Standard License Restrictions

Product licenses may be issued with one or more restrictions. Some types of licenses always require restrictions. Restrictions may be considered on a case-by-case basis for others. Table 1 includes a list of such products. Review this list when licensing new products OR reissuing existing licenses, as the need for certain restrictions may change over time. Table 2 provides the text that corresponds to a particular restriction number.

Table 1.

| Product/Fraction | Required Restrictions (by | Additional Restrictions to Consider (by LSRTIS |
|--|--------------------------------|---|
| A 1 36 ' 1 37 ' | LSRTIS Number) | Number) |
| Anaplasma Marginale Vaccine | 42 | |
| Allergenic Extract | 42 | |
| Antivenom | 42 | |
| Autogenous | 55 | |
| Avian Pneumovirus, live/modified live | 42, 43, 44 | |
| Avian Reovirus Vaccine, Killed | 43 | |
| Avian Influenza Vaccine (turkeys, non-H5, non-H7 | 43, 46 | 60 |
| Avian Influenza vaccine (turkey H5, turkey H7, all chicken | 43, 46, 58 | 60 |
| Babesia caballi | 45 | |
| Babesia equi | 45 | |
| Bovine spongiform encephalopathy | 45, 62, 343 | 46 |
| diagnostics | | |
| Bronchitis Vaccine, Live Virus (except | 43 | |
| Mass & Conn) | | |
| Brucella abortus (vaccines & kits) | 42, 43 | |
| Bursal Disease Vaccine, Live Virus | 43, 47 | |
| Bursal Disease Vaccine, Killed | 43 | |
| Cancer products | 42 | |
| Conditional licenses ("regular") | 43, 46, 48, 49, 50, 56, 480 | 42, 54, 57 |
| Chronic wasting disease diagnostics | 45, 62 | |
| Duck enteritis virus | 43 | |
| Ehrlichia risticii | 43 | |
| Equine arteritis virus, live | 43 | |
| Equine infectious anemia diagnostics | 45 | |
| Feline immunodeficiency virus (vaccine) | 42 | 54 |
| Mycobacterium bovis diagnostics | 42, 43 | |

| Product/Fraction | Required Restrictions (by LSRTIS Number) | Additional Restrictions to Consider (by LSRTIS Number) |
|--|---|--|
| Mycobacterium paratuberculosis | 43 | |
| (vaccines and kits) | | |
| Mycobacterium tuberculosis diagnostics | 42, 43 | |
| Mycoplasma gallisepticum (vaccine & kits) | 43 | |
| Mycoplasma synoviae (vaccines & kits) | 43 | |
| Permitted products | 64, 186 | 186 may be exempted from products for USDA emergency use if the Outline of Production has adequate assurances regarding Ingredients of Animal Origin (IAO) source/testing and a statement that IAO sources/testing will not change without approval of CVB |
| Pigeon Pox Vaccine, live | 60 | |
| Platform Products (conditional) | 43, 46, 48, 50, 56, 71 | Contact PEL Director before licensing a platform product, as some restrictions are still being discussed. |
| Prescription Products | 42, 43, 46, 48, 50, 54, 55, 56, 57, 132, 200, 540 | |
| PRRS virus, live and modified live | 47 | |
| Pseudorabies virus (vaccine & kits) | 43, 45 | |
| Rabies virus | 43 | |
| Scrapie diagnostics | 45, 46, 62 | |
| Streptococcus equi (whole cell bacterin) | 42 | |
| Tenosynovitis virus, live and modified live | 43 | |
| Tuberculin | 42, 43 | |
| Vibrio salmonicida | 43 | |
| Products for disease exotic to U.S. | 52 | 46 |
| Products with wildlife claims | | 46, 58, 61, 62 |
| Components of combination packages in which firm with comb.pkg license does not have a significant step in production of the component | 53 | |
| For further manufacture products | 51 | |

| Product/Fraction | Required Restrictions (by LSRTIS Number) | Additional Restrictions to Consider (by LSRTIS Number) |
|--|--|--|
| Products for diseases appearing on OIE | | 45, 46 |
| list | | |
| Biotechnology-derived products | | 46 |
| containing live vectors that can replicate | | |
| in the vaccinated animal | | |
| Products with Additional Reporting | | 57 |
| Requirement | | |
| Products that are tested at a contract | 70 | |
| facility for any Section V test | | |
| Products with target animal safety testing | 57 (yearly) | |
| exemption | | |

Table 2

| LSRTIS License Restriction Number | Restriction |
|--------------------------------------|---|
| 42 | For use by, or under the supervision of, a veterinarian. |
| 43 | Distribution in each State shall be limited to authorized recipients |
| .0 | designated by proper State officialsunder such additional conditions |
| | as these authorities may require. |
| 44 | The distribution and use of this vaccine shall be limited to <specify< td=""></specify<> |
| | state(s) and/or foreign countries>. |
| 45 | Sale and use in the United States restricted to laboratories approved |
| | by State and Federal (USDA) animal health officials. |
| 46 | Export distribution shall be limited to authorized recipients |
| | designated by proper animal health regulatory officialsunder such |
| | additional conditions as these authorities may require. |
| 47 | Recommended use shall be restricted to premises having a history of |
| | the disease. |
| 48 | Reissuance of this license shall be considered in accordance with 9 |
| | CFR Part 102.6. |
| 49 | The following statement shall appear on all labels: This product |
| | license is conditional. Efficacy and potency test studies in progress. |
| 50 | Trade Names shall not be used with this product. |
| 51 | For further manufacture. |
| 52 | For Export Only. |
| 53 | This product may only be marketed as a component of <other< td=""></other<> |
| | Product Codes>. |
| 54 | Marketing and promotional materials must be reviewed by the CVB |
| | prior to publication or distribution. |
| 55 | This license does not authorize production, distribution, or shipment |
| | of a vaccine/bacterin for foot-and-mouth disease, rinderpest, any H5 |
| | or H7 subtype of avian influenza, any subtype of avian influenza in |
| | chickens, swine vesicular disease, Newcastle disease, African swine |
| | fever, classical swine fever, Brucella abortus, vesicular stomatitis, |
| | and rabbit hemorrhagic disease or any other disease the Administrator |
| | determines may pose a risk to animal or public health. |
| 56 | This license will terminate on <specify date="" length="" of="" or="" time="">.</specify> |
| 57 | Unusual conditions or adverse events linked to vaccinated animals |
| | are to be reported to the CVB <specify interval="" reporting="">.</specify> |
| 58 | Domestic distribution and use shall be under the supervision or |
| | control of USDA, APHIS, Veterinary Services, as part of an official |
| | USDA animal disease control program. |
| 59 | Recommended use shall be restricted to healthy 10- to 17-week-old |
| | broiler breeder replacement chickens and to premises where no other |
| | susceptible chickens are maintained. |
| 60 | For use in <specified species=""> only.</specified> |
| 61 | Restricted to use in State or Federal Rabies Control Programs. |

| 62 | Potency testing and distribution and use shall be under the |
|-----|---|
| | supervision or control of USDA, APHIS, Veterinary Servicesunder |
| | such additional conditions as these authorities may require. |
| 63 | Serials may be released, subject to immediate recall, should the |
| | Master Seed/Cell Stocks, now under test by APHIS, be found |
| | unsatisfactory. |
| 64 | The producer/permittee agrees to submit to periodic reinspections of |
| | the production facility under terms to be specified in a Cooperative |
| | Agreement between the parties. The permittee agrees to be |
| | responsible for all costs associated with these inspections. |
| 70 | The CVB has the authority to inspect TGA Sciences, Inc., No. T100, |
| | that performs the Serology, Clostridium botulinum Type B potency |
| | test contracted by the Licensee. |
| 71 | The license may be renewed upon request by the firm and at the |
| | discretion of APHIS based on product performance, safety profile, |
| | manufacturing consistency, and inspections by the CVB. |
| 132 | Disposition records, maintained according to 9 CFR 116.2, shall be |
| | prepared in a format acceptable to APHIS and submitted to CVB at |
| | intervals determined by APHIS. |
| 186 | Each shipment of biological product distributed and sold under this |
| | permit must be accompanied by an original certificate endorsed by a |
| | full-time, salaried veterinarian of the agency responsible for animal |
| | health of the Government of <specify country="">, or other assurances</specify> |
| | acceptable to APHIS, certifying that: 1.) Ingredients of animal origin |
| | are sourced from the United States or countries considered free, low |
| | risk, or not affected with foreign animal diseases of concern and with |
| | negligible or controlled risk of Bovine Spongiform Encephalopathy |
| | [BSE], according to APHIS' Animal Disease Status designations as |
| | defined in Veterinary Services Memorandum 800.51. 2.) During the |
| | manufacturing process, the manufacturing facility does not receive, |
| | store, or process any ingredients of ruminant origin from countries |
| | with BSE. 3.) The product complies with all other provisions of 9 |
| | CFR 113.53. |
| 200 | The license may be renewed upon request by the firm and at the |
| | discretion of APHIS based on manufacturing consistency and |
| | inspections by the CVB. For use as a veterinary prescription. |
| | Efficacy and potency have not been demonstrated. |
| 343 | Use of this kit is restricted to testing conducted as part of the official |
| | USDA BSE surveillance program. |
| 344 | Product imported under this permit may only be received by USDA |
| | personnel or persons designated by the USDA, as part of an official |
| | USDA animal disease control program. |
| 480 | The licensee shall demonstrate acceptable progress toward |
| | completion of host animal efficacy and potency test studies in |
| | accordance with acceptable protocols filed with the CVB prior to |
| | reissuance. |
| | • |

| 540 | Marketing and promotional materials are restricted to licensed |
|-----|--|
| | veterinarians. |
| 551 | This license restricted to antigens from the VP7 gene from an |
| | individual Rotavirus Strain C isolate. |