In respect of the following General Import Authorisation:

**IMP/GEN/2010/07** - dated 24 February 2010 for Heat treated organs, glands, blood or other tissues, or products derived from these tissues from ungulates, born and bred in captivity and not known or suspected to be infected with specified animal pathogens, intended for educational, research and museum purposes from Australia, Canada, Iceland, Norway, New Zealand, Switzerland and the USA.

The above-mentioned Authorisation has been amended as follows:

**Products are amended to read:**

**IMP/GEN/2010/07** - dated 24 February 2010 for Heat treated organs, glands, blood or other tissues, or products derived from these tissues from ungulates, born and bred in captivity and not known or suspected to be infected with specified animal pathogens, intended for research and diagnostic purposes.

**Condition 2, second bullet point is amended to read:**

commercial/shipping documents providing the name and address of consignor and consignee, description of the product and quantities, the place of origin, place of dispatch and the category of the material as defined in Regulation (EC) No 1069/2009;

**Condition 3, second bullet point is amended to read:**

the products are not derived from animals known or suspected to be infected with a pathogen which causes a notifiable disease to which animals are susceptible in accordance with the Importation of Animal Pathogens Order 1980 (IAPO) or the Specified Animal Pathogens Order 2008 (SAPO) or the animal health legislation of the exporting country;

**Condition 11 is amended to read:**

On completion of the testing, and unless they are kept for reference purposes or re-dispatched to the third country of origin (see General note 7), any residues of the material and any products derived from the use of those samples shall be disposed of in accordance with the requirements of Annex XIV, Chapter III, Section 1 of Regulation (EU) No 142/2011.
General note 1 is amended to read:

Nothing in this authorisation gives exemption from any prohibition or restriction imposed by any other legislation including: the Official Feed and Food Controls (England) Regulations 2009, the provisions of the Food Safety Act 1990 and Regulations made under it, the Animal By Products (Identification) Regulations 1995, the Animal By-Products Regulations 2005, the Marketing Authorisations for Veterinary Medicinal Products Regulations 1994 the Control of Substances Hazardous to Health, International Air Transport Association, the Convention on International Trade in Endangered Species or by any regulation superseding or amending the same.

General notes are amended to include:

9. Definitions

Research and diagnostic samples – as defined in Regulation (EU) No 142/2011 are animal by products and derived products intended for the following purposes: examination in the context of diagnostic activities or analysis for the promotion of progress in science and technology, in the context of educational or research activities

10. Should you wish to import Display items or Trade Samples as defined in Regulation (EU) No 142/2011, please contact AH Chelmsford.

11. Any subsequent use of research and diagnostic samples for purposes other than those referred to in point 38 of Annex 1 shall be prohibited.

12. Please refer to the Importer Information Notes (IINs) on the Defra website for further information.

Dated: 18 March 2011

Officer of the Department for Environment, Food and Rural Affairs