.

4.5.2 Post treatment security

The treatment supplier shall document their method used to transfer the responsibility for the security of treated product to the owner of the product.

4.6 Treatment monitoring

Each critical control point identified in the treatment plan (see clause 4.2) shall be monitored.

i. The procedures for ensuring treatment have been applied to the required standard shall be documented and results recorded by the treatment supplier.

ii. Temperature monitoring and recording methods shall be documented, and equipment used must be accurate to +/- 1°C. Examples of suitable equipment are: thermographs for kiln sterilisation, temperature data loggers for fumigants, temperature sensitive bacteria in steam sterilisation. Mercury maximum/minimum thermometers are not acceptable as the prime method for monitoring temperature.

iii. Where product is being treated, the product’s temperature shall be measured in accordance with the treatment specification for the pests of concern.

iv. For application of fumigants a method shall be used to verify the correct concentration of gas, e.g. detector tubes, thermal conductivity analysers, interference refractometers/gas chromatography, fumiscope, control insects (only of a similar genus to the target insects or in e) below), colour indicator sachets of the correct C:T value. A halide lamp is not acceptable for measuring concentration levels.

v. Where an easily discernible organism is the reason for treatment, to ensure effectiveness of a particular treatment beyond all doubt, a reinspection option may be carried out to verify pest mortality in lieu of the above methods.
4.7 Treatment records

Approved treatment suppliers shall keep and maintain a treatment register that shall be:

i. Kept up to date;
ii. Available at all times for the purpose of MPI or IVA audit;
iii. Kept for a period of at least two years from the date of treatment.

The register shall capture the following information:

i. The unique product identification number;
ii. Treatment date;
iii. Treatment location;
iv. Treatment type (e.g. fumigation, active ingredient or physical action);
v. Dose rate, time, product temperature (specification where declaration is required by importing country);
vi. Product description;

vii. Who carried out the treatment (including signature) at commencement and completion (if different person);
viii. Monitoring results;
ix. Treatment certificate number (if issued);
x. Other marks of identification on the product;
xii. Client;
xiii. Reference to other biosecurity documents (e.g. Biosecurity directive);
xiv. Records of any certification numbers assigned to the facility by MPI;
xv. Traceability records, retained to a level that allows the fate of all treated material to be traced from the treatment stage, right through to storage, and despatch to clients.

4.8 Provision of treatment certificates

The treatment supplier shall sign a “certificate of treatment” with the appropriate (as required to obtain clearance or export certification) information from section 4.7 on completion of each treatment. All monitoring results shall be shown on the treatment suppliers register. The certificate shall be signed by a treatment technician.

When a treatment supplier is treating wood packaging material in accordance to ISPM 15 where the wood is to be on-sold or transferred to a wood packaging manufacturer, and the treatment supplier elects not to apply the certification mark directly to the treated wood, a treatment certificate shall be supplied for each batch or lot of treated timber that is on-sold or transferred to the wood packaging manufacturer.

As a minimum this treatment certificate shall contain the following:

i. Name of treatment supplier;
ii. Type of treatment performed e.g. heat treatment;
iii. Date of treatment;
iv. Details of treatment e.g. core temperature, duration of treatment, etc;
v. Certification number of the heat treatment facility;
vi. Description of wood packaging treated e.g. type of packaging, quantity, etc;
vii. Details of any distinguishing marks present on wood packaging; and 
viii. Must be signed by a treatment technician.

5.0 THE AUDIT PROCESS

The treatment supplier’s nominated IVA shall audit their documented and approved system for ongoing compliance.

The audit frequency and scope applied shall be determined by the level of:
i. Confidence in the validation of the treatment;
ii. Confidence attained through prior audits.

5.1 Risk categorisation of treatment supplier:

Level 1  Treatment suppliers with automated and tamper proof monitoring equipment recognised as validating that the treatment specification has been achieved. Refer to Table 2 for qualifying criteria for reduction in audit frequency.

Level 2  All other treatment suppliers. This category of treatment supplier will be audited at the rate in Table 2 based on the number of treatments per year (n) undertaken. Refer to Table 2 for qualifying criteria for reduction in audit frequency.

Table 2. Audit frequencies

<table>
<thead>
<tr>
<th>Risk category</th>
<th>System audit</th>
<th>Entry level surveillance audit frequency</th>
<th>Qualifying criteria for reduction</th>
<th>Medium reduced surveillance audit frequency</th>
<th>Qualifying criteria for reduction</th>
<th>Low reduced surveillance audit frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Level 1</td>
<td>1 per yr (ongoing)</td>
<td>3 per year</td>
<td>One year without a critical non-compliance.</td>
<td>2 per year</td>
<td>Six months without a critical non-compliance.</td>
<td>1 per year</td>
</tr>
<tr>
<td>Level 2</td>
<td>1 per yr (ongoing)</td>
<td>1.5√n</td>
<td>One year without a critical non-compliance.</td>
<td>1.0√n</td>
<td>Six months without a critical non-compliance.</td>
<td>0.5√n</td>
</tr>
</tbody>
</table>

5.2. Changes in audit frequency

5.2.1 Where a critical non-compliance is identified during any audit, the audit frequency shall immediately increase to:

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i. Daily audits for a maximum of three days, during which time the treatment supplier must identify, implement and have verified agreed corrective action(s).
ii. The treatment supplier’s failure to manage the critical non-compliance in accordance with the IVA audit report shall result in the treatment supplier’s system immediately reverting to supervision until an agreed corrective action strategy has been verified as having been implemented by the IVA auditor through a targeted systems audit.

Following satisfactory completion of the above corrective action process the treatment supplier’s system shall resume at the entry-level audit frequency.

5.2.2 Where a major non-compliance is identified the treatment supplier shall move up one frequency

5.2.3 An IVA may determine that the frequency of audit should be increased or reduced. In such cases they shall justify their reasons in writing to MPI.

Such change is at the sole discretion of MPI and on a case-by-case basis.

6.0 NON-COMPLIANCES

Non-compliances will be graded by an IVA or by MPI as critical, major and minor.

6.1 Critical non-compliance

A critical non-compliance will be caused by actions or inactions that lead to the total loss of confidence in the treatment supplier’s compliance with the requirements of its approved quality management system, or that will lead to treatments not complying with specifications.

Critical non-compliances must be addressed as soon as practical, and in any event within three days.

Examples of critical non-compliance include:

i. Incorrect treatment applied;
ii. Failure to follow approved procedures that will impact on the effectiveness of the treatment;
iii. Registered treatment technicians not meeting the competency criteria;
iv. Undertaking a treatment without a treatment suppliers registered, competent treatment technician;
v. Required treatment monitoring not been undertaken;
vi. Required treatment facilities and/or equipment not used;
vii. Equipment calibration not carried out;
viii. Equipment not working to specification;
ix. Live target pests (above the allowable maximum pest limits) found during inspection of the treated product (once the required mortality time has elapsed).
x. Product being certified without being treated;
xii. Untreated product not segregated or separately identified from treated product;
xiii. Three or more major non-compliance faults detected in the current and immediate past audit;
xiv. Agreed actions to address major non-compliances not being completed within agreed timeframes without justification.

6.2 Major non-compliance

A major non-compliance will be caused by actions or inactions that, if not attended to urgently, will lead to the total loss of confidence in the treatment supplier’s compliance with the requirements of its approved or authorised quality management system, or that will lead to treatments not complying with specifications.

Major non-compliances must be addressed as soon as practicable, and in any event within one week.

Examples of major non-compliance include:
i. Auditee fails to identify, classify or record defects correctly;
ii. Amendments to documented procedural details of treatment supplier’s system not notified to the IVA;
iii. Failure to apply document control procedures;
iv. Actions taken following treatments not recorded;
v. Failure to follow approved procedures that is not immediately impacting on the effectiveness of the treatment, but if left unattended will erode confidence in the treatment suppliers system;
vi. Treatment specifications (when specified) not available to treatment technicians.
vii. Corrective action for a minor non-compliance not implemented within the agreed time frame;
viii. Three or more minor non-compliance in any one audit;
ix. Agreed actions to address minor non-compliances not being completed within agreed timeframes;
x. Failure to notify start or recommencing of seasonal operations (where applicable).

6.3 Minor non-compliance

A minor non-compliance will be caused by actions or inactions that are not considered to result in total loss of confidence in the treatment supplier’s quality management system, or that do not lead to treatments not complying with specifications.

Minor non-compliances must be addressed within a period of one month.
6.4 Three majors makes one critical

Should three major non-compliances be raised during the current and preceding audit a critical non-compliance will be raised on:

i. For treatment suppliers under the supervision programme – Clause 6.1 of this document;

ii. For treatment suppliers under the approved treatment supplier programme – clause 6.1 of the treatment supplier standard;

iii. For IVAs - clause 4.1 of the IVA standard.

6.5 Three minors makes one major

Should any three minor non-compliances be raised during any one audit a Major non-compliance will be raised on:

i. For treatment suppliers under the Supervision Programme – Clause 5.4.1 of this document;

ii. For treatment suppliers seeking under the approved treatment supplier programme – clause 4.1.1 of the treatment supplier standard.

6.6 Not addressing non-compliances

Any non-compliance not satisfactorily addressed by the agreed time will be upgraded to the next level of classification, i.e. minor to major, major to critical.

7.0 COSTS

All costs incurred by MPI or an IVA (including time and any travel associated with evaluation of the approved treatment supplier’s system, audit of the approved treatment supplier, communication and reporting) shall be met by the treatment supplier.

8.0 USE OF MPI LOGO AND CLAIMS OF MPI APPROVED STATUS

The MPI logo or the word MPI is not to be used. When approved a treatment supplier can use the following words "Ministry for Primary Industries approved" provided that the wording is specific as to what the approval is for.

For example a fumigation company could display on its certificate:

 Approved by the Ministry for Primary Industries to carry out official fumigations.

Where an approved treatment supplier provides treatments not required by importing countries or directed by MPI to treat imports, no claim of MPI approved status may be made for that treatment and in no circumstances can a certificate that in any way indicates MPI approval of the treatment be issued.
9.0 SUSPENSION OR TERMINATION OF APPROVAL

9.1 Suspension of a treatment suppliers approval

Approval of a treatment supplier to perform official treatments may be suspended by MPI in full or for a specified period, where one or more of the following occurs:

i. Where a treatment supplier’s system has not been active for more than 12 months, or

ii. A treatment supplier has not contracted the services of an IVA; or

iii. A treatment supplier fails to make full payment of fees to MPI unless in dispute; or

iv. Requested by the treatment supplier.

MPI shall formally advise the treatment supplier of their suspension.

During the period of the suspension, the treatment supplier shall not offer or perform any official treatments on behalf of MPI.

Certification of product that was produced since the last successful audit and the date of suspension are at the sole discretion of MPI.

9.2 Reinstatement of a treatment supplier following suspension

Reinstatement of a treatment supplier’s approval by MPI to perform official treatments shall occur only when all conditions prescribed by MPI and or its representatives have been met.

MPI shall formally advise the treatment supplier of the date from which their approval will be reinstated.

9.3 Process for termination of approval

Termination of a treatment supplier’s approval shall occur:

i. Where falsification of any record is found;

ii. Where the treatment supplier is wrongfully claiming MPI approval;

iii. Where more than two critical non-compliances are identified within any 12 month period;

iv. If the conditions for reinstatement in the suspension notice are not met within the specified time;

v. At the request of the treatment supplier.

MPI shall formally advise the treatment supplier of the reasons for the termination of the approval, and the effective date of the termination.

MPI contract of approval shall be returned to MPI (or through the IVA) within five working days of the approval being terminated. In addition any equipment for the
application of certification marks or material advertising MPI approval status shall be disposed of and verified to MPI.
APPENDIX 1

APPLICATION FOR APPROVAL OF TREATMENT SUPPLIER FOR THE
PROVISION OF TREATMENT SERVICES FOR AND ON BEHALF OF
MINISTRY FOR PRIMARY INDUSTRIES (MPI) FOR IMPORT RISK GOODS
AND EXPORT GOODS

Please complete this application and forward to your chosen IVA.

<table>
<thead>
<tr>
<th>Treatment Supplier Name</th>
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<tbody>
<tr>
<td>Address</td>
<td></td>
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<tr>
<td>Locations of operation</td>
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<tr>
<td>Scope of service</td>
<td>Type of Treatments</td>
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<td>Registered certification mark (ISPM15)</td>
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<td>Email</td>
<td></td>
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<tr>
<td>Name and title of person responsible for Treatment Suppliers System</td>
<td></td>
</tr>
</tbody>
</table>
TREATMENT SUPPLIERS STATEMENT

I, ........................................................... (Treatment Supplier), wish to apply for approval under the requirements set down in Ministry for Primary Industries (MPI) Requirements for the supplier of official treatments.

1. I agree to meet the requirements of MPI Requirements for the supplier of official treatments.

2. I agree to document my Treatment Suppliers System meeting the requirements specified by MPI Requirements for the supplier of official treatments.

3. I agree to operate to the above documented system and procedures as approved by Director Plant, Food, and Environment, MPI.

4. I agree to MPI making enquiries and using the information supplied by me, in connection with this application or any contract entered into as a result of this application, for the following purposes:
   i. To ensure that I am a fit and proper person to hold the approved status conferred by the Contract;
   ii. To ensure that I have appropriate consents, permits, licences and authorities in respect of my business operations and my business premises that are required;
   iii. To notify the public of my approved status.

4. I consent to such enquiries being made to or by the Police, Customs Department, New Zealand Horticultural Export Authority, and any statutory Board involved in import and export of products. I consent to publication of my approved status in any publication available to the public.

5. I agree to afford MPI or MPI's representative’s reasonable co-operation and access necessary to carry out audits.

6. Included with this application is a non-refundable application fee of $480.00 (+ GST) for processing the application.

7. I note that any contract of approval will be subject to desk evaluation and subsequent audits and that the above fee excludes these costs. I agree to pay any reasonable costs of such evaluation and audits as may be charged to me from time to time.

8. I understand that if I fail to provide all or any of the information requested in connection with this application, I may be denied approval.

9. I understand that under the Information Privacy Principles of the Privacy Act 1993, I have rights of access to, and correction of, personal information held in connection with this application.
NOTE TO APPLICANT

MPI means any officer or agent of MPI including MPI’s Representative.

This application does not in itself entitle the applicant to provide treatment services for MPI Biosecurity. Approval may be given by MPI once the requirements of MPI Requirements for the supplier of official treatments have been met.

State here the Independent Verification Agency (IVA) you are contracting to undertake pre-approval evaluation and initial audits of your operator system.

........................................................................................................................................................................

(IVA)

........................................................................................................................................................................

(Signature of Treatment Supplier) (Date)

........................................................................................................................................................................

(Name - please print) (Title)

POST THIS APPLICATION TO:

Senior adviser, Biosecurity & Environment

Ministry for Primary Industries New Zealand

PO Box 2526

Wellington
APPENDIX 2

APPROVAL OF TREATMENT SUPPLIER FOR THE PROVISION OF TREATMENT SERVICES FOR AND ON BEHALF OF MINISTRY FOR PRIMARY INDUSTRIES (MPI) FOR IMPORT RISK GOODS AND EXPORT GOODS

CONTRACT OF APPROVAL

Made this day of 20

BETWEEN: “HER MAJESTY THE QUEEN IN RIGHT OF NEW ZEALAND acting by and through the Ministry for Primary Industries (“MPI”)”
AND: (“the Treatment Supplier”)

WHEREAS:

A. Background

MPI is responsible for ensuring that treatments being applied to imported risk goods provide the best practicable level of control, and that treatments being applied to export goods comply with importing country requirements. MPI is also responsible for ensuring that only competent Treatment Suppliers and individuals are involved with the delivery of treatment activities.

The Treatment Supplier has demonstrated procedural ability and proficiency in the provision of Treatment services in relation to import risk goods and export goods.

MPI desires to approve the Treatment Supplier for the purpose of allowing him/her/it to provide Treatment services in relation to import risk goods and export goods for and on behalf of MPI.

The Treatment Supplier desires to be approved by MPI in order to provide Treatment services in relation to import risk goods and export goods for and on behalf of MPI.

The Treatment Supplier acknowledges that the Treatment Supplier has been advised by MPI to obtain legal advice before signing this contract, and to obtain appropriate and sufficient insurance to meet the Treatment Supplier’s potential liabilities (including liabilities to MPI), whether under this Contract or otherwise.
B. Purpose of this Contract

This Contract sets out the legally binding arrangement entered into by MPI and the Treatment Supplier for the approval of the Treatment Supplier by MPI.

DEFINITIONS

“Treatment services means the services that are provided by suppliers under the attached Standard.

“Contract” means this Contract, including the Standard and any other documents and requirements incorporated by reference.

“MPI” means Ministry for Primary Industries.

“Treatment Supplier Quality Management System” refers to the “Treatment Supplier Quality Management System” defined in Appendix 1 of the part of the Standard entitled “Treatment Supplier Programme: Overview and General Requirements”.

“Standard” means the attached documents entitled “Treatment Supplier Programme: Overview and General Requirements” and “Treatment Supplier Programme: Requirements for supplier of official treatments” dated June 1 2006, subject to any changes to the documents (for example, following periodic review).
PRINCIPAL TERMS AND CONDITIONS

1  Term

1.1 This Contract commences on the date it is signed by the authorised representatives of both parties and will, subject to clauses 6, 7 and 9.3, terminate as per section 9 of the part of the Standard entitled “Treatment Supplier Programme: Requirements for supplier of official treatments”.

2 Correctness of Information

2.1 The Treatment Supplier warrants that the following information (including written and oral information) supplied by the Treatment Supplier to MPI is correct and adequate in all respects:

2.1.1 all information supplied in or in connection with the application form entitled "Application for approval of treatment supplier for the provision of treatment services for and on behalf of MPI for import risk goods and export goods";

2.1.2 all other information supplied in connection with the approval of the Treatment Supplier under this Contract; and

2.1.3 all information required to be supplied under the Standard.

3 Treatment Supplier’s Other Warranties

3.1 The Treatment Supplier warrants that throughout the term of this Contract the Treatment Supplier will maintain its Treatment Supplier Quality Management System and all other relevant practices to substantially correspond with all the information referred to in clause 2.1, except to the extent that any changes made are approved by MPI in accordance with the Standard.

3.2 The Treatment Supplier warrants to notify MPI of any change to the Treatment Supplier’s name.

3.3 The Treatment Supplier warrants that where it is an unlisted company, it will notify MPI as soon as reasonably practicable of any:

3.3.1 change in the legal or beneficial ownership of any of its shares; or
3.3.2 issue of new capital; or
3.3.3 change to the rights and powers attaching to any of its shares; or
3.3.4 change to the composition of the board of directors (as this term is defined in section 127 of the Companies Act 1993).
3.4 The Treatment Supplier warrants to fully comply with all the requirements, and other specifications set out in the Standard.

3.5 The Treatment Supplier warrants to take all reasonable steps to enable and facilitate MPI, and any persons acting for or otherwise associated with MPI, to perform their tasks and functions as envisaged in, or otherwise in connection with, the Standard.

3.6 The Treatment Supplier warrants not providing activities for purposes not covered by this Contract. The Treatment Supplier will take all reasonable steps to ensure that these activities are not provided for such unauthorised purposes, or by unauthorised persons.

4 MPI’s Obligation

4.1 MPI hereby approves the Treatment Supplier for the term of this Contract for the purpose of enabling the Treatment Supplier to provide Treatment services for and on behalf of MPI in relation to import risk goods and export goods.

4.2 The Treatment Supplier accepts that nothing in this Contract or in any dealings of any kind between the Treatment Supplier and MPI, Crown officers, or agents of or other persons associated with MPI or Crown officers, represents to the Treatment Supplier or otherwise creates any kind of expectation on the Treatment Supplier’s part that:

4.2.1 any other approval or any certification of any kind will be granted by MPI or will be granted within a certain time period; or

4.2.2 any plant products, or other things that are accompanied by, or otherwise reliant on any Treatment services provided by the Treatment Supplier on behalf of MPI will be accepted by an importing country’s official control authorities or will be accepted within a certain time period.

4.3 MPI Quarantine Service will, and MPI will ensure that independent verification agencies, only accept valid treatment certificates

5 EXCLUSION OF LIABILITY

5.1 THE TREATMENT SUPPLIER ACCEPTS THAT UNDER NO CIRCUMSTANCES WILL MPI, CROWN OFFICERS, OR AGENTS OF OR OTHER PERSONS ASSOCIATED WITH MPI OR CROWN OFFICERS, BE LIABLE UNDER THE LAW OF TORT, CONTRACT, OR OTHERWISE FOR ANY LOSS, CLAIM, ACTION, DEMAND, EXPENSE, INQUIRY, HARM, OR DAMAGE, HOWEVER CAUSED, ARISING DIRECTLY OR INDIRECTLY FROM OR CONNECTED IN ANY WAY TO:

5.1.1 THE PERFORMANCE, OR AS THE CASE MAY BE, NON-PERFORMANCE OF THE TREATMENT SUPPLIER (OR ANY OF ITS CONTRACTORS, SUB-CONTRACTORS, AGENTS, OR EMPLOYEES THAT ARE NOT A PARTY
5.1.2 THE PROVISION OR NON-PROVISION OF ANY TREATMENT SERVICES BY THE TREATMENT SUPPLIER.

6 **Suspension and Termination by MPI**

6.1 MPI may at any time suspend approval of the Treatment Supplier in accordance with Section 9 of the part of the Standard entitled “Treatment Supplier Programme: Requirements for supplier of official treatments”, in addition to any other rights of suspension provided by law.

6.2 MPI may at any time terminate approval of the Treatment Supplier in accordance with Section 9 of the part of the Standard entitled “Treatment Supplier Programme: Requirements for supplier of official treatments”, in addition to any other rights of termination provided by law.

6.3 MPI may at any time suspend or terminate approval of the Treatment Supplier for breach of the Standard relating to payment of fees in accordance with Section 9.1 iii of the part of the Standard entitled “Treatment Supplier Programme: Requirements for supplier of official treatments”.

6.4 Where a change of a kind that is specified in clause 3.3 occurs, MPI may terminate the approval of the Treatment Supplier.

7 **Extension following Audit of Treatment Supplier**

7.1 Where the results of audits of compliance with the Standard indicate the requirements of the Standard are being complied with, the Contract will be deemed extended, subject to clauses 6 and 9.3, beyond the last audit date.

8 **INDEMNITY AND INSURANCE**

8.1 The Treatment Supplier will **INDEMNIFY AND KEEP INDEMNIFIED** MPI from and against any liability, loss, damage, costs and expenses (including legal costs and any expenses of going to arbitration), which MPI may suffer or incur arising directly or indirectly from:

8.1.1 the performance, or as the case may be, non-performance of the Treatment Supplier (or any of its contractors, sub-contractors, agents, or employees that are not a party to this Contract) of any of its obligations in respect of this Contract;

8.1.2 negligent acts or omissions on the part of the Treatment Supplier (or any of its contractors, sub-contractors, agents, or employees that are not a party to this Contract);
8.1.3 suspension or termination of the Treatment Supplier 's approval in accordance with clause 6; or

8.1.4 the provision or non-provision of Treatment services for MPI by the Treatment Supplier.

8.2 The Treatment Supplier will maintain public liability insurance with a minimum sum insured of $1,000,000 (1 million).

9  **Force Majeure**

9.1 Notwithstanding any other provision of this Contract, neither party shall be liable to the other for any act or omission, or any failure to comply with any warranty or to perform any of its obligations under this Contract, where such, act, omission, or failure is caused by fire, flood, storm, earthquake, civil disturbance, war, act of God, or any other event or circumstances reasonably beyond its control (called “Force Majeure”), **provided that** the party alleging Force Majeure has taken all reasonable precautions to avoid or mitigate the consequences of such occurrence.

9.2 The party unable to fulfil its obligations due to Force Majeure will immediately:

9.2.1 notify the other in writing of the reasons for its failure to comply with the warranty or to perform the obligation, and the effect of such failure; and

9.2.2 use all responsible endeavours to avoid or remove the cause and comply with the warranty or perform the obligation.

9.3 Upon receiving notice pursuant to clause 9.2, or upon otherwise being made aware of any Force Majeure circumstances affecting the Treatment Supplier, MPI may at its absolute discretion suspend approval of the Treatment Supplier until such time as the circumstances have been avoided, removed or abated sufficiently to enable the Treatment Supplier to comply with the warranty or perform the obligation.

10 **Assignment**

10.1 Neither party shall assign all or any of its rights, obligations, or liabilities under this Contract. In the event of a purported assignment in breach of this clause, this Contract shall terminate.
11 **Disputes**

11.1 The parties agree to use their best efforts to resolve any dispute which may arise under the Contract through good faith negotiations. Except as provided in clause 11.4, no party shall commence any arbitration or litigation in relation to this Contract unless it has first invited the chief executive of the other party to meet with its own chief executive for the purpose of endeavouring to resolve the dispute on mutually acceptable terms.

11.2 Should resolution of the dispute not be achieved at chief executive level, the dispute will be submitted to mediation before any litigation is commenced. Any party may initiate mediation by giving written notice to the other party of their intent to do so. Should the parties be unable to agree on a mediator within two (2) working days of receipt of notice of intent to seek mediation, then the mediator will be selected by the President for the time being of the Lawyers Engaged in Alternative Dispute Resolution (LEADR) or its successor.

11.3 Any dispute arising under this Contract which cannot be settled by negotiation or mediation between the parties or their respective representatives shall be submitted to arbitration in accordance with the Arbitration Act 1996.

11.4 In the absence of agreement concerning the appointment of an arbitrator, either party may request the President of the New Zealand Law Society to appoint a suitably qualified independent arbitrator to hear and determine the dispute.

11.5 Nothing in this clause shall preclude either party from taking immediate steps to seek urgent equitable relief before a New Zealand Court.

12 **Application of Biosecurity Act**

12.1 Nothing in this Contract overrides any obligations of MPI and the Treatment Supplier under the Biosecurity Act 1993.
13 **Entire Agreement**

13.1 This Contract sets out the entire agreement between the parties.

Signed for and on behalf of:

**HER MAJESTY THE QUEEN IN RIGHT OF NEW ZEALAND**
(acting by and through the Ministry for Primary Industries)

Name: ____________________  
Position: ____________________  
Date: ____________________

WITNESS:  
Name: ____________________  
Occupation: ____________________  
Address: ____________________

Signed for and on behalf of the treatment supplier  
Name: ____________________  
Position: ____________________  
Date: ____________________

WITNESS:  
Name: ____________________  
Occupation: ____________________  
Address: ____________________
APPENDIX 3

Treatment Technician and other staff Competency

1.0 TREATMENT TECHNICIAN COMPETENCY

The treatment technician shall be physically able to carry out the treatment process being undertaken and provide evidence that they comply with any requirements for testing (e.g. demonstrating full colour vision where colour recognition is required).

The treatment technician shall demonstrate the following competencies (measured during audits) relevant to the treatment being applied:

i. Sound understanding of the treatment processes;
ii. Ability to determine which is the correct treatment to be provided and to obtain necessary up-to-date specifications for that treatment;
iii. Knowledge of the treatment being applied. (e.g. Calculations on dose rate/time/temperature relationships);
iv. Facility and equipment operation and maintenance;
v. Equipment calibration procedures;
vi. Security, segregation and identification of treated product (goods);
vii. Treatment efficacy methods;
viii. Reaction properties on products (goods) for that type of treatment;
ix. Creation and filing of all required records;
x. Where required by the HSNO act, hold an approved handler certificate, and a controlled substance license;
xii. Where relevant, pest management training, must have been completed to the equivalent of TOPNZ Unit Standard 3263;
xiii. Understanding of relevant Legislation and Regulations;
xiv. Understanding of Material Safety Data Sheets (MSDS) of individual chemicals used;
xv. Emergency response procedures (4.9)

2.0 OTHER STAFF

Other staff may be used for treatment work provided that:

i. Their duties are commensurate with their knowledge and experience; and
ii. They are given adequate direction; and
iii. Their work is under direct supervision of a treatment technician.
iv. The treatment technician supervising their work is not involved in the application of the same treatment (therefore treatment technician is supervising only)

Other staff shall be provided with appropriate detailed procedures and checklists.