(4) Any establishment wholesaling any animals (except birds, rats, and mice); and
(5) Any establishment exhibiting pet animals in a room that is separate from or adjacent to the retail pet store, or in an outside area, or anywhere off the retail pet store premises.

Sanctuary area means that area in a primary enclosure for a swim-with-the-dolphin program that is off-limits to the public and that directly abuts the buffer area.

Sanitize means to make physically clean and to remove and destroy, to the maximum degree that is practical, agents injurious to health.

Secretary means the Secretary of Agriculture of the United States or his representative who shall be an employee of the Department.

Sheltered housing facility means a housing facility which provides the animals with shelter; protection from the elements; and protection from temperature extremes at all times. A sheltered housing facility may consist of runs or pens totally enclosed in a barn or building, or of connecting inside/outside runs or pens with the inside pens in a totally enclosed building.

Standards means the requirements with respect to the humane housing, exhibition, handling, care, treatment, temperature, and transportation of animals by dealers, exhibitors research facilities, carriers, intermediate handlers, and operators of auction sales as set forth in part 3 of this subchapter.

State means a State of the United States, the District of Columbia, Commonwealth of Puerto Rico, the Virgin Islands, Guam, American Samoa, or any other territory or possession of the United States.

Study area means any building room, area, enclosure, or other containment outside of a core facility or centrally designated or managed area in which animals are housed for more than 12 hours.

Swim-with-the-dolphin (SWTD) program means any human-cetacean interactive program in which a member of the public enters the primary enclosure in which an SWTD designated cetacean is housed to interact with the animal. This interaction includes, but such inclusions are not limited to, wading, swimming, snorkeling, or scuba diving in the enclosure. This interaction excludes, but such exclusions are not limited to, feeding and petting pools, and the participation of any member(s) of the public audience as a minor segment of an educational presentation or performance of a show.

Transporting device means an interim vehicle or device, other than man, used to transport an animal between the primary conveyance and the terminal facility or in and around the terminal facility of a carrier or intermediate handler.

Transporting vehicle means any truck, car, trailer, airplane, ship, or railroad car used for transporting animals.

Weaned means that an animal has become accustomed to take solid food and has so done, without nursing, for a period of at least 5 days.

Wild animal means any animal which is now or historically has been found in the wild, or in the wild state, within the boundaries of the United States, its territories, or possessions. This term includes, but is not limited to, animals such as: Deer, skunk, opossum, raccoon, mink, armadillo, coyote, squirrel, fox, wolf.

Wild state means living in its original, natural condition; not domesticated.

Zoo means any park, building, cage, enclosure, or other structure or premise in which a live animal or animals are kept for public exhibition or viewing, regardless of compensation.


**EFFECTIVE DATE NOTE:** At 64 FR 15920, Apr. 2, 1999, the definitions of buffer area, interactive area, interactive session, sanctuary area, and swim-with-the-dolphin (SWTD) program were suspended, effective Apr. 2, 1999.

**PART 2—REGULATIONS**

Subpart A—Licensing

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(iii) Valid addresses for all locations, facilities, premises, or sites where animals, animal facilities, equipment, and records are held, kept, or maintained;

(iv) The anticipated maximum number of animals on hand at any one time during the period of licensure;

(v) The anticipated type of animals described in paragraph (b)(2)(ii) of this section to be owned, held, maintained, sold, or exhibited, including those animals leased, during the period of licensure;

(vi) If the person is seeking a license as an exhibitor, whether the person intends to exhibit any animal at any location other than the person’s location(s) listed pursuant to paragraph (a)(1)(iii) of this section; and

(vii) Disclosure of any plea of nolo contendere (no contest) or finding of violation of Federal, State, or local laws or regulations pertaining to animal cruelty or the transportation, ownership, neglect, or welfare of animals.

(2) The completed application form, along with a $120 license fee, shall be submitted to the appropriate Animal Care office.

(3) The following persons are exempt from the licensing requirements under section 2 or section 3 of the Act:

(i) Retail pet stores as defined in part 1 of this subchapter;

(ii) Any person who sells or negotiates the sale or purchase of any animal except wild or exotic animals, dogs, or cats, and who derives no more than $500 gross income from the sale of such animals during any calendar year and is not otherwise required to obtain a license;

(iii) Any person who maintains a total of four or fewer breeding female pet animals as defined in part 1 of this subchapter, small exotic or wild mammals (such as hedgehogs, degus, spiny mice, prairie dogs, flying squirrels, jerboas, domesticated ferrets, chinchillas, and gerbils), and/or domesticated farm-type animals (such as cows, goats, pigs, sheep, llamas, and alpacas) and sells only the offspring of these animals, which were born and raised on his or her premises, for pets or exhibition, and is not otherwise required to obtain a license. This exemption does not extend to any person residing in a household that collectively maintains a total of more than four of these breeding female animals, regardless of ownership, or to any person maintaining such breeding female animals on premises on which more than four of these breeding female animals are maintained, or to any person acting in concert with others where they collectively maintain a total of more than four of these breeding female animals, regardless of ownership;

(iv) Any person who sells fewer than 25 dogs and/or cats per year, which were born and raised on his or her premises, for research, teaching, or testing purposes or to any research facility and is not otherwise required to obtain a license. This exemption does not extend to any person residing in a household that collectively sells 25 or more dogs and/or cats, regardless of ownership, nor to any person acting in concert with others where they collectively sell 25 or more dogs and/or cats, regardless of ownership. The sale of any dog or cat not born and raised on the premises for research purposes requires a license;

(v) Any person who arranges for transportation or transports animals solely for the purpose of breeding, exhibiting in purebred shows, boarding (not in association with commercial transportation), grooming, or medical treatment, and is not otherwise required to obtain a license;

(vi) Any person who buys, sells, transports, or negotiates the sale, purchase, or transportation of any animals used only for the purposes of food or fiber (including fur);

(vii) Any person who maintains a total of eight or fewer pet animals as defined in part 1 of this subchapter, small exotic or wild mammals (such as hedgehogs, degus, spiny mice, prairie dogs, flying squirrels, jerboas, domesticated ferrets, chinchillas, and gerbils), and/or domesticated farm-type animals (such as cows, goats, pigs, sheep, llamas, and alpacas) for exhibition, and is not otherwise required to obtain a license. This exemption does not extend to any person acting in concert with others where they collectively maintain a total of more than eight of these animals for exhibition,
§ 2.2 Acknowledgement of regulations and standards.

Animal Care will supply a copy of the Act and the regulations and standards in this subchapter to an applicant upon request. Signing the application form is an acknowledgement that the applicant has reviewed the Act and the regulations and standards and agrees to comply with them.

(Approved by the Office of Management and Budget under control numbers 0579-0036 and 0579-0470)

§ 2.2 Animal and Plant Health Inspection Service, USDA

regardless of possession and/or ownership;

(viii) Any person who buys animals solely for his or her own use or enjoyment and does not sell or exhibit animals, or is not otherwise required to obtain a license;

(b)(1) No person shall have more than one license. Licenses are issued to specific persons, and are issued for specific activities, types and numbers of animals, and approved sites. A new license must be obtained upon change of ownership, location, activities, or animals. A licensee shall notify Animal Care no fewer than 90 days and obtain a new license before any change in the name, address, substantial control or ownership of his business or operation, locations, activities, and number or type of animals described in paragraph (b)(2) of this section. Any person who is subject to the regulations in this subchapter and who intends to exhibit any animal at any location other than the person’s approved site must provide that information on their application form in accordance with paragraph (a) of this section and submit written itineraries in accordance with §2.126.

(2) Licenses authorize a specific number and specific type(s) of animals, as follows:

(i) Licenses authorize increments of 50 animals on hand at any single point in time during the period of licensure. A licensee must obtain a new license before any change resulting in more than the authorized number of animals on hand at any single point in time during the period of licensure.

(ii) Licenses authorize the use of animals subject to subparts A through F in part 3 of this subchapter, except that, for animals subject to subparts D and F, licenses must specifically authorize the use of each of the following groups of animals: Group 5 (baboons and nonbrachiating species larger than 33 pounds) and Group 6 (great apes over 55 pounds and brachiating species) nonhuman primates; exotic and wild felids (including but not limited to lions, tigers, leopards, cheetahs, jaguars, cougars, lynx, servals, bobcats, and caracals, and any hybrid cross thereof); hyenas and/or exotic and wild canids (including but not limited to wolves, coyotes, foxes, and jackals); bears; and mega-herbivores (including but not limited to elephants, rhinoceroses, hippopotamuses, and giraffes). A licensee must obtain a new license before using any animal beyond those types or numbers of animals authorized under the existing license.

(c) A license will be issued to any applicant, except as provided in §§2.3 through 2.11, when:

(1) The applicant has met the requirements of this section and §§2.2 and 2.3; and

(2) The applicant has paid a $120 license fee to the appropriate Animal Care office. The applicant may pay the fee by certified check, cashier’s check, personal check, money order, or credit card. An applicant whose check is returned by a bank will be charged a fee of $20 for each returned check. If an applicant’s check is returned, subsequent fees must be paid by certified check, cashier’s check, money order, or credit card.

(d) The failure of any person to comply with any provision of the Act, or any of the provisions of the regulations or standards in this subchapter, shall constitute grounds for denial of a license or for its suspension or revocation by the Secretary, as provided in the Act.

(Approved by the Office of Management and Budget under control numbers 0579-0036 and 0579-0470)

§ 2.2 Animal and Plant Health Inspection Service, USDA

Animal Care will supply a copy of the Act and the regulations and standards in this subchapter to an applicant upon request. Signing the application form is an acknowledgement that the applicant has reviewed the Act and the regulations and standards and agrees to comply with them.

(Approved by the Office of Management and Budget under control numbers 0579-0036 and 0579-0470)

§ 2.2 Animal and Plant Health Inspection Service, USDA

Animal Care will supply a copy of the Act and the regulations and standards in this subchapter to an applicant upon request. Signing the application form is an acknowledgement that the applicant has reviewed the Act and the regulations and standards and agrees to comply with them.

(Approved by the Office of Management and Budget under control numbers 0579-0036 and 0579-0470)
§ 2.3 Demonstration of compliance with standards and regulations.

(a) Each applicant for a license must demonstrate that his or her location(s) and any animals, facilities, vehicles, equipment, or other locations used or intended for use in the business comply with the Act and the regulations and standards set forth in parts 2 and 3 of this subchapter. Each applicant must make his or her animals, locations, facilities, vehicles, equipment, and records available for inspection during business hours and at other times mutually agreeable to the applicant and APHIS, to ascertain the applicant’s compliance with the Act and the regulations and standards.

(b) Each applicant for a license must be inspected by APHIS and demonstrate compliance with the Act and the regulations and standards, as required in paragraph (a) of this section, before APHIS will issue a license. If the first inspection reveals that the applicant’s animals, premises, facilities, vehicles, equipment, locations, or records do not meet the applicable requirements of this subchapter, APHIS will advise the applicant of existing deficiencies and the corrective measures that must be completed to come into compliance with the regulations and standards. An applicant who fails the first inspection may request up to two more inspections by APHIS to demonstrate compliance with the Act and the regulations and standards. The applicant must request the second inspection, and if applicable, the third inspection, within 60 days following the first inspection.

(c) Any applicant who fails the third and final prelicense inspection may appeal all or part of the inspection findings to the Deputy Administrator. To appeal, the applicant must send a written statement contesting the inspection finding(s) and include any documentation or other information in support of the appeal. To receive consideration, the appeal must be received by the Deputy Administrator within 7 days of the date the applicant received the third prelicense inspection report. Within 7 days of receiving a timely appeal, the Deputy Administrator will issue a written response to notify the applicant whether APHIS will issue a license or deny the application.

(d) If an applicant fails inspection or fails to request reinspections within the 60-day period, or fails to submit a timely appeal of the third prelicense inspection report as described in paragraph (c) of this section, the applicant cannot reapply for a license for a period of 6 months from the date of the failed third inspection or the expiration of the time to request a third inspection. No license will be issued until the applicant pays the license fee and demonstrates upon inspection that the animals, premises, facilities, vehicles, equipment, locations, and records are in compliance with all applicable requirements in the Act and the regulations and standards in this subchapter.

(Approved by the Office of Management and Budget under control number 0579–0036)

§ 2.4 Non-interference with APHIS officials.

A licensee or applicant for an initial license shall not interfere with, threaten, abuse (including verbally abuse), or harass any APHIS official in the course of carrying out his or her duties.

§ 2.5 Duration of license and termination of license.

(a) A license issued under this part shall be valid and effective for 3 years unless:

1. The license has been revoked or suspended pursuant to section 19 of the Act or terminated pursuant to § 2.12.

2. The license is voluntarily terminated upon request of the licensee, in writing, to the Deputy Administrator.

3. The license has expired, except that:

   (1) The Deputy Administrator may issue a temporary license, which automatically expires after 120 days, to an applicant whose immediately preceding 3-year license has expired, if: 

   A. The applicant submits the appropriate application form before the expiration date of a preceding license; and

   B. The applicant had no noncompliances with the Act and the regulations and standards in parts 2 and 3 of this subchapter documented in any inspection report during the preceding period of licensure.
§ 2.11 Denial of license application.

(a) A license will not be issued to any applicant who:

(1) Has not complied with the requirements of §§ 2.1 through 2.4 and has not paid the fees indicated in § 2.1;

(2) Is not in compliance with the Act or any of the regulations or standards in this subchapter;

(3) Has had a license revoked or whose license is suspended, as set forth in § 2.1(d);

(4) Was an officer, agent, or employee of a licensee whose license has been suspended or revoked and who was responsible for or participated in the activity upon which the order of suspension or revocation was based, as set forth in § 2.9;

(5) Has pled nolo contendere (no contest) or has been found to have violated any Federal, State, or local laws or regulations pertaining to animal cruelty within 3 years of application, or after 3 years if the Administrator determines that the circumstances render the applicant unfit to be licensed;

(6) Is or would be operating in violation or circumvention of any Federal, State, or local laws; or

(7) Has made any false or fraudulent statements or provided any false or fraudulent records to the Department or other government agencies, or has pled nolo contendere (no contest) or has

(b) Any person whose license has been revoked shall not be licensed or registered, in his or her own name or in any other manner, and no partnership, firm, corporation, or other legal entity in which any such person has a substantial interest, financial or otherwise, will be licensed or registered.

(c) Any person whose license has been suspended or revoked shall not buy, sell, transport, exhibit, or deliver for transportation, any animal during the period of suspension or revocation, under any circumstances, whether on his or her behalf or on the behalf of another licensee or registrant.

[85 FR 28796, May 13, 2020]
§ 2.12 Termination of a license.

A license may be terminated at any time for any reason that a license application may be denied pursuant to § 2.11 after a hearing in accordance with the applicable rules of practice in 7 CFR part 1.

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§ 2.13 Appeal of inspection report.

Except as otherwise provided in § 2.3(c), any licensee or registrant may appeal all or part of the inspection findings in an inspection report to the Deputy Administrator. To appeal, the licensee or registrant must send a written statement contesting the inspection finding(s) and include any documentation or other information in support of the appeal. To receive consideration, the appeal must be received by the Deputy Administrator within 21 days of the date the licensee or registrant received the inspection report that is the subject of the appeal.

[85 FR 28797, May 13, 2020]

Subpart B—Registration

§ 2.25 Requirements and procedures.

(a) Each carrier and intermediate handler, and each exhibitor not required to be licensed under section 3 of the Act and the regulations of this subchapter, shall register with the Secretary by completing and filing a properly executed form which will be furnished, upon request, by the Deputy
Administrator. The registration form shall be filed with the Deputy Administrator for the State in which the registrant has his or her principal place of business, and shall be updated every 3 years by the completion and filing of a new registration form which will be provided by the Deputy Administrator.

(b) A subsidiary of a business corporation, rather than the parent corporation, will be registered as an exhibitor unless the subsidiary is under such direct control of the parent corporation that the Secretary determines that it is necessary that the parent corporation be registered to effectuate the purposes of the Act.

(c) No registrant or person required to be registered shall interfere with, threaten, abuse (including verbally abuse), or harass any APHIS official who is in the course of carrying out his or her duties.


§ 2.26 Acknowledgment of regulations and standards.

APHIS will supply a copy of the regulations and standards in this subchapter with each registration form. The registrant shall acknowledge receipt of and shall agree to comply with the regulations and standards by signing a form provided for this purpose by APHIS, and by filing it with the Deputy Administrator.


§ 2.27 Notification of change of operation.

(a) A registrant shall notify the Deputy Administrator by certified mail of any change in the name, address, or ownership, or other change in operations affecting its status as an exhibitor, carrier, or intermediate handler, within 10 days after making such change.

(b)(1) A registrant which has not used, handled, or transported animals for a period of at least 2 years may be placed in an inactive status by making a written request to the Deputy Administrator a registrant shall notify the Deputy Administrator in writing at least 10 days before using, handling, or transporting animals again after being in an inactive status.

(2) A registrant which goes out of business or which ceases to function as a carrier, intermediate handler, or exhibitor, or which changes its method of operation so that it no longer uses, handles, or transports animals, and which does not plan to use, handle, or transport animals again at any time in the future, may have its registration canceled by making a written request to the Deputy Administrator. The former registrant is responsible for re-registering and demonstrating its compliance with the Act and regulations should it start using, handling, or transporting animals at any time after its registration is canceled.


Subpart C—Research Facilities

§ 2.30 Registration.

(a) Requirements and procedures. (1) Each research facility, other than a Federal research facility, shall register with the Secretary by completing and filing a properly executed form which will be furnished, upon request, by the Deputy Administrator. The registration form shall be filed with the Deputy Administrator. Except as provided in paragraph (a)(2) of this section, where a school or department of a university or college uses or intends to use live animals for research, tests, experiments, or teaching, the university or college rather than the school or department will be considered the research facility and will be required to register with the Secretary. An official who has the legal authority to bind the parent organization shall sign the registration form.

(2) In any situation in which a school or department of a university or college demonstrates to the Secretary that it is a separate legal entity and its operations and administration are independent of those of the university or college, the school or department will be registered rather than the university or college.
(3) A subsidiary of a business corporation, rather than the parent corporation, will be registered as a research facility unless the subsidiary is under such direct control of the parent corporation that the Secretary determines that it is necessary that the parent corporation be registered to effectuate the purposes of the Act.

(b) Acknowledgment of regulations and standards. APHIS will supply a copy of the regulations and standards in this subchapter with each registration form. The research facility shall acknowledge receipt of and shall agree to comply with the regulations and standards by signing a form provided for this purpose by APHIS, and by filing it with the Deputy Administrator.

(c) Notification of change of operation. A research facility shall notify the Deputy Administrator in writing of any change in the name, address, or ownership, or other change in operations affecting its status as a research facility, within 10 days after making such change. The Notification of Change form (APHIS Form 7033) may be used to provide the information.

(d) Duration of a registration and conditions for cancellation of a registration. (1) A research facility that goes out of business or ceases to function as a research facility, or that changes its method of operation so that it no longer uses, handles, or transports animals, and does not plan to use, handle, or transport animals at any time in the future, may have its registration canceled by making a written request to the Deputy Administrator.

(2) If the Deputy Administrator has sufficient evidence showing that a research facility has ceased to function as a research facility, then the Deputy Administrator may cancel the registration on its own, without a written request from the research facility.

(3) If a research facility plans to resume regulated activity, the facility is responsible for submitting a form (APHIS Form 7011A) to reregister at least 10 days prior to it using, handling, or transporting animals. There are no fees associated with such reregistration.

(e) Non-interference with APHIS officials. No research facility shall interfere with, threaten, abuse (including verbally abuse), or harass any APHIS official who is in the course of carrying out his or her duties.

§ 2.31 Institutional Animal Care and Use Committee (IACUC).

(a) The Chief Executive Officer of the research facility shall appoint an Institutional Animal Care and Use Committee (IACUC), qualified through the experience and expertise of its members to assess the research facility’s animal program, facilities, and procedures. Except as specifically authorized by law or these regulations, nothing in this part shall be deemed to permit the IACUC to prescribe methods or set standards for the design, performance, or conduct of actual research or experimentation by a research facility.

(b) IACUC membership. (1) The members of each Committee shall be appointed by the Chief Executive Officer of the research facility;

(2) The Committee shall be composed of a Chairman and at least two additional members;

(3) Of the members of the Committee:

(i) At least one shall be a Doctor of Veterinary Medicine, with training or experience in laboratory animal science and medicine, who has direct or delegated program responsibility for activities involving animals at the research facility;

(ii) At least one shall not be affiliated in any way with the facility other than as a member of the Committee, and shall not be a member of the immediate family of a person who is affiliated with the facility. The Secretary intends that such person will provide representation for general community interests in the proper care and treatment of animals;

(4) If the Committee consists of more than three members, not more than three members shall be from the same administrative unit of the facility.

(c) IACUC functions. With respect to activities involving animals, the IACUC, as an agent of the research facility, shall:
(1) Review, at least once every six months, the research facility’s program for humane care and use of animals, using title 9, chapter I, subchapter A—Animal Welfare, as a basis for evaluation;

(2) Inspect, at least once every six months, all of the research facility’s animal facilities, including animal study areas, using title 9, chapter I, subchapter A—Animal Welfare, as a basis for evaluation; Provided, however, That animal areas containing free-living wild animals in their natural habitat need not be included in such inspection;

(3) Prepare reports of its evaluations conducted as required by paragraphs (c)(1) and (2) of this section, and submit the reports to the Institutional Official of the research facility; Provided, however, That the IACUC may determine the best means of conducting evaluations of the research facility’s programs and facilities; and Provided, further, That no Committee member wishing to participate in any evaluation conducted under this subpart may be excluded. The IACUC may use subcommittees composed of at least two Committee members and may invite ad hoc consultants to assist in conducting the evaluations, however, the IACUC remains responsible for the evaluations and reports as required by the Act and regulations. The reports shall be reviewed and signed by a majority of the IACUC members and must include any minority views. The reports shall be updated at least once every six months upon completion of the required semiannual evaluations and shall be maintained by the research facility and made available to APHIS and to officials of funding Federal agencies for inspection and copying upon request. The reports must contain a description of the nature and extent of the research facility’s adherence to this subchapter, must identify specifically any departures from the provisions of title 9, chapter I, subchapter A—Animal Welfare, and must state the reasons for each departure. The reports must distinguish significant deficiencies from minor deficiencies. A significant deficiency is one which, with reference to Subchapter A, and, in the judgment of the IACUC and the Institutional Official, is or may be a threat to the health or safety of the animals. If program or facility deficiencies are noted, the reports must contain a reasonable and specific plan and schedule with dates for correcting each deficiency. Any failure to adhere to the plan and schedule that results in a significant deficiency remaining uncorrected shall be reported in writing within 15 business days by the IACUC, through the Institutional Official, to APHIS and any Federal agency funding that activity;

(4) Review, and, if warranted, investigate concerns involving the care and use of animals at the research facility resulting from public complaints received and from reports of noncompliance received from laboratory or research facility personnel or employees;

(5) Make recommendations to the Institutional Official regarding any aspect of the research facility’s animal program, facilities, or personnel training;

(6) Review and approve, require modifications in (to secure approval), or withhold approval of those components of proposed activities related to the care and use of animals, as specified in paragraph (d) of this section;

(7) Review and approve, require modifications in (to secure approval), or withhold approval of proposed significant changes regarding the care and use of animals in ongoing activities; and

(8) Be authorized to suspend an activity involving animals in accordance with the specifications set forth in paragraph (d)(6) of this section.

d IACUC review of activities involving animals. (1) In order to approve proposed activities or proposed significant changes in ongoing activities, the IACUC shall conduct a review of those components of the activities related to the care and use of animals and determine that the proposed activities are in accordance with this subchapter unless acceptable justification for a departure is presented in writing; Provided, however, That field studies as defined in part 1 of this subchapter are exempt from this requirement. Further, the IACUC shall determine that the proposed activities or significant changes in ongoing activities meet the following requirements:
§2.31

(i) Procedures involving animals will avoid or minimize discomfort, distress, and pain to the animals;

(ii) The principal investigator has considered alternatives to procedures that may cause more than momentary or slight pain or distress to the animals, and has provided a written narrative description of the methods and sources, e.g., the Animal Welfare Information Center, used to determine that alternatives were not available;

(iii) The principal investigator has provided written assurance that the activities do not unnecessarily duplicate previous experiments;

(iv) Procedures that may cause more than momentary or slight pain or distress to the animals will:
   (A) Be performed with appropriate sedatives, analgesics or anesthetics, unless withholding such agents is justified for scientific reasons, in writing, by the principal investigator and will continue for only the necessary period of time;
   (B) Involve, in their planning, consultation with the attending veterinarian or his or her designee;
   (C) Not include the use of paralytics without anesthesia;

(v) Animals that would otherwise experience severe or chronic pain or distress that cannot be relieved will be painlessly euthanized at the end of the procedure or, if appropriate, during the procedure;

(vi) The animals’ living conditions will be appropriate for their species in accordance with part 3 of this subchapter, and contribute to their health and comfort. The housing, feeding, and nonmedical care of the animals will be directed by the attending veterinarian or other scientist trained and experienced in the proper care, handling, and use of the species being maintained or studied;

(vii) Medical care for animals will be available and provided as necessary by a qualified veterinarian;

(viii) Personnel conducting procedures on the species being maintained or studied will be appropriately qualified and trained in those procedures;

(ix) Activities that involve surgery include appropriate provision for preoperative and post-operative care of the animals in accordance with established veterinary medical and nursing practices. All survival surgery will be performed using aseptic procedures, including surgical gloves, masks, sterile instruments, and aseptic techniques. Major operative procedures on non-rodents will be conducted only in facilities intended for that purpose which shall be operated and maintained under aseptic conditions. Non-major operative procedures and all surgery on rodents do not require a dedicated facility, but must be performed using aseptic techniques. Operative procedures conducted at field sites need not be performed in dedicated facilities, but must be performed using aseptic procedures;

(x) No animal will be used in more than one major operative procedure from which it is allowed to recover, unless:
   (A) Justified for scientific reasons by the principal investigator, in writing;
   (B) Required as routine veterinary procedure or to protect the health or well-being of the animal as determined by the attending veterinarian; or
   (C) In other special circumstances as determined by the Administrator on an individual basis. Written requests and supporting data should be sent to the Animal and Plant Health Inspection Service, Animal Care, 4700 River Road, Unit 84, Riverdale, Maryland 20737–1234;

(xi) Methods of euthanasia used must be in accordance with the definition of the term set forth in 9 CFR part 1, §1.1 of this subchapter, unless a deviation is justified for scientific reasons, in writing, by the investigator.

(2) Prior to IACUC review, each member of the Committee shall be provided with a list of proposed activities to be reviewed. Written descriptions of all proposed activities that involve the care and use of animals shall be available to all IACUC members, and any member of the IACUC may obtain, upon request, full Committee review of those activities. If full Committee review is not requested, at least one member of the IACUC, designated by the chairman and qualified to conduct the review, shall review those activities, and shall have the authority to approve, require modifications in (to
secure approval), or request full Commit-
tee review of any of those activi-
ties. If full Committee review is re-
quested for a proposed activity, ap-
proval of that activity may be granted
only after review, at a convened meet-
ing of a quorum of the IACUC, and with
the approval vote of a majority of the
quorum present. No member may par-
ticipate in the IACUC review or ap-
proval of an activity in which that
member has a conflicting interest (e.g.,
is personally involved in the activity),
except to provide information re-
quested by the IACUC, nor may a mem-
ber who has a conflicting interest con-
tribute to the constitution of a
quorum;

(3) The IACUC may invite consult-
ants to assist in the review of complex
issues arising out of its review of pro-
posed activities. Consultants may not
approve or withhold approval of an ac-
tivity, and may not vote with the
IACUC unless they are also members of
the IACUC;

(4) The IACUC shall notify principal
investigators and the research facility
in writing of its decision to approve or
withhold approval of those activities
related to the care and use of animals,
or of modifications required to secure
IACUC approval. If the IACUC decides
to withhold approval of an activity, it
shall include in its written notification
a statement of the reasons for its deci-
sion and give the principal investigator
an opportunity to respond in person or
in writing. The IACUC may reconsider
its decision, with documentation in
Committee minutes, in light of the in-
formation provided by the principal in-
vestigator;

(5) The IACUC shall conduct com-
plete reviews of activities covered by
this subchapter at appropriate inter-
vals as determined by the IACUC, but
not less than every 3 years. The com-
plete review shall address all require-
m ents related to the care and use of
animals under paragraphs (d) and (e) of
this section. The IACUC shall be pro-
vided a written description of all ac-
tivities that involve the care and use of
animals for review and approval at the
end of the term.

(6) The IACUC may suspend an activ-
ity that it previously approved if it de-
termines that the activity is not being
conducted in accordance with the de-
scription of that activity provided by
the principal investigator and approved
by the Committee. The IACUC may
suspend an activity only after review
of the matter at a convened meeting of
a quorum of the IACUC and with the
suspension vote of a majority of the
quorum present;

(7) If the IACUC suspends an activity
involving animals, the Institutional Of-
ficial, in consultation with the IACUC,
shall review the reasons for suspension,
take appropriate corrective action, and
report that action with a full expla-
nation to APHIS and any Federal agen-
cy funding that activity; and

(8) Proposed activities and proposed
significant changes in ongoing activi-
ties that have been approved by the
IACUC may be subject to further ap-
propriate review and approval by offi-
cials of the research facility. However,
those officials may not approve an ac-
tivity involving the care and use of
animals if it has not been approved by
the IACUC.

(e) A proposal to conduct an activity
involving animals, or to make a sig-
nificant change in an ongoing activity
involving animals, must contain the
following:

(1) Identification of the species and
the approximate number of animals to
be used;

(2) A rationale for involving animals,
and for the appropriateness of the spe-
cies and numbers of animals to be used;

(3) A complete description of the pro-
posed use of the animals;

(4) A description of procedures de-
digned to assure that discomfort and
pain to animals will be limited to that
which is unavoidable for the conduct of
scientifically valuable research, includ-
ing provision for the use of analgesic,
anesthetic, and tranquilizing drugs
where indicated and appropriate to
minimize discomfort and pain to ani-
mals; and

(5) A description of any euthanasia
method to be used.

§ 2.32 Personnel qualifications.

(a) It shall be the responsibility of
the research facility to ensure that all

[54 FR 36147, Aug. 31, 1989, as amended at 59
FR 67611, Dec. 30, 1994; 63 FR 62926, Nov. 10,
1998; 86 FR 66926, Nov. 24, 2021]
scientists, research technicians, animal technicians, and other personnel involved in animal care, treatment, and use are qualified to perform their duties. This responsibility shall be fulfilled in part through the provision of training and instruction to those personnel.

(b) Training and instruction shall be made available, and the qualifications of personnel reviewed, with sufficient frequency to fulfill the research facility’s responsibilities under this section and §2.31.

(c) Training and instruction of personnel must include guidance in at least the following areas:

(1) Humane methods of animal maintenance and experimentation, including:
   (i) The basic needs of each species of animal;
   (ii) Proper handling and care for the various species of animals used by the facility;
   (iii) Proper pre-procedural and post-procedural care of animals; and
   (iv) Aseptic surgical methods and procedures;

(2) The concept, availability, and use of research or testing methods that limit the use of animals or minimize animal distress;

(3) Proper use of anesthetics, analgesics, and tranquilizers for any species of animals used by the facility;

(4) Methods whereby deficiencies in animal care and treatment are reported, including deficiencies in animal care and treatment reported by any employee of the facility. No facility employee, Committee member, or laboratory personnel shall be discriminated against or be subject to any reprisal for reporting violations of any regulation or standards under the Act;

(5) Utilization of services (e.g., National Agricultural Library, National Library of Medicine) available to provide information:
   (i) On appropriate methods of animal care and use;
   (ii) On alternatives to the use of live animals in research;
   (iii) That could prevent unintended and unnecessary duplication of research involving animals; and
   (iv) Regarding the intent and requirements of the Act.

§ 2.33 Attending veterinarian and adequate veterinary care.

(a) Each research facility shall have an attending veterinarian who shall provide adequate veterinary care to its animals in compliance with this section:

(1) Each research facility shall employ an attending veterinarian under formal arrangements. In the case of a part-time attending veterinarian or consultant arrangements, the formal arrangements shall include a written program of veterinary care and regularly scheduled visits to the research facility;

(2) Each research facility shall assure that the attending veterinarian has appropriate authority to ensure the provision of adequate veterinary care and to oversee the adequacy of other aspects of animal care and use; and

(3) The attending veterinarian shall be a voting member of the IACUC; Provided, however, That a research facility with more than one Doctor of Veterinary Medicine (DVM) may appoint to the IACUC another DVM with delegated program responsibility for activities involving animals at the research facility.

(b) Each research facility shall establish and maintain programs of adequate veterinary care that include:

(1) The availability of appropriate facilities, personnel, equipment, and services to comply with the provisions of this subchapter;

(2) The use of appropriate methods to prevent, control, diagnose, and treat diseases and injuries, and the availability of emergency, weekend, and holiday care;

(3) Daily observation of all animals to assess their health and well-being; Provided, however, That daily observation of animals may be accomplished by someone other than the attending veterinarian; and Provided, further, That a mechanism of direct and frequent communication is required so that timely and accurate information on problems of animal health, behavior, and well-being is conveyed to the attending veterinarian;

(4) Guidance to principal investigators and other personnel involved in the care and use of animals regarding handling, immobilization, anesthesia,
analgesia, tranquilization, and euthanasia; and
(5) Adequate pre-procedural and post-procedural care in accordance with current established veterinary medical and nursing procedures.

§ 2.34 [Reserved]

§ 2.35 Recordkeeping requirements.
(a) The research facility shall maintain the following IACUC records:
(1) Minutes of IACUC meetings, including records of attendance, activities of the Committee, and Committee deliberations;
(2) Records of proposed activities involving animals and proposed significant changes in activities involving animals, and whether IACUC approval was given or withheld; and
(3) Records of semiannual IACUC reports and recommendations (including minority views), prepared in accordance with the requirements of § 2.31(c)(3) of this subpart, and forwarded to the Institutional Official.
(b) Every research facility shall make, keep, and maintain records or forms which fully and correctly disclose the following information concerning each live dog or cat purchased or otherwise acquired, owned, held, or otherwise in their possession or under their control, transported, euthanized, sold, or otherwise disposed of by the research facility. The records shall include any offspring born of any animal while in the research facility’s possession or under its control:
(1) The name and address of the person from whom a dog or cat was purchased or otherwise acquired, whether or not the person is required to be licensed or registered under the Act;
(2) The USDA license or registration number of the person if he or she is licensed or registered under the Act;
(3) The vehicle license number and State, and the driver’s license number (or photographic identification card for nondrivers issued by a State) and State of the person, if he or she is not licensed or registered under the Act;
(4) The date of acquisition of each dog or cat;
(5) The official USDA tag number or tattoo assigned to each dog or cat under § 2.38(g) of this subpart;
(6) A description of each dog or cat which shall include:
(i) The species and breed or type of animal;
(ii) The sex;
(iii) The date of birth or approximate age; and
(iv) The color and any distinctive markings;
(7) Any identification number or mark assigned to each dog or cat by the research facility;
(8) If dogs or cats are acquired from any person not licensed or registered under the Act and not a pound or shelter, the research facility must obtain a certification that the animals were born and raised on the person’s premises and that the person has sold fewer than 25 dogs and/or cats that year.
(c) In addition to the information required to be kept and maintained by every research facility concerning each live dog or cat under paragraph (a) of this section, every research facility transporting, selling, or otherwise disposing of any live dog or cat to another person, shall make and maintain records or forms which fully and correctly disclose the following information:
(1) The name and address of the person to whom a live dog or cat is transported, sold, or otherwise disposed of;
(2) The date of transportation, sale, euthanasia, or other disposition of the animal; and
(3) The method of transportation, including the name of the initial carrier or intermediate handler, or if a privately owned vehicle is used to transport the dog or cat, the name of the owner of the privately owned vehicle.
(d)(1) The USDA Interstate and International Certificate of Health Examination for Small Animals (APHIS Form 7001) and Record of Acquisition and Dogs and Cats on Hand (APHIS Form 7005) are forms which may be used by research facilities to keep and maintain the information required by paragraph (b) of this section.
(2) The USDA Interstate and International Certificate of Health Examination for Small Animals (APHIS Form 7001) and Record of Disposition of Dogs and Cats (APHIS Form 7006) are forms which may be used by research
facilities to keep and maintain the information required by paragraph (c) of this section.

(e) One copy of the record containing the information required by paragraphs (b) and (c) of this section shall accompany each shipment of any live dog or cat sold or otherwise disposed of by a research facility; Provided, however, That, except as provided in §2.133 of this part, information that indicates the source and date of acquisition of any dog or cat need not appear on the copy of the record accompanying the shipment. One copy of the record containing the information required by paragraphs (b) and (c) of this section shall be retained by the research facility.

(f) All records and reports shall be maintained for at least three years. Records that relate directly to proposed activities and proposed significant changes in ongoing activities reviewed and approved by the IACUC shall be maintained for the duration of the activity and for an additional three years after completion of the activity. All records shall be available for inspection and copying by authorized APHIS or funding Federal agency representatives at reasonable times. APHIS inspectors will maintain the confidentiality of the information and will not remove the materials from the research facilities' premises unless there has been an alleged violation, they are needed to investigate a possible violation, or for other enforcement purposes. Release of any such materials, including reports, summaries, and photographs that contain trade secrets or commercial or financial information that is privileged or confidential will be governed by applicable sections of the Freedom of Information Act. Whenever the Administrator notifies a research facility in writing that specified records shall be retained pending completion of an investigation or proceeding under the Act, the research facility shall hold those records until their disposition is authorized in writing by the Administrator.

(Approved by the Office of Management and Budget under control number 0579–0036)


§ 2.36 Annual report.

(a) The reporting facility shall be that segment of the research facility, or that department, agency, or instrumentation of the United States that uses or intends to use live animals in research, tests, experiments, or for teaching. Each reporting facility shall submit an annual report to the Deputy Administrator on or before December 1 of each calendar year. The report shall cover the previous Federal fiscal year. The Annual Report of Research Facility (APHIS Form 7023), Continuation Sheet for Annual Report of Research Facility (APHIS Form 7023A), and Annual Report of Research Facility Column E Explanation (APHIS Form 7023B) are forms which may be used to submit the information required by paragraph (b) of this section.

(b) The annual report shall:

(1) Assure that professionally acceptable standards governing the care, treatment, and use of animals, including appropriate use of anesthetic, analgesic, and tranquilizing drugs, prior to, during, and following actual research, teaching, testing, surgery, or experimentation were followed by the research facility;

(2) Assure that each principal investigator has considered alternatives to painful procedures;

(3) Assure that the facility is adhering to the standards and regulations under the Act, and that it has required that exceptions to the standards and regulations be specified and explained by the principal investigator and approved by the IACUC. A summary of all such exceptions must be attached to the facility's annual report. In addition to identifying the IACUC-approved exceptions, this summary must include a brief explanation of the exceptions, as well as the species and number of animals affected;

(4) State the location of all facilities where animals were housed or used in
actual research, testing, teaching, or experimentation, or held for these purposes;

(5) State the common names and the numbers of animals upon which teaching, research, experiments, or tests were conducted involving no pain, distress, or use of pain-relieving drugs. Routine procedures (e.g., injections, tattooing, blood sampling) should be reported with this group;

(6) State the common names and the numbers of animals upon which experiments, teaching, research, surgery, or tests were conducted involving accompanying pain or distress to the animals and for which appropriate anesthetic, analgesic, or tranquillizing drugs were used;

(7) State the common names and the numbers of animals upon which teaching, experiments, research, surgery, or tests were conducted involving accompanying pain or distress to the animals and for which the use of appropriate anesthetic, analgesic, or tranquillizing drugs would have adversely affected the procedures, results, or interpretation of the teaching, research, experiments, surgery, or tests. An explanation of the procedures producing pain or distress in these animals and the reasons such drugs were not used shall be attached to the annual report;

(8) State the common names and the numbers of animals being bred, conditioned, or held for use in teaching, testing, experiments, research, or surgery but not yet used for such purposes.


§ 2.37 Federal research facilities.

Each Federal research facility shall establish an Institutional Animal Care and Use Committee which shall have the same composition, duties, and responsibilities required of nonfederal research facilities by §2.31 with the following exceptions:

(a) The Committee shall report deficiencies to the head of the Federal agency conducting the research rather than to APHIS; and

(b) The head of the Federal agency conducting the research shall be responsible for all corrective action to be taken at the facility and for the granting of all exceptions to inspection protocol.

§ 2.38 Miscellaneous.

(a) Information as to business: furnishing of same by research facilities. Each research facility shall furnish to any APHIS official any information concerning the business of the research facility which the APHIS official may request in connection with the enforcement of the provisions of the Act, the regulations, and the standards in this subchapter. The information shall be furnished within a reasonable time and as may be specified in the request for information.

(b) Access and inspection of records and property. (1) Each research facility shall, during business hours, allow APHIS officials:

(i) To enter its place of business;

(ii) To examine records required to be kept by the Act and the regulations in this part;

(iii) To make copies of the records;

(iv) To inspect the facilities, property, and animals, as the APHIS officials consider necessary to enforce the provisions of the Act, the regulations, and the standards in this subchapter; and

(v) To document, by the taking of photographs and other means, conditions and areas of noncompliance.

(2) The use of a room, table or other facilities necessary for the proper examination of the records and for inspection of the property or animals shall be extended to APHIS officials by the research facility.

(c) Publication of lists of research facilities subject to the provisions of this part. APHIS will publish on its website lists of research facilities registered in accordance with the provisions of this subpart. The lists may also be obtained upon request from the Deputy Administrator.

(d) Inspection for missing animals. Each research facility shall allow, upon request and during business hours, police or officers of other law enforcement agencies with general law enforcement authority (not those agencies whose duties are limited to enforcement of local animal regulations) to enter its place of business to inspect
§ 2.38 9 CFR Ch. I (1–1–22 Edition)

animals and records for the purpose of seeking animals that are missing, under the following conditions:

(1) The police or other law officer shall furnish to the research facility a written description of the missing animal and the name and address of its owner before making a search;

(2) The police or other law officer shall abide by all security measures required by the research facility to prevent the spread of disease, including the use of sterile clothing, footwear, and masks where required, or to prevent the escape of an animal.

(e) Confiscation and destruction of animals. (1) If an animal being held by a research facility is not being used to carry out research, testing, or experimentation, and is found by an APHIS official to be suffering as a result of the failure of the research facility to comply with any provision of the regulations or the standards set forth in this subchapter, the APHIS official shall make a reasonable effort to notify the research facility of the condition of the animal(s) and request that the condition be corrected and that adequate care be given to alleviate the animal’s suffering or distress, or that the animal(s) be destroyed by euthanasia. In the event that the research facility refuses to comply with this request, the APHIS official may confiscate the animal(s) for care, treatment, or disposal as indicated in paragraph (e)(2) of this section, if, in the opinion of the Administrator, the circumstances indicate the animal’s health is in danger.

(2) In the event that the APHIS official is unable to locate or notify the research facility as required in this section, the APHIS official shall contact a local police or other law officer to accompany him or her to the premises and shall provide for adequate care when necessary to alleviate the animal’s suffering. If, in the opinion of the Administrator, the condition of the animal(s) cannot be corrected by this temporary care, the APHIS official shall confiscate the animal(s).

(3) Confiscated animals may be placed, by sale or donation, with other registrants or licensees that comply with the standards and regulations and can provide proper care, or they may be euthanized. The research facility from which the animals were confiscated shall bear all costs incurred in performing the placement or euthanasia activities authorized by this section.

(f) Handling. (1) Handling of all animals shall be done as expeditiously and carefully as possible in a manner that does not cause trauma, overheating, excessive cooling, behavioral stress, physical harm, or unnecessary discomfort.

(2)(i) Physical abuse shall not be used to train, work, or otherwise handle animals.

(ii) Deprivation of food or water shall not be used to train, work, or otherwise handle animals; Provided, however: That the short-term withholding of food or water from animals, when specified in an IACUC-approved activity that includes a description of monitoring procedures, is allowed by these regulations.

(g) Identification of dogs and cats. (1) All live dogs or cats, including those from any exempt source, delivered for transportation, transported, purchased or otherwise acquired, sold, or disposed of by a research facility, shall be identified at the time of such delivery for transportation, purchase, sale, disposal, or acquisition in one of the following ways:

(i) By the official tag or tattoo which was affixed to the animal at the time it was acquired by the research facility, as required by this section; or

(ii) By a tag, tattoo, or collar, applied to the live dog or cat by the research facility and which individually identifies the dog or cat by number.

(2) All official tag or tattoo numbers shall be correctly listed in the records of purchase, acquisition, disposal, or sale which shall be maintained in accordance with §2.35.

(3) Unweaned puppies or kittens need not be individually identified while they are maintained as a litter with their dam in the same primary enclosure, provided the dam has been individually identified.

(4) The official tag shall be made of a durable alloy such as brass, bronze, or steel, or of a durable plastic. Aluminum of a sufficient thickness to assure the tag is durable and legible may also be used. The tag may be circular
A list of the commercial manufacturers who produce these tags and are known to the Department may be obtained from the Deputy Administrator. Any manufacturer who desires to be included in the list should notify the Administrator.

1 A list of the commercial manufacturers who produce these tags and are known to the Department may be obtained from the Deputy Administrator. Any manufacturer who desires to be included in the list should notify the Administrator.

(5) Each tag shall have the following information embossed or stamped on so that it is easily readable:

(i) The letters “USDA”;

(ii) Numbers identifying the State and dealer, exhibitor, or research facility (e.g., 39-AB); and

(iii) Numbers identifying the animal (e.g., 82488).

(6) Official tags shall be serially numbered and shall be applied to dogs or cats in the manner set forth in this section in as close to consecutive numerical order as possible. No tag number shall be used to identify more than one animal or shall be reused within a 5-year period.

(7) Research facilities may obtain, at their own expense, official tags from commercial tag manufacturers.1 At the time the research facility is registered, the Department will assign identification letters and numbers to be used on the official tags.

(8) Each research facility shall be held accountable for all official tags acquired. In the event an official tag is lost from a dog or cat while in the possession of a research facility, the facility shall make a diligent effort to locate and reapply the tag to the proper animal. If the lost tag is not located, the research facility shall affix another official tag to the animal in the manner prescribed in this section and record the tag number on the official records.

(9) When a dog or cat wearing or identified by an official tag arrives at a research facility, the facility may continue to use that tag to identify the dog or cat or the tag may be replaced as indicated in paragraph (g)(1) of this section. All tags removed by a research facility shall be retained and disposed of as indicated in this section.

(10) Where a dog or cat to which is affixed or which is identified by an official tag is euthanized, or dies from other causes, the research facility shall remove and retain the tag for the required period, as set forth in paragraph (g)(11) of this section.

(11) All official tags removed and retained by a research facility shall be held until called for by an APHIS official or for a period of 1 year.

(12) When official tags are removed from animals for disposal, the tags must be disposed of so as to preclude their reuse for animal identification. No animal identification number shall be used within any 5-year period following its previous use.

(h) Health certification. (1) No research facility, including a Federal research facility, shall deliver to any intermediate handler or carrier for transportation, in commerce, or shall transport in commerce any dog, cat, or nonhuman primate unless the dog, cat, or nonhuman primate is accompanied by a health certificate executed and issued by a licensed veterinarian. The health certificate shall state that:

(i) The licensed veterinarian inspected the dog, cat, or nonhuman primate on a specified date which shall not be more than 10 days prior to the delivery of the dog, cat, or nonhuman primate for transportation; and

(ii) When so inspected, the dog, cat, or nonhuman primate appeared to the licensed veterinarian to be free of any infectious disease or physical abnormality which would endanger the animal(s) or other animals or endanger public health.

(2) The Secretary may provide exceptions to the health certification requirement on an individual basis for animals shipped to a research facility for purposes of research, testing, or experimentation when the research facility requires animals not eligible for certification. Requests should be addressed to the Animal and Plant Health Inspection Service, Animal Care, 4700 River Road, Unit 84, Riverdale, Maryland 20737–1234.

(3) The U.S. Interstate and International Certificate of Health Examination for Small Animals (APHIS Form 7001) may be used for health certification by a licensed veterinarian as required by this section.

(1) Holding of animals. If any research facility obtains prior approval of the
Deputy Administrator, it may arrange to have another person hold animals: Provided, That:

(1) The other person agrees, in writing, to comply with the regulations in this part and the standards in part 3 of this subchapter, and to allow inspection of the premises by an APHIS official during business hours;

(2) The animals remain under the total control and responsibility of the research facility; and

(3) The Institutional Official agrees, in writing, that the other person or premises is a recognized animal site under its research facility registration. APHIS Form 7009 shall be used for approval.

(4) The other person or premises must either be directly included in the research facility’s contingency plan required under paragraph (l) of this section or develop its own contingency plan in accordance with paragraph (l) of this section.

(j) Holding period. Research facilities that obtain dogs and cats from sources other than dealers, exhibitors, and exempt persons shall hold the animals for 5 full days, not including the day of acquisition, before they may be used by the facility. Research facilities shall comply with the identification of animals requirements set forth in §2.38(g) during this period.

(k) Compliance with standards and prohibitions. (1) Each research facility shall comply in all respects with the regulations set forth in subpart C of this part and the standards set forth in part 3 of this subchapter for the humane handling, care, treatment, housing, and transportation of animals; Provided, however. That exceptions to the standards in part 3 and the provisions of subpart C of this part may be made only when such exceptions are specified and justified in the proposal to conduct the activity and are approved by the IACUC.

(2) No person shall obtain live dogs or cats by use of false pretenses, misrepresentation, or deception.

(3) No person shall acquire, buy, sell, exhibit, use for research, transport, or offer for transportation, any stolen animal.

(4) Each research facility shall comply with the regulations set forth in §2.133 of subpart I of this part.

(l) Contingency planning. (1) Research facilities must develop, document, and follow an appropriate plan to provide for the humane handling, treatment, transportation, housing, and care of their animals in the event of an emergency or disaster (one which could reasonably be anticipated and expected to be detrimental to the good health and well-being of the animals in their possession). Such contingency plans must:

(i) Identify situations the facility might experience that would trigger the need for the measures identified in a contingency plan to be put into action including, but not limited to, emergencies such as electrical outages, faulty HVAC systems, fires, and animal escapes, as well as natural disasters the facility is most likely to experience.

(ii) Outline specific tasks required to be carried out in response to the identified emergencies or disasters including, but not limited to, detailed animal evacuation instructions or shelter-in-place instructions and provisions for providing backup sources of food and water as well as sanitation, ventilation, bedding, veterinary care, etc.;

(iii) Identify a chain of command and who (by name or by position title) will be responsible for fulfilling these tasks; and

(iv) Address how response and recovery will be handled in terms of materials, resources, and training needed.

(2) For current registrants, the contingency plan must be in place by July 29, 2013. For research facilities registered after this date, the contingency plan must be in place prior to conducting regulated activities. The plan must be reviewed by the research facility on at least an annual basis to ensure that it adequately addresses the criteria listed in paragraph (l)(1) of this section. Each registrant must maintain documentation of their annual reviews, including documenting any amendments or changes made to their plan since the previous year’s review, such as changes made as a result of recently predicted, but historically unforeseen, circumstances (e.g., weather extremes).
Contingency plans, as well as all annual review documentation and training records, must be made available to APHIS and any funding Federal agency representatives upon request. Facilities maintaining or otherwise handling marine mammals in captivity must also comply with the requirements of §3.101(b) of this subchapter.

(3) The facility must provide and document participation in and successful completion of training for its personnel regarding their roles and responsibilities as outlined in the plan. For current registrants, training of facility personnel must be completed within 60 days of the research facility putting their plan in place; for research facilities registered after July 5, 2022, training of facility personnel must be completed within 60 days of the facility putting its contingency plan in place. This deadline applies to employees hired before and up to 30 days after the facility puts its contingency plan in place. For employees hired more than 30 days after the facility puts its contingency plan in place, training must be conducted within 30 days of their start date. Any substantive changes to the plan as a result of the annual review must be communicated to employees through training which must be conducted within 30 days of making the changes.

(Approved by the Office of Management and Budget under control number 0579–0479)

Subpart D—Attending Veterinarian and Adequate Veterinary Care

§ 2.40 Attending veterinarian and adequate veterinary care (dealers and exhibitors).

(a) Each dealer or exhibitor shall have an attending veterinarian who shall provide adequate veterinary care to its animals in compliance with this section.

(1) Each dealer and exhibitor shall employ an attending veterinarian under formal arrangements. In the case of a part-time attending veterinarian or consultant arrangements, the formal arrangements shall include a written program of veterinary care and regularly scheduled visits to the premises of the dealer or exhibitor; and

(2) Each dealer and exhibitor shall assure that the attending veterinarian has appropriate authority to ensure the provision of adequate veterinary care and to oversee the adequacy of other aspects of animal care and use.

(b) Each dealer or exhibitor shall establish and maintain programs of adequate veterinary care that include:

(1) The availability of appropriate facilities, personnel, equipment, and services to comply with the provisions of this subchapter;
In general, well fitted collars made of leather or plastic will be acceptable under this provision. The use of certain types of chains presently used by some dealers may also be deemed acceptable. APHIS will determine the acceptability of a material proposed for usage as collars from the standpoint of humane considerations on an individual basis in consultation with the dealer or exhibitor involved. The use of materials such as wire, elastic, or sharp metal that might cause discomfort or injury to the dogs or cats is not acceptable.

2 In general, well fitted collars made of leather or plastic will be acceptable under this provision. The use of certain types of chains presently used by some dealers may also be deemed acceptable. APHIS will determine the acceptability of a material proposed for usage as collars from the standpoint of humane considerations on an individual basis in consultation with the dealer or exhibitor involved. The use of materials such as wire, elastic, or sharp metal that might cause discomfort or injury to the dogs or cats is not acceptable.
(ii) A distinctive and legible tattoo marking approved by the Administrator; or
(iii) A plastic-type collar acceptable to the Administrator which has legibly placed thereon the information required for an official tag pursuant to §2.51.

(4) When any dealer has made a reasonable effort to affix an official tag to a cat, as set forth in paragraphs (a) and (b) of this section, and has been unable to do so, or when the cat exhibits serious distress from the attachment of a collar and tag, the dealer shall attach the collar and tag to the door of the primary enclosure containing the cat and take measures adequate to maintain the identity of the cat in relation to the tag. Each primary enclosure shall contain no more than one weaned cat without an affixed collar and official tag, unless the cats are identified by a distinctive and legible tattoo or plastic-type collar approved by the Administrator.

(c) A class “C” exhibitor shall identify all live dogs and cats under his or her control or on his or her premises, whether held, purchased, or otherwise acquired:
(1) As set forth in paragraph (b)(1) or (b)(3) of this section, or
(2) By identifying each dog or cat with:
   (i) An official USDA sequentially numbered tag that is kept on the door of the animal’s cage or run;
   (ii) A record book containing each animal’s tag number, a written description of each animal, the data required by §2.75(a), and a clear photograph of each animal; and
   (iii) A duplicate tag that accompanies each dog or cat whenever it leaves the compound or premises.

(e) Unweaned puppies or kittens need not be individually identified as required by paragraphs (a) and (b) of this section while they are maintained as a litter with their dam in the same primary enclosure, provided the dam has been individually identified.

(ii) All animals, except dogs and cats, delivered for transportation, transported, purchased, sold, or otherwise acquired or disposed of by any dealer or exhibitor shall be identified by the dealer or exhibitor at the time of delivery for transportation, purchase, sale, acquisition or disposal, as provided for in this paragraph and in records maintained as required in §§2.75 and 2.77.

(2) When one or more animals, other than dogs or cats, are confined in a primary enclosure, the animal(s) shall be identified by:
   (i) A label attached to the primary enclosure which shall bear a description of the animals in the primary enclosure, including:
      (A) The number of animals;
      (B) The species of the animals;
      (C) Any distinctive physical features of the animals; and
   (ii) Marking the primary enclosure with a painted or stenciled number which shall be recorded in the records of the dealer or exhibitor together with:
      (A) A description of the animal(s);
      (B) The species of the animal(s); and
      (C) Any distinctive physical features of the animal(s); or
   (iii) A tag or tattoo applied to each animal in the primary enclosure by the dealer or exhibitor which individually identifies each animal by description or number.

(3) When any animal, other than a dog or cat, is not confined in a primary enclosure, it shall be identified on a record, as required by §2.75, which shall accompany the animal at the time it is delivered for transportation, transported, purchased, or sold, and shall be kept and maintained by the dealer or exhibitor as part of his or her records.

§2.51 Form of official tag.

(a) The official tag shall be made of a durable alloy such as brass, bronze, or steel, or of a durable plastic. Aluminum of a sufficient thickness to assure the tag is durable and legible may also be used. The tag shall be one of the following shapes:
(1) Circular in shape and not less than 1¼ inches in diameter, or
(2) Oblong and flat in shape, not less than 2 inches by ¾ inch and riveted to an acceptable collar.

(b) Each tag shall have the following information embossed or stamped on so that it is easily readable:
subsection 2.52 How to obtain tags.

Dealers or exhibitors may obtain, at their own expense, official tags from commercial tag manufacturers.\(^4\) At the time the dealer or exhibitor is issued a license or is registered, the Department will assign identification letters and numbers and inform them of the identification letters and numbers to be used on the official tags.

subsection 2.53 Use of tags.

Official tags obtained by a dealer, exhibitor, or research facility, shall be applied to dogs or cats in the manner set forth in §2.50 and in as close to consecutive numerical order as possible. No tag number shall be used to identify more than one animal. No number shall be repeated within a 5-year period.

subsection 2.54 Lost tags.

Each dealer or exhibitor shall be held accountable for all official tags acquired. In the event an official tag is lost from a dog or cat while in the possession of a dealer or exhibitor, the dealer or exhibitor shall make a diligent effort to locate and reapply the tag to the proper animal. If the lost tag is not located, the dealer or exhibitor shall affix another official tag to the animal in the manner prescribed in §2.50, and record the tag number on the official records.

\(^4\) A list of the commercial manufacturers who produce these tags and are known to the Department may be obtained from the Deputy Administrator. Any manufacturer who desires to be included in the list should notify the Administrator.

subsection 2.55 Removal and disposal of tags.

(a) Where a dog or cat to which is affixed or which is identified by an official tag is euthanized, or dies from other causes, the dealer or exhibitor shall remove and retain the tag for the required period, as set forth in paragraph (b) of this section.

(b) All official tags removed and retained by a dealer or exhibitor shall be held until called for by an APHIS official or for a period of 1 year.

(c) When official tags are removed from animals for disposal, the tags must be disposed of so as to preclude their reuse for animal identification. No animal identification number shall be used within any 5-year period following its previous use.

subsection 2.60 Prohibition on the purchase, sale, use, or transportation of stolen animals.

No person shall buy, sell, exhibit, use for research, transport, or offer for transportation, any stolen animal.

subsection 2.75 Records: Dealers and exhibitors.

(a)(1) Each dealer, other than operators of auction sales and brokers to whom animals are consigned, and each exhibitor shall make, keep, and maintain records or forms which fully and correctly disclose the following information concerning each dog or cat purchased or otherwise acquired whether or not the person is required to be licensed or registered under the Act:

(i) The name and address of the person from whom a dog or cat was purchased or otherwise acquired whether or not the person is required to be licensed or registered under the Act;

(ii) The USDA license or registration number of the person if he or she is licensed or registered under the Act;

(iii) The vehicle license number and State, and the driver’s license number...
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(or photographic identification card for nondrivers issued by a State) and State of the person, if he or she is not licensed or registered under the Act;

(iv) The name and address of the person to whom a dog or cat was sold or given and that person’s license or registration number if he or she is licensed or registered under the Act;

(v) The date a dog or cat was acquired or disposed of, including by euthanasia;

(vi) The official USDA tag number or tattoo assigned to a dog or cat under §§2.50 and 2.54;

(vii) A description of each dog or cat which shall include:

(A) The species and breed or type;
(B) The sex;
(C) The date of birth or approximate age; and
(D) The color and any distinctive markings;

(viii) The method of transportation including the name of the initial carrier or intermediate handler or, if a privately owned vehicle is used to transport a dog or cat, the name of the owner of the privately owned vehicle;

(ix) The date and method of disposition of a dog or cat, e.g., sale, death, euthanasia, or donation.

(2) Each dealer and exhibitor shall use Record of Acquisition and Dogs and Cats on Hand (APHIS Form 7005) and Record of Disposition of Dogs and Cats (APHIS Form 7006) to make, keep, and maintain the information required by paragraph (a)(1) of this section: Provided, that if a dealer or exhibitor who uses a computerized recordkeeping system believes that APHIS Form 7005 and APHIS Form 7006 are unsuitable for him or her to make, keep, and maintain the information required by paragraph (a)(1) of this section, APHIS will advise the person as to the disposition of his or her request for a variance from the requirement to use APHIS Form 7005 and APHIS Form 7006.

(i) The request for a variance must consist of a written statement describing why APHIS Form 7005 and APHIS Form 7006 are unsuitable for the dealer or exhibitor to make, keep, and maintain the information required by paragraph (a)(1) of this section, and a description of the computerized recordkeeping system the person would use in lieu of APHIS Form 7005 and APHIS Form 7006 to make, keep, and maintain the information required by paragraph (a)(1) of this section. APHIS will advise the person as to the disposition of his or her request for a variance from the requirement to use APHIS Form 7005 and APHIS Form 7006.

(ii) A dealer or exhibitor whose request for a variance has been denied may request a hearing in accordance with the applicable rules of practice for the purpose of showing why the request for a variance should not be denied. The denial of the variance shall remain in effect until the final legal decision has been rendered.

(3) The USDA Interstate and International Certificate of Health Examination for Small Animals (APHIS Form 7001) may be used by dealers and exhibitors to make, keep, and maintain the information required by §2.78.

(4) One copy of the record containing the information required by paragraph (a)(1) of this section shall accompany each shipment of any dog or cat purchased or otherwise acquired by a dealer or exhibitor. One copy of the record containing the information required by paragraph (a)(1) of this section shall accompany each shipment of any dog or cat sold or otherwise disposed of by a dealer or exhibitor: Provided, however, that, except as provided in §2.133(b) of this part for dealers, information that indicates the source and date of acquisition of a dog or cat need not appear on the copy of the record accompanying the shipment. One copy of the record containing the information required by paragraph (a)(1) of this section shall be retained by the dealer or exhibitor.

(b) Every dealer other than operators of auction sales and brokers to whom animals are consigned, and exhibitor shall make, keep, and maintain records or forms which fully and correctly disclose the following information concerning animals other than dogs and cats, purchased or otherwise acquired, owned, held, leased, or otherwise in his or her possession or under his or her control, or which is transported, sold, euthanized, or otherwise disposed of by that dealer or exhibitor.

(b)(1) The records shall include any offspring born of any animal while in his or her possession or under his or her control.
§ 2.76 Records: Operators of auction sales and brokers.

(a) Every operator of an auction sale or broker shall make, keep, and maintain records or forms which fully and correctly disclose the following information concerning each animal consigned for auction or sold, whether or not a fee or commission is charged:

(1) The name and address of the person who owned or consigned the animal(s) for sale;

(2) The name and address of the buyer or consignee who received the animal;

(3) The USDA license or registration number of the person(s) selling, consigning, buying, or receiving the animals if he or she is licensed or registered under the Act;

(4) The vehicle license number and State, and the driver’s license number (or photographic identification card for non-drivers issued by a State) and State of the person, if he or she is not licensed or registered under the Act;

(5) The date of the consignment;

(6) The official USDA tag number or tattoo assigned to the animal under §§2.50 and 2.54;

(7) A description of the animal which shall include:
   (i) The species and breed or type of animal;
   (ii) The sex of the animal; and
   (iii) The date of birth or approximate age; and
   (iv) The color and any distinctive markings;

(8) The auction sales number or records number assigned to the animal.

(b) One copy of the record containing the information required by paragraph (a) of this section shall be given to the consignor of each animal, one copy of the record shall be given to the purchaser of each animal: Provided, however, That information which indicates the source and date of consignment of any animal need not appear on the copy of the record given the purchaser of any animal. One copy of the record containing the information required by paragraph (a) of this section shall be retained by the operator of such auction sale, or broker, for each animal sold by the auction sale or broker.


§ 2.76 Operators of auction sales.

(i) The name and address of the person from whom the animals were purchased or otherwise acquired;

(ii) The USDA license or registration number of the person if he or she is licensed or registered under the Act;

(iii) The vehicle license number and State, and the driver’s license number (or photographic identification card for non-drivers issued by a State) and State of the person, if he or she is not licensed or registered under the Act;

(iv) The name and address of the person to whom an animal was sold or given;

(v) The date of purchase, acquisition, sale, or disposal of the animal(s);

(vi) The species of the animal(s); and

(vii) The number of animals in the shipment.

(2) Record of Animals on Hand (other than dogs and cats) (APHIS Form 7019) and Record of Acquisition, Disposition, or Transport of Animals (other than dogs and cats) (APHIS Form 7020) are forms which may be used by dealers and exhibitors to keep and maintain the information required by paragraph (b)(1) of this section concerning animals other than dogs and cats except as provided in §2.78.

(3) One copy of the record containing the information required by paragraph (b)(1) of this section shall accompany each shipment of any animal other than a dog or cat purchased or otherwise acquired by a dealer or exhibitor. One copy of the record containing the information required by paragraph (b)(1) of this section shall accompany each shipment of any animal other than a dog or cat sold or otherwise disposed of by a dealer or exhibitor: Provided, however, That information which indicates the source and date of acquisition of any animal other than a dog or cat need not appear on the copy of the record accompanying the shipment. The dealer or exhibitor shall retain one copy of the record containing the information required by paragraph (b)(1) of this section.

§ 2.77 Records; Carriers and intermediate handlers.

(a) In connection with all live animals accepted for shipment on a C.O.D. basis or other arrangement or practice under which the cost of an animal or the transportation of an animal is to be paid and collected upon delivery of the animal to the consignee, the accepting carrier or intermediate handler, if any, shall keep and maintain a copy of the consignor’s written guarantee for the payment of transportation charges for any animal not claimed as provided in § 2.80, including, where necessary, both the return transportation charges and an amount sufficient to reimburse the carrier for out-of-pocket expenses incurred for the care, feeding, and storage of the animal. The carrier or intermediate handler at destination shall also keep and maintain a copy of the shipping document containing the time, date, and method of each attempted notification and the final notification to the consignee and the name of the person notifying the consignee, as provided in § 2.80.

(b) In connection with all live dogs, cats, or nonhuman primates delivered for transportation, in commerce, to any carrier or intermediate handler, by any dealer, research facility, exhibitor, broker, operator of an auction sale, or department, agency, or instrumentality of the United States or of any State or local government, the accepting carrier or intermediate handler shall keep and maintain a copy of the health certification completed as required by § 2.78, tendered with each live dog, cat, or nonhuman primate.

§ 2.78 Health certification and identification.

(a) No dealer, exhibitor, operator of an auction sale, broker, or department, agency, or instrumentality of the United States or of any State or local government shall deliver to any intermediate handler or carrier for transportation, in commerce, or shall transport in commerce any dog, cat, or nonhuman primate unless and until it is accompanied by a health certificate issued by a licensed veterinarian in accordance with paragraph (a) of this section, or an exemption issued by the Secretary in accordance with paragraph (b) of this section.

(b) The U.S. Interstate and International Certificate of Health Examination for Small Animals (APHIS Form 7001) may be used for health certification by a licensed veterinarian as required by this section.

§ 2.79 C.O.D. shipments.

(a) No carrier or intermediate handler shall accept any animal for transportation, in commerce, upon any
C.O.D. or other basis where any money is to be paid and collected upon delivery of the animal to the consignee, unless the consignor guarantees in writing the payment of all transportation, including any return transportation, if the shipment is unclaimed or the consignee cannot be notified in accordance with paragraphs (b) and (c) of this section, including reimbursing the carrier or intermediate handler for all out-of-pocket expenses incurred for the care, feeding, and storage or housing of the animal.

(b) Any carrier or intermediate handler receiving an animal at a destination on a C.O.D. or other basis where any money is to be paid and collected upon delivery of the animal to the consignee shall attempt to notify the consignee at least once every 6 hours for a period of 24 hours after arrival of the animal at the animal holding area of the terminal cargo facility. The carrier or intermediate handler shall record the time, date, and method of each attempted notification and the final notification to the consignee, and the name of the person notifying the consignee, on the shipping document and on the copy of the shipping document accompanying the C.O.D. shipment. If the consignee cannot be notified of the C.O.D. shipment within 24 hours after its arrival, the carrier or intermediate handler shall return the animal to the consignor, or to whomever the consignor has designated, on the next practical available transportation, in accordance with the written agreement required in paragraph (a) of this section.

(c) It is the responsibility of any carrier or intermediate handler to hold, feed, and care for any animal accepted for transportation, in commerce, under a C.O.D. or other arrangement where any money is to be paid and collected upon delivery of the animal until the consignee accepts shipment at destination or until returned to the consignor or his or her designee should the consignee fail to accept delivery of the animal or if the consignee could not be notified as prescribed in paragraph (b) of this section.

(d) Nothing in this section shall be construed as prohibiting any carrier or intermediate handler from requiring any guarantee in addition to that required in paragraph (a) of this section for the payment of the cost of any transportation or out-of-pocket or other incidental expenses incurred in the transportation of any animal.

§ 2.80 Records, disposition.

(a) No dealer, exhibitor, broker, operator of an auction sale, carrier, or intermediate handler shall, for a period of 1 year, destroy or dispose of, without the consent in writing of the Administrator, any books, records, documents, or other papers required to be kept and maintained under this part.

(b) Unless otherwise specified, the records required to be kept and maintained under this part shall be held for 1 year after an animal is euthanized or disposed of and for any period in excess of one year as necessary to comply with any applicable Federal, State, or local law. Whenever the Administrator notifies a dealer, exhibitor, broker, operator of an auction sale, carrier, or intermediate handler in writing that specified records shall be retained pending completion of an investigation or proceeding under the Act, the dealer, exhibitor, broker, operator of an auction sale, carrier, or intermediate handler in writing that specified records shall be retained pending completion of an investigation or proceeding under the Act, the dealer, exhibitor, broker, operator of an auction sale, carrier, or intermediate handler shall hold those records until their disposition is authorized by the Administrator.

Subpart H—Compliance With Standards and Holding Period

§ 2.100 Compliance with standards.

(a) Each dealer, exhibitor, operator of an auction sale, and intermediate handler shall comply in all respects with the regulations set forth in part 2 and
the standards set forth in part 3 of this
subchapter for the humane handling,
care, treatment, housing, and transpor-
tation of animals.

(b) Each carrier shall comply in all
respects with the regulations in part 2
and the standards in part 3 of this sub-
chapter setting forth the conditions
and requirements for the humane
transportation of animals in commerce
and their handling, care, and treat-
ment in connection therewith.

§ 2.101 Holding period.

(a) Any live dog or cat acquired by a
dealer or exhibitor shall be held by
him or her, under his or her super-
vision and control, for a period of not
less than 5 full days, not including the
day of acquisition, after acquiring the
animal, excluding time in transit: Pro-
vided, however:

(1) That any live dog or cat acquired
by a dealer or exhibitor from any pri-
vate or contract animal pound or shel-
ter shall be held by that dealer or ex-
hibitor under his or her supervision
and control for a period of not less
than 10 full days, not including the day
of acquisition, after acquiring the ani-
mal, excluding time in transit;

(2) Live dogs or cats which have com-
pleted a 5-day holding period with an-
other dealer or exhibitor, or a 10-day
holding period with another dealer or
exhibitor if obtained from a private or
contract shelter or pound, may be sold
or otherwise disposed of by subsequent
dealers or exhibitors after a minimum
holding period of 24 hours by each sub-
sequent dealer or exhibitor excluding
time in transit;

(3) Any dog or cat suffering from dis-
ease, emaciation, or injury may be de-
stroyed by euthanasia prior to the
completion of the holding period re-
quired by this section; and

(4) Any live dog or cat, 120 days of
age or less, that was obtained from the
person that bred and raised such dog or
cat, may be exempted from the 5-day
holding requirement and may be dis-
posed of by dealers or exhibitors after a
minimum holding period of 24 hours,
excluding time in transit. Each subse-
quent dealer or exhibitor must also
hold each such dog or cat for a 24-hour
period excluding time in transit.

(b) During the period in which any
dog or cat is being held as required by
this section, the dog or cat shall be un-
loaded from any means of conveyance
in which it was received, for food,
water, and rest, and shall be handled,
cared for, and treated in accordance
with the standards set forth in part 3,
subpart A, of this subchapter and
§2.131.

§ 2.102 Holding facility.

(a) If any dealer or exhibitor obtains
the prior approval of the Deputy Ad-
ministrator, he may arrange to have
another person hold animals for the re-
quired period provided for in paragraph
(a) of §2.101: Provided, That:

(1) The other person agrees in writing
to comply with the regulations in part
2 and the standards in part 3 of this
subchapter and to allow inspection of
his premises by an APHIS official dur-
ing business hours; and

(2) The animals remain under the
total control and responsibility of the
dealer or exhibitor.

(3) Approval will not be given for a
dealer or exhibitor holding a license as
set forth in §2.1 to have animals held
for purposes of this section by another
licensed dealer or exhibitor. APHIS
Form 7009 shall be used for approval.

(4) The other person or premises
must either be directly included in the
dealer’s or exhibitor’s contingency plan
required under §2.134 or must develop
its own contingency plan in accordance
with §2.134.

(b) If any intermediate handler ob-
tains prior approval of the Deputy Ad-
ministrator, it may arrange to have
another person hold animals: Provided,
That:

(1) The other person agrees in writing
to comply with the regulations in part
2 and the standards in part 3 of this
subchapter and to allow inspection of
the premises by an APHIS official dur-
ing business hours; and

(2) The animals remain under the
total control and responsibility of the
research facility or intermediate han-
dler.

(3) The other person or premises
must either be directly included in the

5 An operator of an auction sale is not con-
considered to have acquired a dog or cat which
is sold through the auction sale.
§ 2.125 Information as to business; furnishing of same by dealers, exhibitors, operators of auction sales, intermediate handlers, and carriers.

Each dealer, exhibitor, operator of an auction sale, intermediate handler, and carrier shall furnish to any APHIS official any information concerning the business of the dealer, exhibitor, operator of an auction sale, intermediate handler or carrier which the APHIS official may request in connection with the enforcement of the provisions of the Act, the regulations and the standards in this subchapter. The information shall be furnished within a reasonable time and as may be specified in the request for information.

§ 2.126 Access and inspection of records and property; submission of itineraries.

(a) Each dealer, exhibitor, intermediate handler, or carrier, shall, during business hours, allow APHIS officials:
   (1) To enter its place of business;
   (2) To examine records required to be kept by the Act and the regulations in this part;
   (3) To make copies of the records;
   (4) To inspect and photograph the facilities, property and animals, as the APHIS officials consider necessary to enforce the provisions of the Act, the regulations and the standards in this subchapter; and
   (5) To document, by the taking of photographs and other means, conditions and areas of noncompliance.

(b) The use of a room, table, or other facilities necessary for the proper examination of the records and inspection of the property or animals must be extended to APHIS officials by the dealer, exhibitor, intermediate handler or carrier, and a responsible adult shall be made available to accompany APHIS officials during the inspection process.

(c) Any person who is subject to the Animal Welfare regulations and who intends to exhibit any animal at any location other than the person’s approved site (including, but not limited to, circuses, traveling educational exhibits, animal acts, and petting zoos), except for travel that does not extend overnight, shall submit a written itinerary to the Deputy Administrator. The itinerary shall be received by the Deputy Administrator no fewer than 2 days in advance of any travel and shall contain complete and accurate information concerning the whereabouts of any animal intended for exhibition at any location other than the person’s approved site. If the exhibitor accepts an engagement for which travel will begin with less than 48 hours’ notice, the exhibitor shall immediately contact the Deputy Administrator in writing with the required information. APHIS expects such situations to occur infrequently, and exhibitors who repeatedly provide less than 48 hours’ notice will, after notice by APHIS, be subject to increased scrutiny under the Act.

(1) The itinerary shall include the following:
   (i) The name of the person who intends to exhibit the animal and transport the animal for exhibition purposes, including any business name and current Act license or registration number and, in the event that any animal is leased, borrowed, loaned, or under some similar arrangement, the name of the person who owns such animal;
   (ii) The name, identification number or identifying characteristics, species (common or scientific name), sex and age of each animal; and
   (iii) The names, dates, and locations (with addresses) where the animals will travel, be housed, and be exhibited, including all anticipated dates and locations (with addresses) for any stops and layovers that allow or require removal of the animals from the transport enclosures. Unanticipated delays of such length shall be reported to the Deputy Administrator the next APHIS business day. APHIS Regional offices are
available each weekday, except on Federal holidays, from 8 a.m. to 5 p.m.  
(2) The itinerary shall be revised as necessary, and the Deputy Administrator shall be notified of any changes. If initial notification of a change due to an emergency is made by a means other than email or facsimile, it shall be followed by written documentation at the earliest possible time. For changes that occur after normal APHIS business hours, the change shall be conveyed to the Deputy Administrator no later than the following APHIS business day. APHIS Regional offices are available each weekday, except on Federal holidays, from 8 a.m. to 5 p.m.

(Approved by the Office of Management and Budget under control number 0579–0361)


§ 2.127 Publication of lists of persons subject to the provisions of this part.

APHIS will publish on its website lists of persons licensed or registered in accordance with the provisions of this part. The lists may also be obtained upon request from the Deputy Administrator.

[85 FR 28798, May 13, 2020]

§ 2.128 Inspection for missing animals.

Each dealer, exhibitor, intermediate handler and carrier shall allow, upon request and during business hours, police or officers of other law enforcement agencies with general law enforcement authority (not those agencies whose duties are limited to enforcement of local animal regulations) to enter his or her place of business to inspect animals and records for the purpose of seeking animals that are missing, under the following conditions:

(a) The police or other law officer shall furnish to the dealer, exhibitor, intermediate handler or carrier a written description of the missing animal and the name and address of its owner before making a search.

(b) The police or other law officer shall abide by all security measures required by the dealer, exhibitor, intermediate handler or carrier to prevent the spread of disease, including the use of sterile clothing, footwear, and masks where required, or to prevent the escape of an animal.

§ 2.129 Confiscation and destruction of animals.

(a) If an animal being held by a dealer, exhibitor, intermediate handler, or by a carrier is found by an APHIS official to be suffering as a result of the failure of the dealer, exhibitor, intermediate handler, or carrier to comply with any provision of the regulations or the standards set forth in this subchapter, the APHIS official shall make a reasonable effort to notify the dealer, exhibitor, intermediate handler, or carrier of the condition of the animal(s) and request that the condition be corrected and that adequate care be given to alleviate the animal’s suffering or distress, or that the animal(s) be destroyed by euthanasia. In the event that the dealer, exhibitor, intermediate handler, or carrier refuses to comply with this request, the APHIS official may confiscate the animal(s) for care, treatment, or disposal as indicated in paragraph (b) of this section, if, in the opinion of the Administrator, the circumstances indicate the animal’s health is in danger.

(b) In the event that the APHIS official is unable to locate or notify the dealer, exhibitor, intermediate handler, or carrier as required in this section, the APHIS official shall contact a local police or other law officer to accompany him to the premises and shall provide for adequate care when necessary to alleviate the animal’s suffering. If in the opinion of the Administrator, the condition of the animal(s) cannot be corrected by this temporary care, the APHIS official shall confiscate the animals.

(c) Confiscated animals may be:

(1) Placed, by sale or donation, with other licensees or registrants that comply with the standards and regulations and can provide proper care; or

(2) Placed with persons or facilities that can offer a level of care equal to or exceeding the standards and regulations, as determined by APHIS, even if the persons or facilities are not licensed by or registered with APHIS; or
§ 2.130 Minimum age requirements.

No dog or cat shall be delivered by any person to any carrier or intermediate handler for transportation, in commerce, or shall be transported in commerce by any person, except to a registered research facility, unless such dog or cat is at least eight (8) weeks of age and has been weaned.

§ 2.131 Handling of animals.

(a) All licensees who maintain wild or exotic animals must demonstrate adequate experience and knowledge of the species they maintain.

(b)(1) Handling of all animals shall be done as expeditiously and carefully as possible in a manner that does not cause trauma, overheating, excessive cooling, behavioral stress, physical harm, or unnecessary discomfort.

(2)(i) Physical abuse shall not be used to train, work, or otherwise handle animals.

(ii) Deprivation of food or water shall not be used to train, work, or otherwise handle animals; Provided, however, that the short-term withholding of food or water from animals by exhibitors is allowed by these regulations as long as each of the animals affected receives its full dietary and nutrition requirements each day.

(c)(1) During public exhibition, any animal must be handled so there is minimal risk of harm to the animal and to the public, with sufficient distance and/or barriers between the animal and the general viewing public so as to assure the safety of animals and the public.

(2) Performing animals shall be allowed a rest period between performances at least equal to the time for one performance.

(3) Young or immature animals shall not be exposed to rough or excessive public handling or exhibited for periods of time which would be detrimental to their health or well-being.

(d) Animals shall be exhibited only for periods of time and under conditions consistent with their good health and well-being.

(2) A responsible, knowledgeable, and readily identifiable employee or attendant must be present at all times during periods of public contact.

(3) During public exhibition, dangerous animals such as lions, tigers, wolves, bears, or elephants must be under the direct control and supervision of a knowledgeable and experienced animal handler.

(4) If public feeding of animals is allowed, the food must be provided by the animal facility and shall be appropriate to the type of animal and its nutritional needs and diet.

(e) When climatic conditions present a threat to an animal’s health or well-being, appropriate measures must be taken to alleviate the impact of those conditions. An animal may never be subjected to any combination of temperature, humidity, and time that is detrimental to the animal’s health or well-being, taking into consideration such factors as the animal’s age, species, breed, overall health status, and acclimation.

§ 2.132 Procurement of dogs, cats, and other animals; dealers.

(a) A class “B” dealer may obtain live random source dogs and cats only from:

(1) Other dealers who are licensed under the Act and in accordance with the regulations in part 2;

(2) State, county, or city owned and operated animal pounds or shelters; and

(3) A legal entity organized and operated under the laws of the State in which it is located as an animal pound or shelter, such as a humane shelter or contract pound.

(b) No person shall obtain live dogs, cats, or other animals by use of false
pretenses, misrepresentation, or deception.

(c) Any dealer, exhibitor, research facility, carrier, or intermediate handler who also operates a private or contract animal pound or shelter shall comply with the following:

(1) The animal pound or shelter shall be located on premises that are physically separated from the licensed or registered facility. The animal housing facility of the pound or shelter shall not be adjacent to the licensed or registered facility.

(2) Accurate and complete records shall be separately maintained by the licensee or registrant and by the pound or shelter. The records shall be in accordance with §§2.75 and 2.76, unless the animals are lost or stray. If the animals are lost or stray, the pound or shelter records shall provide:

(i) An accurate description of the animal;
(ii) How, where, from whom, and when the dog or cat was obtained;
(iii) How long the dog or cat was held by the pound or shelter before being transferred to the dealer; and
(iv) The date the dog or cat was transferred to the dealer.

(3) Any dealer who obtains or acquires a live dog or cat from a private or contract pound or shelter, including a pound or shelter he or she operates, shall hold the dog or cat for a period of at least 10 full days, not including the day of acquisition and excluding time in transit. This holding period shall include at least one Saturday. The provisions of this paragraph apply to:

(a) Each of the entities listed in paragraphs (a)(1) through (a)(3) of this section that acquire any live dog or cat shall, before selling or providing the live dog or cat to a dealer, hold and care for the dog or cat for a period of not less than 5 full days after acquiring the animal, not including the date of acquisition and excluding time in transit. This holding period shall include at least one Saturday. The provisions of this paragraph apply to:

(1) Each pound or shelter owned and operated by a State, county, or city;
(2) Each private pound or shelter established for the purpose of caring for animals, such as a humane society, or other organization that is under contract with a State, county, or city, that operates as a pound or shelter, and that releases animals on a voluntary basis; and
(3) Each research facility licensed by USDA as a dealer.

(b) A dealer shall not sell, provide, or make available to any person a live random source dog or cat unless the dealer provides the recipient of the dog or cat with certification that contains the following information:

(1) The name, address, USDA license number, and signature of the dealer;
(2) The name, address, USDA license or registration number, if such number exists, and signature of the recipient of the dog or cat;
(3) A description of each dog or cat being sold, provided, or made available that shall include:

(i) The species and breed or type (for mixed breeds, estimate the two dominant breeds or types);
(ii) The sex;
(iii) The date of birth or, if unknown, then the approximate age;
(iv) The color and any distinctive markings; and...
(v) The Official USDA-approved identification number of the animal. However, if the certification is attached to a certificate provided by a prior dealer which contains the required description, then only the official identification numbers are required.

(4) The name and address of the person, pound, or shelter from which the dog or cat was acquired by the dealer, and an assurance that the person, pound, or shelter was notified that the cat or dog might be used for research or educational purposes;

(5) The date the dealer acquired the dog or cat from the person, pound, or shelter referred to in paragraph (b)(4) of this section; and

(6) If the dealer acquired the dog or cat from a pound or shelter, a signed statement by the pound or shelter that it met the requirements of paragraph (a) of this section. This statement must at least describe the animals by their official USDA identification numbers. It may be incorporated within the certification if the dealer makes the certification at the time that the animals are acquired from the pound or shelter or it may be made separately and attached to the certification later. If made separately, it must include the same information describing each animal as is required in the certification. A photocopy of the statement will be regarded as a duplicate original.

(c) The original certification required under paragraph (b) of this section shall accompany the shipment of a live dog or cat to be sold, provided, or otherwise made available by the dealer.

(d) A dealer who acquires a live dog or cat from another dealer must obtain from that dealer the certification required by paragraph (b) of this section and must attach that certification (including any previously attached certification) to the certification which he or she provides pursuant to paragraph (b) of this section (a photocopy of the original certification will be deemed a duplicate original if the dealer does not dispose of all of the dogs or cats in a single transaction).

(e) A dealer who completes, provides, or receives a certification required under paragraph (b) of this section shall keep, maintain, and make available for APHIS inspection a copy of the certification for at least 1 year following disposition.

(f) A research facility which acquires any live random source dog or cat from a dealer must obtain the certification required under paragraph (b) of this section and shall keep, maintain, and make available for APHIS inspection the original for at least 3 years following disposition.

(g) In instances where a research facility transfers ownership of a live random source dog or cat acquired from a dealer to another research facility, a copy of the certification required by paragraph (b) of this section must accompany the dog or cat transferred. The research facility to which the dog or cat is transferred shall keep, maintain, and make available for APHIS inspection the copy of the certification for at least 3 years following disposition.

§2.134 Contingency planning.

(a) Dealers, exhibitors, intermediate handlers, and carriers must develop, document, and follow an appropriate plan to provide for the humane handling, treatment, transportation, housing, and care of their animals in the event of an emergency or disaster (one which could reasonably be anticipated and expected to be detrimental to the good health and well-being of the animals in their possession). Such contingency plans must:

(1) Identify situations the licensee or registrant might experience that would trigger the need for the measures identified in a contingency plan to be put into action including, but not limited to, emergencies such as electrical outages, faulty HVAC systems, fires, mechanical breakdowns, and animal escapes, as well as natural disasters most likely to be experienced;

(2) Outline specific tasks required to be carried out in response to the identified emergencies or disasters including, but not limited to, detailed animal evacuation instructions or shelter-in-place instructions and provisions for providing backup sources of food and water as well as sanitation, ventilation, bedding, veterinary care, etc.;

(3) Identify a chain of command and who (by name or by position title) will
be responsible for fulfilling these tasks; and

(4) Address how response and recovery will be handled in terms of materials, resources, and training needed.

(b) For current licensees and registrants, the contingency plan must be in place by July 29, 2013. For new dealers, exhibitors, intermediate handlers, and carriers licensed or registered after this date, the contingency plan must be in place prior to conducting regulated activities. The plan must be reviewed by the dealer, exhibitor, intermediate handler, or carrier on at least an annual basis to ensure that it adequately addresses the criteria listed in paragraph (a) of this section. Each licensee and registrant must maintain documentation of their annual reviews, including documenting any amendments or changes made to their plan since the previous year’s review, such as changes made as a result of recently predicted, but historically unforeseen, circumstances (e.g., weather extremes). Contingency plans, as well as all annual review documentation and training records, must be made available to APHIS upon request. Traveling entities must carry a copy of their contingency plan with them at all times and make it available for APHIS inspection while in travel status. Dealers, exhibitors, intermediate handlers, and carriers maintaining or otherwise handling marine mammals in captivity must also comply with the requirements of §3.101(b) of this subchapter.

(c) Dealers, exhibitors, intermediate handlers, and carriers must provide training for their personnel regarding their roles and responsibilities as outlined in the plan. For current licensees and registrants, training of dealer, exhibitor, intermediate handler, and carrier personnel must be completed by September 27, 2013. For new dealers, exhibitors, intermediate handlers, or carriers licensed or registered after July 29, 2013, training of personnel must be completed within 60 days of the dealer, exhibitor, intermediate handler, or carrier putting their contingency plan in place. Employees hired 30 days or more before their contingency plan is put in place must also be trained by that date. For employees hired less than 30 days before that date or after that date, training must be conducted within 30 days of their start date. Any changes to the plan as a result of the annual review must be communicated to employees through training which must be conducted within 30 days of making the changes.

[77 FR 76823, Dec. 31, 2012]

EFFECTIVE DATE NOTE: At 78 FR 46255, July 31, 2013, §2.134 was amended by lifting the stay on the section published at 78 FR 46255, July 31, 2013; removing the date “July 29, 2013” from the first sentence of paragraph (b) and adding “July 5, 2022” in its place; removing the words “and training records” from the first sentence of paragraph (b); revising the last sentence of paragraph (b); adding paragraph (c); and adding an OMB citation at the end of the section, effective Jan. 3, 2022. For the convenience of the user, the added and revised text is set forth as follows:

§ 2.134  Contingency planning.

* * * * *

(b) * * * The APHIS Contingency Plan form may be used to keep and maintain the information required by § 2.38(b)(1) and (2).

(c) Dealers, exhibitors, intermediate handlers, and carriers must provide training for their personnel regarding their roles and responsibilities as outlined in the plan. For current licensees and registrants, training of dealer, exhibitor, intermediate handler, and carrier personnel must be completed within 60 days of the licensee and registrant putting their contingency plan in place; for new dealers, exhibitors, intermediate handlers, or carriers licensed or registered after July 5, 2022, training of personnel must be completed within 60 days of the dealer, exhibitor, intermediate handler, or carrier putting their contingency plan in place. This deadline applies to employees hired before and up to 30 days after the date the licensee or registrant puts its contingency plan in place. For employees hired more than 30 days after the date the licensee or registrant puts its contingency plan in place, training must be conducted within 30 days of their start date. Any substantive changes to the plan as a result of the annual review must be communicated to employees through training which must be conducted within 30 days of making the changes.

(Approved by the Office of Management and Budget under control number 0579–0479)
§ 2.150 Import permit.

(a) No person shall import a live dog from any part of the world into the States for purposes of resale unless the dog is accompanied by an import permit issued by APHIS and is imported into the States within 30 days after the proposed date of arrival stated in the import permit.

(b) An application for an import permit must be submitted to the Animal and Plant Health Inspection Service, Animal Care, 4700 River Road Unit 84, Riverdale, MD 20737–1234 or though Animal Care’s Web site (http://www.aphis.usda.gov/animal_welfare/). Application forms for import permits may be obtained from Animal Care at the address listed above.

(c) The completed application must include the following information:

1. The name and address of the person intending to export the dog(s) to the States;
2. The name and address of the person intending to import the dog(s) into the States;
3. The number of dogs to be imported and the breed, sex, age, color, markings, and other identifying information of each dog;
4. The purpose of the importation;
5. The port of embarkation and the mode of transportation;
6. The port of entry in the United States;
7. The proposed date of arrival in the States; and
8. The name and address of the person to whom the dog(s) will be delivered in the States and, if the dog(s) is or are imported for resale for research purposes, the USDA registration number of the research facility where the dog will be used for research, tests, or experiments.

(d) After receipt and review of the application by APHIS, an import permit indicating the applicable conditions for importation under this subpart may be issued for the importation of the dog(s) described in the application if such dog(s) appears to be eligible to be imported. Even though an import permit has been issued for the importation of a dog, the dog may only be imported if all applicable requirements of this subpart and any other applicable regulations of this subchapter and any other statute or regulation of any State or of the United States are met.

(Approved by the Office of Management and Budget under control number 0579–0379)


§ 2.151 Certifications.

(a) Required certificates. Except as provided in paragraph (b) of this section, no person shall import a live dog from any part of the world into the States for purposes of resale unless the following conditions are met:

1. Health certificate. Each dog is accompanied by an original health certificate issued in English by a licensed veterinarian with a valid license to practice veterinary medicine in the country of export that:
   i. Specifies the name and address of the person intending to import the dog into the States;
   ii. Identifies the dog on the basis of breed, sex, age, color, markings, and other identifying information;
   iii. States that the dog is at least 6 months of age;
   iv. States that the dog was vaccinated, not more than 12 months before the date of arrival at the U.S. port, for distemper, hepatitis, leptospirosis, parvovirus, and parainfluenza virus (DHLPP) at a frequency that provides continuous protection of the dog from those diseases and is in accordance with currently accepted practices as cited in veterinary medicine references guides;
   v. States that the dog is in good health (i.e., free of any infectious disease or physical abnormality which would endanger the dog or other animals or endanger public health, including, but not limited to, parasitic infections, lesions of the skin, nervous system disturbances, jaundice, or diarrhea); and
   vi. Bears the signature and the license number of the veterinarian issuing the certificate.

2. Rabies vaccination certificate. Each dog is accompanied by a valid rabies
vaccination certificate\(^6\) that was issued in English by a licensed veterinarian with a valid license to practice veterinary medicine in the country of export for the dog not less than 3 months of age at the time of vaccination that:

   (i) Specifies the name and address of the person intending to import the dog into the States;
   (ii) Identifies the dog on the basis of breed, sex, age, color, markings, and other identifying information;
   (iii) Specifies a date of rabies vaccination at least 30 days before the date of arrival of the dog at a U.S. port;
   (iv) Specifies a date of expiration of the vaccination which is after the date of arrival of the dog at a U.S. port; and
   (v) Bears the signature and the license number of the veterinarian issuing the certificate.

(b) Exceptions—(1) Research. The provisions of paragraphs (a)(1)(iii), (a)(1)(iv), (a)(1)(v), and/or (a)(2) of this section do not apply to any person who imports a live dog from any part of the world into the States for resale for use in research, tests, or experiments at a research facility, provided that: Such person submits satisfactory evidence to Animal Care at the time of his or her application for an import permit that the specific provision(s) would interfere with the dog's use in such research, tests, or experiments in accordance with a research proposal and the proposal has been approved by the research facility IACUC.

(2) Veterinary care. The provisions of paragraphs (a)(1)(iii) through (a)(1)(v) and (a)(2) of this section do not apply to any person who imports a live dog from any part of the world into the States for veterinary treatment by a licensed veterinarian and subsequent resale, provided that:

   (i) The original health certificate required in paragraph (a)(1) of this sec-

\(^6\) Alternatively, this requirement can be met by providing an exact copy of the rabies vaccination certificate if so required under the Public Health Service regulations in 42 CFR 71.51.
seized and the person intending to import the dog shall provide for the care (including appropriate veterinary care), forfeiture, and adoption of the dog, at his or her expense.

(79 FR 48659, Aug. 18, 2014, as amended at 85 FR 28798, May 13, 2020)

PART 3—STANDARDS

Subpart A—Specifications for the Humane Handling, Care, Treatment, and Transportation of Dogs and Cats

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ANIMAL HEALTH AND HUSBANDRY STANDARDS

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3.61 Primary enclosures used to transport live rabbits.
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