2013 AAFP Feline Vaccination Advisory Panel Report

Rationale: This Report was developed by the Feline Vaccination Advisory Panel of the American Association of Feline Practitioners (AAFP) to provide practical recommendations to help clinicians select appropriate vaccination schedules for their feline patients based on risk assessment. The recommendations rely on published data as much as possible, as well as consensus of a multidisciplinary panel of experts in immunology, infectious disease, internal medicine and clinical practice.

Introduction

The AAFP produced the first organization-driven vaccination guidelines in 1998. These were updated in 2000 and again in 2006.¹ Each version has offered a comprehensive review of the literature and has provided recommendations for vaccine protocols based on known science along with some extrapolation between studies and between species when feline studies were not available. This Report has used the same criteria.

The practicing veterinarian is in the best position to determine how to put these Guidelines into practice for an individual patient. The veterinarian should undertake a clinical risk/benefit assessment for each animal and discuss recommended vaccination schedules with the owner so that they can make an informed choice. The assessment should include discussion on the likelihood of exposure, the health and lifestyle of the animal, and the risks related to vaccination.

The Advisory Panel recognizes that situations differ in different countries, and that every country will have slightly different issues and priorities; thus these Guidelines will not necessarily be applicable to every country and the practitioner must interpret accordingly.

The three international panels that have produced feline vaccination guidelines (AAFP, World Small Animal Veterinary Association and European Advisory Board on Cat Diseases) recommend that an annual health examination be performed irrespective of whether vaccines are administered. While the optimal frequency of health examinations for cats is unknown, it is generally

CONTENTS

<table>
<thead>
<tr>
<th>CONTENTS</th>
<th>page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Introduction</td>
<td>785</td>
</tr>
<tr>
<td>Vaccination principles</td>
<td>786</td>
</tr>
<tr>
<td>General information on types of feline vaccines</td>
<td>787</td>
</tr>
<tr>
<td>Risk/benefit assessment</td>
<td>788</td>
</tr>
<tr>
<td>Vaccination recommendations for specific situations</td>
<td>789</td>
</tr>
<tr>
<td>– household pet cats</td>
<td>791</td>
</tr>
<tr>
<td>– shelter-housed cats</td>
<td>791</td>
</tr>
<tr>
<td>– cats in trap-neuter-return programs</td>
<td>793</td>
</tr>
<tr>
<td>– cats in breeding catteries</td>
<td>793</td>
</tr>
<tr>
<td>Vaccine adverse events</td>
<td>794</td>
</tr>
<tr>
<td>Pre-vaccination testing</td>
<td>795</td>
</tr>
<tr>
<td>Injectable vaccine administration site recom-</td>
<td>798</td>
</tr>
<tr>
<td>mendations</td>
<td></td>
</tr>
<tr>
<td>Legal considerations associated with vaccina-</td>
<td>799</td>
</tr>
<tr>
<td>tion</td>
<td></td>
</tr>
<tr>
<td>Abbreviations used in the Report and Disease Information Fact Sheets</td>
<td>799</td>
</tr>
<tr>
<td>Appendix 1: Frequently asked questions</td>
<td></td>
</tr>
<tr>
<td>– General FAQs</td>
<td>802</td>
</tr>
<tr>
<td>– Shelter FAQs</td>
<td>803</td>
</tr>
<tr>
<td>– Trap-neuter-return FAQs</td>
<td>804</td>
</tr>
<tr>
<td>– Adverse event FAQs</td>
<td>805</td>
</tr>
<tr>
<td>Appendix 2: Vaccinations for Your Cat</td>
<td></td>
</tr>
<tr>
<td>– Pet Owner Guide</td>
<td>807</td>
</tr>
</tbody>
</table>

See page 799 for list of Disease Information Fact Sheets and other resources available online as Supplementary Files.


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accepted that healthy adult cats should be examined at least once a year. In the past, annual veterinary visits were structured around vaccinations as the primary focus. With the increasing body of knowledge about duration of immunity (DOI) from vaccinations, their potential adverse effects, and the increased awareness of pet owners about these issues, it is clear that vaccination no longer justifies the need for annual visits.

Practitioners are encouraged to help cat owners understand the value of regular health care and that it ideally should be proactive rather than reactive. A useful approach is for health care to be tailored to the various feline life stages, which improves early recognition of potential health-related issues and can facilitate treatment.2

A Pet Owner Guide, discussing the risks and benefits of vaccination, is included as Appendix 2 (pages 807 and 808).

Vaccination principles

Vaccination plays an important role in the control of infectious diseases, both for an individual as well as for the cat population (ie, herd health). Some vaccine antigens are also used to lessen the potential for zoonotic spread of disease (eg, rabies). The benefits of routine and widespread vaccination are clear: the incidence of serious disease caused by highly pathogenic organisms, such as feline parvovirus (panleukopenia), can be reduced in populations in which widespread vaccination is practised. However, the level of protection conferred by a particular vaccine in an individual patient varies. The quality of vaccine-induced immunity in any patient is influenced by a complex interaction of factors unique to the individual patient, the patient’s environment, and the nature of the vaccine and pathogen. Precisely predicting either the outcome of vaccination or subsequent exposure to a pathogen is difficult (or impossible) and, therefore, vaccination should never be offered as a guarantee of protection.

The risk of infection and subsequent development of disease varies with a number of factors including the age and health of the cat, magnitude of exposure to the infectious agent, the pathogenicity of individual agents, the geographic prevalence of infection and the vaccination history of the cat. Some of the factors that negatively affect an individual animal’s ability to respond to vaccination include interference from maternally derived antibodies (MDA), congenital or acquired immunodeficiency, concurrent disease or infection, inadequate nutrition, immunosuppressive medications, chronic stress and an aging immune response. Additionally, some vaccinal agents (eg, FPV) will induce a much stronger protective immune response than others (eg,

Patient risk variables to take into consideration

- Age of cat
- Health of cat
- Magnitude of exposure to agent
- Agent pathogenicity
- Geographic prevalence
- History
- MDA interference
- Congenital or acquired immunodeficiency
- Immunosuppressive therapy
- Concurrent disease
- Nutritional status
- Chronic stress
- Aging immune response
feline herpesvirus [FHV-1]). As vaccine-afforded protection against both infection and disease is thus variable and not absolute, exposure to infected animals and infectious agents should be minimized, even after vaccination.

Kittens are generally more susceptible to infections than adult cats and are typically develop more severe disease (Figure 1). Thus, they represent a principal primary target population for vaccination. As part of a routine health care program, the vaccination needs of all cats, including adults, should be assessed at least once a year, in conjunction with a comprehensive physical examination and consultation, modifying vaccination recommendations as necessary on the basis of altered risk/benefit ratio.

Vaccination is a medical procedure, and the decision to vaccinate, even with core vaccines (see box above), should be based on a risk/benefit assessment for each cat and for each vaccine antigen. Vaccination may indeed be beneficial, but it is not innocuous, and the benefit of vaccinating an animal (eg, the induction of clinically meaningful immunity) must be balanced against the risk of adverse events, likelihood of exposure and severity of disease. Where practical, every effort should be made to ensure that cats are healthy prior to vaccination; however, concurrent illness should not necessarily preclude vaccination.

The overall objectives of vaccination are shown on the right.

General information on types of feline vaccines

Vaccines, including different products licensed to protect against the same pathogen, are not necessarily alike. Different vaccine technologies may directly influence efficacy, safety, DOI and route of administration of individual products. Awareness of fundamental differences is necessary.

The following terminology is used throughout these Guidelines to describe types of vaccines: inactivated (killed), modified-live (attenuated) and recombinant. The attributes of each vaccine type are summarized in Table 1.

Characteristics of vaccine types have been reviewed as recently as 2011.4 All veterinary vaccines, prior to licensing, are subjected to testing for efficacy, safety, potency and purity. Testing methods may vary among different manufacturers and licensing authorities. While all licensed vaccines need to meet minimum efficacy standards, the level of protection induced can vary depending on many factors, including the method used to manufacture the product. For further information on licensing, readers should refer to the 2006 Guidelines (see box on page 786) and to individual licensing authorities (United States Department of Agriculture [USDA]; Canadian Food Inspection Agency [CFIA]; Veterinary Medicines Directorate [VMD], Department for Environment, Food, and Rural Affairs [DEFRA], UK; European Medicines Agency [EMA], EU).

The principal differences between inactivated, modified-live and recombinant vaccines are discussed below.

- **Inactivated vaccines** Vaccinal pathogens can be completely inactivated (ie, killed) by various means, eliminating risk of replication post-inoculation or ‘reversion to virulence’. For these reasons, inactivated vaccines have historically been regarded as the safest vaccines. However, the inclusion of a variety of extraneous chemicals (stabilizers, preservatives), antibiotics, adjuvants and excipient proteins has been implicated as a cause of both acute and delayed adverse reactions in cats.4
- **Modified-live vaccines** For some agents, intact pathogens can be modified so that they
Examples of different types of feline vaccines and their attributes

<table>
<thead>
<tr>
<th>Examples</th>
<th>Inactivated (killed)</th>
<th>Modified-live (attenuated)</th>
<th>Recombinant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Replication following administration</td>
<td>Does not replicate (non-infectious)</td>
<td>May replicate locally and in sites beyond the inoculation site (infectious)</td>
<td></td>
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<tr>
<td>Initial vaccination, in the absence of maternal antibody</td>
<td>Two initial doses are required, 3–4 weeks apart. Protective immunity is expected within 7–10 days following the second dose. Rabies vaccine is the exception as only one initial dose is required; protective immunity is expected to develop by 28 days</td>
<td>Two initial doses are required, 3–4 weeks apart. Protective immunity is expected within 7–10 days following the second dose</td>
<td>rRabies, rFeLV</td>
</tr>
</tbody>
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| Route(s) of administration as stipulated by the manufacturer | Injectable: SC or IM* | Injectable: SC or IM* Mucosal: Intranasal (IN)*** | Injectable: SC |
| Adjuvanted | Yes – the majority | Not required | Some individual products contain an adjuvant |
| Therapeutic indications | Dermatophytosis (in some European countries) | None | None |
| Reversion to virulence | Not possible | Theoretically possible, but highly unlikely | Not possible |

NB Availability of different vaccines (type, antigen and route of administration) varies among countries

*The Advisory Panel recommends that when a vaccine is designed for either subcutaneous (SC) or intramuscular (IM) use, the SC route is used, both for patient comfort as well as for earlier detection of injection-site sarcomas

**Several products (two FHV-1, FCV; one FPV, FHV-1, FCV; *Bordetella*; FIP) are licensed for intranasal administration, though availability varies among countries

MDA = maternally derived antibodies, r = recombinant

retain the ability to replicate in the host and provoke an immune response, but not cause clinical disease. Altered pathogenicity effectively induces subclinical infection and can result in a more rapid onset of immunity for some vaccine antigens than with comparable inactivated vaccines.5,6 All bacterial and viral vaccines licensed for mucosal (intranasal) administration are modified-live, as are a number of injectable vaccines.

**Recombinant vaccines** Discrete genetic sequences can be isolated from a pathogenic virus or bacterium that encode immunogenic proteins. These sequences can either be recombined with the DNA of a live, non-pathogenic virus, which can then be administered as a vaccine (vectored vaccine), or they may be inserted in bacterial plasmids to enable in vitro production of antigens that can be harvested and purified for incorporation into a vaccine (ie, subunit vaccine). Examples of both types of vaccines are licensed for use in veterinary medicine.

Risk/benefit assessment

In assessing the risk for an individual cat, information about the cat, the environment and infectious agents to which the cat will be realistically exposed needs to be considered. Specifically, questions need to be asked that address the cat’s lifestyle as well as the lifestyle of any other cats in the same household. Queries should also be posed regarding other sources of exposure, such as excursions outside the home, boarding and travel.

**Patient**

Age is an important element in assessing an individual’s risk profile. Most infectious diseases are more prevalent in kittens, and kittens less than 6 months old are generally more susceptible to infection and disease than adult cats are. Kittens, therefore, represent a principal primary target population for vaccination. MDA provide important protection for the kitten, but may also interfere with, or neutralize, vaccines. As the level of MDA varies among individuals, the age at which a kitten may be able to respond to vaccination will also vary, and in some cases may be 16 weeks or older. While information is available on the variability of MDA as pertains to FHV-1, FCV and FPV, limited data is available for other antigens; thus the role of MDA in interference with vaccination against rabies, FeLV or other pathogens is unknown. Stopping a vaccination course too early (when MDA are still interfering) is thought to be the single most common cause of vaccination failure in kittens.
Patient’s environment
Population density and opportunity for exposure to other cats (eg, whether the cat is free-roaming or has access to the outdoors) are among the most critical issues affecting risk of exposure to an infectious agent. Cats and kittens living in multiple-cat households and environments (eg, boarding, breeding, foster or shelter facilities) are likely to have a substantially higher risk of infection than are cats living indoors in one- or two-cat households. Furthermore, the introduction of new cats into a household poses a potential risk – not only to the cat entering the household, but also to the whole group because of possible exposure to new infectious agents. The immunosuppressive effects of stress inherent in the change of social demographics may also result in recrudescence and an increased susceptibility to infection and disease. Conversely, cats that are naturally exposed to infectious agents after vaccination may have an opportunity for ‘natural boosting of immunity’ that may not be afforded to cats kept alone.

Indoor cats generally have a low risk of exposure to infectious agents, particularly where the agent in question is only transmitted by direct contact among cats. However, they may also be exposed to infection from other cats in the household (ie, subclinically infected or carrier cats), or by indirect transmission of pathogens brought in from outside on owners’ clothing, shoes, etc. In theory, strictly indoor cats may be more susceptible to developing panleukopenia because they do not receive boosting through the possibility of natural exposure. It is important to ask owners about other exposure that indoor cats may have, such as supervised visits out of doors (eg, on harness/leash, in the garden, etc), visiting other cats in an apartment building, balconies or roof gardens, visiting cats that belong to other family members, and staying in boarding facilities. Fostering shelter cats alters the risk for the resident cats, both through potential direct exposure to infectious agents as well as through stress-induced immunosuppression.

Veterinarians should reassess risk factors for exposure to infectious disease at least once a year, as changes in the health of the animal or its lifestyle may dictate modifications in vaccinations needed.

Trap–neuter–return (TNR) and other special situations are discussed on pages 791–794.

Geographic distribution of infectious agents may result in substantially different risks of exposure for cats living in different areas (eg, rabies). Questions regarding future travel should be included in determining the risk of exposure to specific infectious agents. Periodic housing in boarding facilities, shelters or breeding facilities or other multiple-cat households also places cats at increased risk of exposure to a variety of infectious agents, although the risk will vary substantially between different situations.

Infectious agent
Independent agent-associated variables, such as virulence, strain variation and mutation, challenge dose and stability in the environment, influence the outcome of infection. These are difficult to assess objectively.

Recommendations for vaccination of household pet cats
Developing universal guidelines for vaccination of household pet cats is complicated by the lack of a clear definition of what is, and what is not, a ‘pet cat’. What follows are reasonable recommendations, based on scientific evidence and expert advice, applicable to most cats presented to private practitioners. Differences in cat population density, introduction of new cats, and exposure risk are dynamic variables that the veterinarian must take into consideration when recommending any vaccine for any cat. It is advised that veterinarians reassess risk factors for exposure to infectious disease at each visit (at least once a year), as changes in factors such as the health of the animal or its lifestyle may dictate changes to vaccination needs.

Table 2 summarizes vaccination recommendations for household pet cats.
Additional considerations when vaccinating household pet cats

Because vaccination requirements and risk of exposure to infectious agents vary among household pet cats, individual vaccination protocols will vary. The following recommendations address some alternative situations and offer insights on vaccination of pet cats using non-core vaccines.

> **Vaccination of pet cats in indoor/outdoor households** Cats housed exclusively indoors generally do not require vaccination beyond the aforementioned vaccines (ie, FPV, FHV-1, FCV ± FeLV, rabies). However, in multiple-cat households where some cats are housed exclusively indoors, yet other cats are permitted outside unmonitored, the entire household may be at risk of exposure to additional agents. Veterinarians should consider recommending vaccination of the entire household for selected diseases (eg, FeLV ± rabies) if exposure risk is deemed significant.

Pet cats that spend most (or all) of their lives outdoors are at greater risk of exposure to most infectious diseases compared with predominantly indoor pet cats (Figure 2).

Offsetting this is the natural boosting of immunity they may receive if they are exposed to infectious agents. Among outdoor adult cats, exposure risk for rabies, FeLV and FIV is generally higher than for indoor cats. In addition to the conventional vaccines recommended in Table 2, FIV vaccination could be considered for outdoor cats. (See accompanying Disease Information Fact Sheet on FIV – details on page 799.)

> **Vaccination of pet cats entering boarding facilities** Although, in general, healthy adult cats only require boosters to FPV, FHV-1, FCV vaccines every 3 years, an additional booster 7–10 days prior to boarding may be warranted (and may be required by some catteries), particularly if the cat has not been vaccinated in the previous year. Boarding may be stressful for a cat and also, depending on the cattery and the situation at the time, may lead to exposure to infectious agents. However, disease control measures vary between facilities, with many providing individual housing, sneeze barriers and good hygiene, whereas others permit co-mingling of cats, which will clearly facilitate disease transmission.
transmission. In the event that kittens must enter a boarding facility, it is recommended that they should have received at least two doses of FPV, FHV-1, FCV vaccine, with the last dose 7–10 days prior to entry. In addition, it is strongly recommended that kittens be isolated from the general population of adult cats at all times while boarding.

- **Vaccination during pregnancy and lactation** Vaccination of pregnant or lactating cats is generally not recommended. Whenever possible, queens should be vaccinated before breeding. Vaccines are not evaluated for use in pregnant queens unless specifically stated on the label. However, the benefits of vaccination may outweigh the risks in endemic disease situations. Modified-live FPV vaccines should not be administered to pregnant queens as this has been associated with cerebellar hypoplasia in the kittens.29 (For a more comprehensive discussion, see ‘Recommendations for vaccination of cats housed in breeding catteries’, page 793.)

- **Overdue for vaccination** If the cat has been vaccinated previously and is overdue for revaccination (irrespective of the interval), generally a single vaccination is all that is required. If prior vaccination status is unknown, the cat should be treated as unvaccinated.

- **Bordetella bronchiseptica, Chlamydia felis, FIP and FIV vaccination** For information on the use of these vaccines, see accompanying Disease Information Fact Sheets (details on page 799).

- **Dermatophytosis vaccination** At the time of writing, a monovalent (*Microsporum canis*) and a multivalent (*Microsporum* and *Trichophyton* species) inactivated product are licensed for the prevention and treatment of dermatophytosis in cats in some countries in Europe. None are currently available in the USA or Canada. Limited evidence exists to support the safe use of these products as part of a comprehensive treatment protocol in cats with proven infection, but little evidence is available to support their use for prevention of infection.20,21

### Recommendations for vaccination of shelter-housed cats

Generally, shelter-housed cats (Figure 3) can be considered to be at especially high risk of exposure to infectious disease. Endemic disease, high rates of turnover, stress and sustained exposure are contributing factors. Vaccination in shelters should be limited to those diseases that are likely to be transmitted within the shelter itself. For diseases of concern in shelters (notably FPV and upper respiratory infections), vaccines may be indicated at an earlier age, and be administered at shorter intervals compared with schedules for pet cats. Rapid onset of protection is critical; therefore, administration of FPV, FHV-1, FCV vaccines should be considered for all cats at the time of (or ideally, before) intake.

Table 3 summarizes vaccination recommendations for shelter-housed cats.

### Additional considerations when vaccinating shelter-housed cats

- **Bordetella bronchiseptica and Chlamydia felis vaccination** The benefit of routine vaccination of shelter-housed cats against these disease agents is limited. The association between *B bronchiseptica* isolation and disease in shelters is inconsistent31–34 and *C felis* is not commonly isolated from shelter cats with upper respiratory infection.31 These vaccines should only be considered if the pathogens have been demonstrated as a current problem by laboratory diagnostics. *B bronchiseptica* vaccination should also be used where there is potential direct or indirect contact between cats and dogs on the same site, and the dogs have a recent or current history of infectious respiratory disease.

- **FIV and FIP vaccination** Vaccination of shelter-housed cats against these agents is not generally recommended.

- **Dermatophytosis vaccination** See comments in the household pet cats section.
For diseases of concern in shelters, vaccines may be indicated at an earlier age and administered at shorter intervals compared with schedules for pet cats.

**Table 3** Recommendations for vaccination of shelter-housed cats

<table>
<thead>
<tr>
<th>Vaccine</th>
<th>First inoculation</th>
<th>Subsequent inoculations</th>
<th>Comments</th>
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<tr>
<td>Panleukopenia + herpesvirus-1 + calicivirus (FPV, FHV-1, FCV)</td>
<td>Administer a single dose at intake or, where possible, at least 1 week prior to shelter entry. In kittens, administer the first dose as early as 4–6 weeks of age</td>
<td>Revaccinate every 2–3 weeks until 16–20 weeks of age&lt;sup&gt;8-10&lt;/sup&gt;</td>
<td>Recent studies show that ML SC vaccination may provide better protection in the face of MDA than inactivated vaccines do, and may protect against illness even when cats are placed in a contaminated environment soon after vaccination.&lt;sup&gt;19,23&lt;/sup&gt; ML injectable or IN vaccines containing FPV should not be given to kittens less than 4 weeks of age due to the risk of cerebellar hypoplasia&lt;sup&gt;19&lt;/sup&gt; or clinical panleukopenia (see Appendix 1 [Shelter FAQs] ‘Are there special considerations for vaccinating and housing very young kittens in shelters?’ page 803). For pregnant queens, risk of exposure versus risk of vaccination should be balanced (see Appendix 1 [Shelter FAQs] ‘Should pregnant queens in shelters be vaccinated?’ page 804). Inactivated multivalent calicivirus vaccines exist and may provide broader cross-protection against calicivirus infection than single strain vaccines.&lt;sup&gt;24,25&lt;/sup&gt; Calicivirus may be more prevalent in shelters housing cats long term in group settings.&lt;sup&gt;26,27&lt;/sup&gt; A multivalent vaccine may be preferable in this context. If FCV disease occurs in fully vaccinated cats housed in groups, changing to a product with a different vaccine strain(s) may be of benefit.&lt;sup&gt;28&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Revaccinate once, 2–3 weeks following administration of the initial vaccine</td>
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**Note:** Unless otherwise stipulated, all parenteral vaccines should be administered by the subcutaneous (SC) route.

<sup>1</sup>IN vaccination may provide protection against herpesvirus infection within 4–6 days, providing a hypothetical benefit in shelters.<sup>8,28</sup> However, results of IN vaccination for respiratory viruses in addition to parenteral vaccination in shelters are mixed, showing a modest reduction in upper respiratory disease in one shelter<sup>29</sup> but no difference in another.<sup>30</sup> Although simultaneous use of IN and parenteral vaccination is not generally tested by manufacturers and licensed for such use, there was no evidence in either study of reduced efficacy of the parenteral vaccine due to concurrent IN vaccine administration.<sup>29,30</sup> No information on safety was reported in these studies; however, there was no significant increase in respiratory signs within the first 7 days of administration in cats receiving the IN with the parenteral vaccine versus the parenteral vaccine alone, suggesting that vaccine-induced respiratory signs were not a significant concern. ML = modified-live, IN = intranasal, MDA = maternally derived antibodies.

Modified-live (use of inactivated vaccine is not generally recommended except where panleukopenia risk is low). Recommended for all cats.

Inactivated or recombinant

Rabies

Administer a single dose at the time of entry or release from the facility, depending on risk and length of stay

As for household pet cats (Table 2). NB Rabies vaccine should not be administered to kittens less than 12 weeks/3 months old

Necessary for all cats where legally mandated or in an endemic region. For shelters adopting out virtually all cats, or where the length of stay is commonly months or longer, rabies vaccine should be administered on intake. For shelters with shorter lengths of stay or where not all cats are adopted, rabies vaccination at the time of release is acceptable. If local regulations prohibit issuance of a rabies certificate for vaccines administered at the shelter, cats should receive a rabies vaccination from a local veterinarian within 4 weeks of adoption

Feline leukemia (FeLV)

Administer a single dose of vaccine at the time of intake if group-housed. If group (rather than individual) housing for kittens is used, vaccinate as early as 8 weeks of age

Revaccinate with a second dose 2–3 weeks later

Unlike group-housed cats, risk of FeLV transmission is very low for individually housed cats. FeLV vaccination is recommended for cats in long-term shelters or in group-housing of unrelated cats. Vaccination is not a substitute for testing and segregation of infected cats

Administer a single dose at intake or, where possible, at least 1 week prior to shelter entry. In kittens, administer the first dose as early as 4–6 weeks of age

Revaccinate every 2–3 weeks until 16–20 weeks of age<sup>8-10</sup> | Revaccinate once, 2–3 weeks following administration of the initial vaccine |

IN vaccination may result in onset of protection as early as 4–6 days post-inoculation.<sup>6,24</sup> Study results have been mixed<sup>6</sup> regarding reduction in risk for upper respiratory tract infection in shelters from IN vaccination.<sup>29,30</sup> When using IN vaccination, use only products licensed and approved for administration by this route. Transient, mild signs of upper respiratory infection may develop following administration of vaccine by the IN route.

Modified-live If IN vaccination is used for control of respiratory viruses, all shelter cats over 4–6 weeks of age should simultaneously receive a SC ML FPV vaccine (with or without respiratory viral antigens)
Recommendations for vaccination of cats in trap–neuter–return programs

Most community cats (Figure 4, ie, free-roaming unowned feral and stray cats) lack protective antibody titers against FPV, FHV-1 and rabies.\(^9,35\) In one study, the vast majority of feral cats vaccinated once at the time of TNR surgery developed protective antibody titers against FPV and FCV by the time they were re-trapped for testing 2–3 months later, regardless of whether inactivated or modified-live vaccines were used.\(^35\) In contrast, only inactivated vaccines resulted in a high rate of protective antibodies against FHV-1.\(^35\)

In the same study, nearly all cats developed high antibody titers against rabies after a single dose of inactivated rabies vaccine.\(^35\) Vaccine licensing studies have demonstrated 3–4 year DOI following a single vaccine administered to laboratory kittens. This suggests that, while the first rabies vaccine may only be recognized by regulatory agencies as valid for a single year, it is likely that vaccinated cats are protected for much longer.

It is the recommendation of the Advisory Panel that cats in TNR programs receive FPV, FHV-1, FCV and rabies vaccines at the time of surgery.

Recommendations for vaccination of cats housed in breeding catteries

Breeding catteries are variable in size, population and the nature of available facilities. The cat population may number less than 10 individuals or more than 50. Cats of various ages and life stages are typically present and many catteries continue to house retired breeding individuals that have been neutered. Some also contain household pets that may or may not have access to outdoors. The facilities may be sophisticated enough to allow for segregation of subpopulations or all individuals may be housed together. Generally, the medical and vaccination history of the residents is well known, but some diseases, such as upper respiratory tract disease, may be endemic.

Vaccination programs should be limited to those diseases that are relevant to the cattery and should be determined by analysis of risk factors. When assessing the level of disease risk in catteries, factors to consider include:

- Rate of population turnover.
- Population size and density.
- Number of litters/year (Figure 5).
- Presence of endemic disease.

Transmission of infectious diseases is facilitated by group living, young kittens mixing with older kittens and adults, contact during mating, introduction of new cats, and movement of cats into and out of the cattery (eg, queens going to other catteries for breeding, return of previously sold cats, travel for cat shows or other exhibitions). Catteries assessed as low risk would be considered similar to pet homes (Table 2), whereas catteries assessed as high risk would be considered similar to shelters (Table 3), pet stores, etc. In high-risk environments, vaccines may be used at an earlier age than in pet cats, particularly for control of endemic upper respiratory tract disease.

In general, vaccination may be started at an earlier age than in the pet cat population and revaccination intervals may be shortened. Breeders should be encouraged to work with a veterinarian to develop a comprehensive wellness program that includes appropriate vaccinations for their specific situation. Vaccination records should be kept for each individual in the cattery that include all relevant information (eg, antigen, brand, date, vaccination site, adverse events, etc). Management and husbandry have an important impact on the health of individual cats in catteries. Relevant references and resources should be consulted.\(^36,37\)
Additional considerations when vaccinating cats in breeding catteries

- **FeLV and FIV vaccination** Vaccination of cats in breeding catteries against these agents is not generally recommended. Vaccinate if necessary by analyzing risk, as for household pet cats and kittens (Table 2). The retrovirus status of all cats should be known: vaccination is not a substitute for testing and isolation. Vaccination may be unnecessary if a good testing program is in place and no cats have access to the outdoors. If queens are routinely sent to another cattery for breeding, vaccination of breeding queens may be considered.

- **Rabies vaccination** Cats in breeding catteries in the USA must be vaccinated against rabies according to state regulations. Elsewhere, vaccination against rabies is not generally recommended. Vaccinate if necessary by analyzing risk, as for household pet cats and kittens (Table 2).

- **Bordetella bronchiseptica and Chlamyphila felis vaccination** The benefit of routine vaccination of cats in breeding catteries against these disease agents is limited. These vaccines should only be considered if the pathogens have been demonstrated as a current problem by laboratory diagnostics. When used, the primary series should be administered according to the manufacturer’s instructions, with annual revaccination if the problem remains endemic. In some countries, the manufacturer states that Bordetella vaccination is considered safe for pregnant queens. However, in other countries, datasheets advise that the vaccine should not be used in pregnant or lactating queens or in kittens less than 1 month of age.

- **FIP vaccination** Vaccination of cats in breeding catteries against FIP is generally not recommended as there is insufficient evidence that the vaccine induces clinically relevant protection. (See accompanying Disease Information Fact Sheet – details on page 799.)

- **Dermatophytosis vaccination** See earlier comments in the household pet cats section (page 791).

### Vaccine adverse events

Although the administration of biological products can never be entirely free of risk, in general currently available feline vaccines have an excellent safety record. It is important to report any known or suspected negative events associated with vaccination, recognizing that a temporal relationship between an event and vaccine administration does not necessarily imply causality.
Pre-vaccination testing

Because antibody titers may not reliably correlate with, or predict, the degree of protection or susceptibility for an individual cat, the Advisory Panel recommends employing defined revaccination intervals to assure protection.

FeLV and FIV
The retrovirus status of all cats should be known and this is important if administration of FeLV or FIV vaccines is being considered. There is no recognized clinical benefit in administering vaccine against the retrovirus a cat is infected with, nor are there any known harmful effects. However, when the true retrovirus status of a vaccinated and infected cat eventually becomes known, not having known the cat’s status before vaccination could result in questions about failure to recommend testing before vaccination and vaccine efficacy.

Use of serology
The use of serology (serum antibody titers) to assess protective immunity has been reviewed. It is important to be aware that a variety of methods (immunofluorescence assay, ELISA, virus neutralization, haemagglutination inhibition, etc) are utilized to determine titers. The methodology used may not be reported with the test results. Titer results in individual cats determined at the same point in time, therefore, may vary depending on the methodology used. When electing to submit serum for antibody titers, it needs to be appreciated that a ‘positive’ antibody titer result obtained on one day is not necessarily predictive of a ‘positive’ titer at any point in the future. In general, cats having a ‘positive’ antibody titer against FPV are immune. In fact, the protective immunity that develops following FPV vaccination is expected to be sustained for several years. By contrast, serum antibody titers for FHV-1 and FCV may not necessarily correlate well with protective immunity and should not be used to predict protection in the future. Antibody titers to FeLV and FIV do not correlate with immunity and should not be used to determine the need for vaccination. Although feline rabies titers can be determined (by a certificated laboratory) in individual animals, a rabies titer is only an indication of serological response to vaccination. Rabies titers are not recognized as an index of immunity.

In addition, the absence of significant levels of antibody (a ‘negative’ titer) is not necessarily an indication of susceptibility. For example, a previously vaccinated cat may, over time, lose antibody. Immunologic ‘memory’, however, may prevail. In the event this individual is exposed to a virulent virus, a rapid anamnestic and protective response could result. In some diseases (eg, FHV-1), cell-mediated immunity is important and a cat may be immune even though no antibodies are detectable.

Because antibody titers may not reliably correlate with, or predict, the degree of protection or susceptibility for an individual cat, the Advisory Panel recommends employing defined revaccination intervals rather than measuring antibody titers to assure protection.

Prevalence and type of adverse reactions
Although post-vaccinal adverse events in cats are considered rare, the true prevalence is likely to be underestimated due to under-reporting by both veterinarians and owners. In the most substantial survey to date, adverse reactions were reported for all cats presented to Banfield Pet Hospitals in the United States between 2002 and 2005. During this period, more than 1.25 million doses of various vaccines were administered to nearly 0.5 million cats. Adverse reactions within 30 days of vaccination were reported at a rate of 51.6/10,000 cats vaccinated (0.52%), with 92% of these reactions occurring within the first 3 days. Clinical signs described for 1699 of 2560 cats with vaccination-associated adverse events included lethargy (± pyrexia) in 54%, local pain or swelling at the vaccine site (25%), vomiting (10%), facial or periorbital edema (6%) and generalized pruritus (2%). Death was reported in four cats, and in at least two of these it was attributed to anaphylaxis.

Although the vaccines used were predominantly from one manufacturer, no vaccine type was found to be significantly more likely to cause local reactions. Administration of multivalent FPV, FHV-1, FCV and Chlamydophila vaccines was significantly more likely to be associated with lethargy (± pyrexia) than administration of vaccines without the Chlamydophila component. The risk of an adverse reaction was greatest in cats around 1 year of age and/or increased as the number of vaccines administered concurrently increased.

In another extensive study specifically investigating local post-vaccine reactions, a prevalence of 0.23% was reported. Previous large
studies have suggested adverse vaccine reaction rates of around 1–3%,46–48 but some variation in prevalence can be expected with the use of different products, administration of multiple vaccines at the same appointment, and surveillance methods.

Hypersensitivity reactions
Anaphylaxis and allergic reactions
Anaphylaxis is perhaps the best characterized immune-mediated hypersensitivity (type I) reaction to vaccination, but it is rare (approximately 1–5/10,000 vaccines).4,6 In cats it may manifest as vomiting, diarrhea, respiratory distress, facial or generalized pruritus, facial swelling and collapse.1,43,49

A careful risk assessment is needed when considering the revaccination of cats with a history of anaphylaxis. In cats that have experienced an allergic reaction with true anaphylaxis, revaccination should usually be avoided. Vaccine excipients (inactive ingredients) are thought to cause most type I hypersensitivity reactions.1 Hence, where revaccination is considered necessary, using a different vaccine formulation and premedicating with an antihistamine and glucocorticoids 20–30 mins prior to vaccine administration is recommended, followed by close observation of the patient for several hours.1,4

Depending on geographic location, the requirement to vaccinate cats for rabies may take precedence over medical considerations. Veterinarians are urged to contact the appropriate authorities to determine what the local status is when concerns arise and whether the individual may be excused from vaccination. (See also 2006 Guidelines, Appendix 1: Certificate of Exemption from Rabies Vaccination – details on page 786.)

Other reactions
While other forms of hypersensitivity reactions (types II, III and IV) almost certainly occur in cats after vaccination, these are rarely documented. Some forms of local reaction probably reflect type IV reactions. Polyarthritis is occasionally seen after FCV vacci-

SPECIAL ARTICLE / 2013 AAFP feline vaccination guidelines

Figure 6 While vaccines are not uniquely implicated in sarcoma formation, there are certain actions that can be taken to reduce the risk of FISS, as summarized in Table 5. Courtesy of Albert Lloret

nation. Rarely it may represent a form of type III reaction, but it is mainly due to co-infection with field virus or vaccine virus itself.50,51 (See Appendix 1 [General FAQs] ‘What is the cause of lameness occasionally seen after FCV vaccination?’ page 802.)

Update on feline injection-site sarcomas (FISS)
Vaccine-associated sarcoma was first recognized as an issue in cats in the early 1990s. While initial studies suggested a risk of sarcoma development in around 2/10,000 doses of vaccine administered,52 which increased to 13–36/10,000 doses in other studies,33–35 current estimates based on larger epidemiologic studies (published between 2002 and 200745,56) suggest that the risk of sarcoma development following vaccination is actually very low (probably well below 1/10,000 doses of vaccine).4,45

Although initial reports linked development of sarcomas at vaccination sites with the use of inactivated rabies57 or FeLV vaccines,52 and aluminum-based adjuvants, more recent studies found no relationship between vaccine type, brand or use of inactivated versus modified-live vaccines and the risk of subsequent sarcoma formation.56,58,59 The impact of using the canarypox-vectored rabies vaccine is still unclear. One retrospective study of histopathology samples showed no reduction in the prevalence of FISS after the introduction of this vaccine; however, the types of vaccine used were not reported.8 In a recently published case control study it was suggested that there may be a lower risk of inducing sarcomas with this vaccine than with other rabies vaccines.59 Many of these studies have also clearly shown that injections other than vaccines also have the ability to induce sarcoma formation.

No studies have been published that define objective methods for reducing the risk of FISS in individual cats presented for routine vaccination. Based on our current understanding of this problem, it is likely that vaccines are not uniquely implicated in the development of injection site sarcomas in cats.56,60 FISS risk following vaccination likely results from a complex interaction of multiple extrinsic (eg, frequency and num-
by the use of modified-live vaccines. There are er that, on balance, risk might be mitigated between modified-live and inactivated vac-

mation as outlined above does not clearly injectable products. Although current infor-

risk of inducing FISS, as do at least some other mations demonstrate that all vaccines carry some ries demonstrate that all vaccines carry some

take into consideration. Recent stud-

Advisory Panel recommends that the follow-

prevent or cure FISS. (see Table 6 and Appendix 1 [General FAQs],

occur in some cats

inflammation in the pathogenesis of FISS is not clear.

The role of adjuvants (including those containing aluminum) and local

inflammation in the pathogenesis of FISS is not clear. Both

adjuvanted and non-adjuvanted vaccines induce local inflammation,

although the magnitude and type of inflammation varies among vaccines,

adjuvants and individual cats. However, some authors recommend

considering non-adjuvanted vaccines to try to reduce local inflammation

ML = modified-live, DOI = duration of immunity

Vaccination of preg-
nant queens

To avoid the risk of fetal/ neonatal infection with ML FPV

ML FPV may replicate in cerebellar tissue of fetal and/or neonatal kittens, leading to clinical signs associated with cerebellar hypoplasia.

Vaccination of cats known to be retrovirus positive

To avoid unlikely, but potential consequences of exposing an immune-
suppressed cat to ML, replicating vaccine virus

The risks associated with administering ML vaccine virus to an immune-suppressed (retrovirus-positive) cat are unknown

High-density housing environments where upper respiratory infections are not known to be present

To avoid the risk of accidental or inadvertent oral/nasal exposure to ML FPV, FHV-1, and FCV vaccines

ML FPV, FHV-1 and FCV virus, if inadvertently aerosolized or inoculated by the oral or nasal routes, may cause upper respiratory signs associated with vaccine virus replication on mucosal surfaces

Rabies vaccine: when 3-year DOI is indicated or required

To provide licensed 3-year DOI

The only rabies vaccines currently licensed to provide 3-year DOI are inactivated vaccines

ML = modified-live, DOI = duration of immunity
Post-vaccination monitoring

The Advisory Panel recommends that clinicians and their staff instruct clients to monitor the vaccine site for swelling or lumps in order to detect potential sarcomas while they may still be removed successfully. Biopsy of any mass present is warranted if it (a) remains present 3 months after vaccination; (b) is larger than 7 mm; (c) persists longer than 3 months; (d) increases in size or becomes firm; (e) becomes irregular in shape; (f) is painful; (g) is associated with redness or swelling around the injection site; (h) has a warm or hot skin fold; or (i) is seen to be growing.
2 cm in diameter; or (c) is increasing in size 1 month after vaccination (the ‘3-2-1 rule’ – see Table 5). It is recommended that multiple needle biopsies or an incisional wedge biopsy are obtained to reduce the risk of harvesting non-representative biopsy material and to minimize the risk of tracking tumor cells outside of the future surgical field.

**Legal considerations associated with vaccination**

Veterinarians in most countries are permitted to use professional judgment in the selection and use of licensed vaccines. Reference to these Guidelines, therefore, is appropriate when developing vaccination protocols for individual patients even though the guidance may vary from the manufacturer’s label recommendations or data sheet (eg, annual revaccination vs triennial revaccination for core vaccines).

### Rabies vaccination of cats

Where rabies vaccination of cats is required, veterinarians may not have discretion to vary from the manufacturer’s recommendations or from requirements set forth by regulatory agencies. Rabies vaccination requirements vary from country to country and can vary significantly within individual countries. In locations where feline rabies vaccination is required by law, veterinarians are obligated to be familiar with and follow legal requirements when administering rabies vaccines. Rabies vaccination recommendations contained in these Guidelines do not constitute vaccination requirements.

### Medical record documentation of vaccination

At the time of vaccine administration, the following information should be recorded in the patient’s permanent medical record:

- Vaccine(s) recommended for this patient.
- Date of vaccine administration.
- Identity (name, initials or code) of the person administering the vaccine(s).
- Vaccine name, lot or serial number, expiration date, and manufacturer of vaccine(s) actually administered.
- Site and route of vaccine administration.
- Concurrent medications/therapy.
- Recommendations for future vaccinations.

Adverse events should be recorded in a manner that will clearly alert all staff members during future visits. Risks and benefits of vaccination should be discussed with the owner so that they can make an informed choice. Consent should be documented in the medical record to demonstrate that relevant information was provided to the client and that the client authorized the procedure.
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Conflict of interest

RBF consults with IDEXX Laboratories (Westbrook, ME), Merial Ltd (Duluth, GA) and Elanco (Greenfield, IN). RMG has worked with various pharmaceutical, vaccine and diagnostic test companies over the years to complete research studies, student and house officer training, to present at veterinary conferences, and to serve on expert advisory panels.

References

32. Veir JK, Ruch-Gallie R, Spindel ME and Lappin MR. Prevalence of select-


Appendix 1: Frequently asked questions

General FAQs

- **What is the optimal interval between vaccines?**
The minimum vaccination interval during the primary series is 2 weeks, and the maximum recommended interval is 4 weeks. Kittens presented 6 weeks or longer following administration of the previous dose of vaccine should receive at least two doses of vaccine, 3–4 weeks apart. Although feline-specific data do not exist, extrapolation from mice and humans suggests that a 3-week interval is optimal for induction of memory T cells after administration of a modified-live virus.¹ ²

- **When should kittens be vaccinated?**
The primary vaccination series in kittens is scheduled between 6 and 16 weeks of age; vaccines should be administered at an interval of 3–4 weeks. Under high-risk circumstances (eg, in shelters and catteries with endemic upper respiratory tract disease or panleukopenia risk), vaccination may begin as early as 4–6 weeks of age and be repeated every 2–3 weeks until 16 weeks of age.

- **How often should senior/geriatric cats be vaccinated?**
  - **How should I vaccinate cats with stable chronic disease?**
  Whether older cats respond to vaccination in the same manner as younger animals do is inadequately studied.³ In the absence of data, the Advisory Panel recommends that healthy older cats and those with chronic but stable disease conditions receive vaccines in the same manner as younger adults. Less frequent vaccination is not advised due to inherent immunosenescence. Further, more frequent immunization is not warranted in aged patients with a lifelong history of immunization as data from other species suggests the memory response remains intact throughout life and protective immunity can be effectively maintained between boosts.⁴ ⁵

- **Should cats be vaccinated against rabies in areas where it is not required by law?**
  With the exception of Hawaii, cats in all of the states of the United States and cats in all countries or counties with endemic rabies of any species should be vaccinated against rabies, even if not required in that jurisdiction.

- **Should I vaccinate immunocompromised cats?**
  Patients with impaired immune responses, either due to infection with FIV/FeLV or the use of immunosuppressive therapies, are at increased risk of infection and may be candidates for vaccination. Although there is limited feline-specific data, inactivated vaccines are generally regarded as safer in patients with underlying immunosuppression.⁷ Because immune responses are hampered in immunocompromised patients, vaccination should ideally be updated before immunosuppressive therapies are started. Retrovirus-infected cats should not be vaccinated against the retrovirus they are infected with.

- **Should I vaccinate a kitten or cat with mild illness such as chronic upper respiratory tract infection or diarrhea?**
  In kittens and cats with mild illness, vaccination does not need to be delayed if the patient is eating and is not febrile. Should the illness be severe enough to result in fever or significant inappetence, vaccination should be delayed until these clinical signs have resolved. If the interval between vaccinations is delayed to greater than 6 weeks in primary immunizations, the series should be re-initiated. (See Shelter/Trap–Neuter–Return FAQs, pages 803–804, for alternate recommendations for those settings.)

- **Does it matter if the brand of vaccine used for revaccination is different from the brand administered previously?**
  While there are no studies comparing all vaccines for a particular antigen (or group of antigens), based on the available information the Advisory Panel believes that, subsequent to the initial series, booster vaccinations do not have to be of the same brand or vaccine type.

- **Can a parenteral FPV, FHV-1, FCV vaccine that is meant to be administered via the subcutaneous route be administered intranasally?**
  No, nor should it be administered by any other mucosal route. It will not stimulate an appropriate immune response and may cause clinical disease.

- **What is the cause of lameness occasionally seen after FCV vaccination?**
  In most cases, the lameness and pyrexia are due to coincidental infection with field feline caliciviruses, though a small proportion may be due to vaccine virus itself.⁶ ⁷ There is also some evidence of immune complex formation in the joints of cats infected with some strains of feline calicivirus,¹⁰ ¹¹ but this is likely to be an uncommon cause of lameness after vaccination.

- **How useful are the dermatophyte vaccines available in some European countries?**
  Dermatophyte vaccines have been marketed for years, but scientific studies to prove their efficacy have been unsatisfactory. Although some fungal vaccines may improve clinical signs compared with placebo, there does not seem to be a difference in infection rate between vaccinated and unvaccinated cats, and reliable protection has not been documented.¹² ¹³

- **Is it safe to mix and administer vaccine from one manufacturer with vaccine from another manufacturer?**
  No. Only vaccines from the same manufacturer, and only when stipulated on the product label/data sheet, may be mixed in the same syringe and administered simultaneously. Mixing vaccines from different manufacturers in the same syringe at the same time carries significant risk that one product (due to such factors as pH, osmolality, etc) may rapidly and completely inactivate the immunogenic antigen in the other product, or even in both products.

  For this reason, it is recommended that when administering...
vaccine from two different manufacturers to the same patient at the same appointment, separate inoculation sites should be selected (eg, left side vs right side).

- **How long is a vaccine stable for after being reconstituted?**
  Vaccines should always be stored and handled according to the manufacturer’s instructions. After reconstitution, a vaccine should ideally be used immediately, but certainly within 1 hour.

- **Should the skin be disinfected before administering a vaccine?**
  No, the disinfectant could potentially inactivate modified-live vaccine antigens. Cleaning and reusing of syringes is discouraged for this reason, as well as the risk of contamination.

- **Under what circumstances might an adjuvanted vaccine be preferable to vaccines that do not contain added adjuvants?**
  There are a number of factors that may play a role in the causation of FISS. Although initial reports linked development of sarcomas at vaccination sites with the use of inactivated vaccines and aluminum-based adjuvants, recent studies show that all vaccines carry some risk of inducing FISS, as do at least some other injectable products.
  
  Although current information does not clearly show differences in risk of FISS development between modified-live and inactivated vaccines, some Advisory Panel members feel that risk might be mitigated by the use of modified-live vaccines. However, overall the Advisory Panel concluded that, at the current time, there is insufficient information to make definitive recommendations to use particular vaccine types to reduce the risk of FISS.

  There are indications for using an inactivated vaccine. The risks and benefits should be discussed fully with the client. Some examples of situations where this might be considered are given in Table 6 of the Guidelines (page 797).

### Shelter FAQs

- **Are there special considerations for vaccinating and housing very young kittens in shelters?**
  It is preferable that kittens younger than 8 weeks of age be kept in foster care in clean homes. Interference from MDA and lack of immune competence has a negative impact on the ability of vaccines to induce a protective immune response, and kittens placed in shelters are at high risk of disease. If kittens younger than 8 weeks of age must be kept in shelters, they should be kept in areas isolated from the general population.
  
  When challenge dose is high and exposure is unavoidable, FHV-1 and FCV intranasal or injectable vaccines may be administered to kittens younger than 4–6 weeks of age. Some facilities administer one or two drops of intranasal vaccine rather than the entire dose to each kitten.
  
  However, unless specifically stated on the label, manufacturers have not evaluated the safety and efficacy of these vaccines when used in this manner, nor have such practices been independently evaluated. As such, use of a partial dose is not recommended. As in older cats, signs of upper respiratory disease may be caused by the vaccine. Nonetheless, in environments with endemic upper respiratory disease where the risk of serious disease is high, the benefits of vaccinating in this manner may outweigh the risks.

- **Injectable or intranasal modified-live FPV vaccine may cause cerebellar hypoplasia if administered to kittens prior to 4 weeks of age.**
  Kittens in high-risk shelters should, therefore, be vaccinated with a modified-live, injectable FPV vaccine no earlier than 4–6 weeks of age. Vaccination should be repeated every 2–3 weeks until 16–20 weeks of age.
  
  The shorter end of the inter-vaccination interval and earlier age of first vaccination is appropriate when risk of infectious disease is high, such as during an outbreak or in a known contaminated environment.

- **Are there any special vaccine considerations for cats living long term (months or years) in shelters or sanctuaries?**
  Cats entering a long-term care facility (or any cat for which a long-term shelter stay is anticipated) should be vaccinated against rabies at the time of admission, unless in a rabies-free region. FeLV vaccination is recommended for cats that will be group housed, with the two vaccine primary series ideally completed prior to placement in group housing. Other non-core vaccines (ie, other than FPV, FHV-1, FCV) should be considered as for household pet cats (see Table 2, page 790), depending on risk profile. In the event that a cat resides in the facility for a sufficiently long period to justify booster vaccination, it is recommended that the same schedule for revaccination be followed as is recommended for pet cats. There is no indication for more frequent vaccination in a long-term shelter facility with a stable population. Cats in long-term care facilities are at increased risk of calicivirus infections. Use of dual- or multi-strain calicivirus vaccines in such facilities may be indicated.
  
  If FCV disease occurs in fully vaccinated cats, changing to a product with different vaccine strain(s) may be beneficial.

- **Are vaccine recommendations different for shelter cats that are ill or injured?**
  The great majority of shelter kittens and cats should be vaccinated regardless of physical condition. If the cat’s immune system is so weakened that a modified-live vaccine will induce disease, exposure to the wide variety of infectious pathogens present in most shelters will very likely be fatal. In general, if a cat cannot be safely vaccinated, it cannot safely remain in an animal shelter except in strict isolation. Cats that were injured or ill at the time of initial vaccination should be revaccinated when healthy (no sooner than 2 weeks after recovery).
Should pregnant queens in shelters be vaccinated?
In general, vaccines are not licensed for use in pregnant queens unless specifically stated on the product label. The use of modified-live vaccines in naive queens (ie, those that have never been naturally exposed or vaccinated) during pregnancy is particularly not recommended due to potential adverse effects of FPV on developing fetuses. Nonetheless, the likelihood of exposure to FPV is very high in many shelters, and infection may result in the death of the mother as well as her offspring. Therefore, the risks posed by modified-live vaccination must be weighed against the risks of not vaccinating (ie, maternal, fetal or neonatal infection and death). When pregnant queens are being placed into shelters where FPV exposure is likely, the Advisory Panel believes that the overall benefits of modified-live FPV vaccination outweigh the risks and are preferable in the shelter environment due to the more rapid onset of protection. (See also Tables 3 and 6, on pages 792 and 797, respectively.)

Vaccination against FHV-1 and FCV during pregnancy may actually be beneficial for both mother and offspring, although vaccines are not actually licensed for such use. Vaccines administered early in pregnancy will not only protect the mother, but may provide the offspring with higher levels of MDA to protect them during the first few weeks of life. Reduced morbidity and mortality from feline upper respiratory infection was seen in kittens born to queens vaccinated with an inactivated vaccine against FHV-1 and FCV during early pregnancy, compared with offspring of queens not vaccinated during pregnancy. There was no increase in abortions or stillbirths associated with this practice.

Should previously vaccinated cats receive booster vaccines at the time of shelter intake?
In theory there is no reason to administer vaccines at the time of shelter admission if clear documentation of previous vaccination within the timeframe recommended by these Guidelines can be provided. An exception may be for the respiratory viruses (FHV-1 and FCV). While protection generally persists for 3 years, the degree of protection may wane over time. It may be helpful to revaccinate cats for FHV-1 and FCV if they have not received a vaccine in the previous year. If there is any question about the vaccine history, re-administering vaccines is preferable to reliance on uncertain records.

Should cats be vaccinated even if most of them are likely to be euthanased a few days after intake?
Yes, the primary reason to vaccinate cats in high-euthanasia shelters is to prevent the development of endemic FPV transmission. Protection against FPV develops in a high proportion of cats within the first few days of vaccination (if there is no MDA interference). Vaccinating all cats at intake is associated with a decreased risk of widespread FPV outbreaks.

Should cats be vaccinated on intake even if they are stray and, therefore, not the property of the shelter?
Yes, stray cats should be vaccinated on intake. Many community cats have no antibodies to protect against serious illness; thus, the benefit of vaccination for population and individual health generally greatly outweighs the risk of vaccination. To decrease risks associated with vaccination, antigens and vaccines should be limited to those that present a threat in a given shelter environment, and the clinical signs and procedure for responding to an adverse vaccine reaction should be prominently posted in all areas where vaccines are administered.

Does performing spay/neuter surgery at the time of vaccination diminish immune responses?
Kittens sterilized a week before, a week after, or at the time of vaccination had similar antibody titers to kittens that were vaccinated without surgery. Anesthesia and surgery do not appear to impede serological responses to vaccination.

Trap-neuter-return FAQs

Does the susceptibility of feral cats to common infectious diseases justify investment in vaccination during trap-neuter-return programs?
The majority of feral cats admitted to one TNR program lacked protective antibody titers against FPV, FHV-1 and rabies. The fact that some cats were seropositive suggests that this population of cats was exposed to infectious diseases and would benefit from immunization.

Are feral cats that receive only a single vaccination during the stressful experience of trapping and neutering effectively immunized?
The vast majority of feral cats vaccinated at the time of surgery developed protective antibody titers against FPV-1 and FCV by the time they were re-trapped for testing 2–3 months later, regardless of whether inactivated or modified-live vaccines were used. In contrast, only inactivated vaccines resulted in a high rate of protective antibodies against FHV-1. Nearly all cats developed high antibody titers against rabies after a single dose of inactivated rabies vaccine.

How long does a single rabies vaccine protect feral cats against infection?
Although there are no reports of long-term evaluation in feral cats, vaccine licensing studies have demonstrated 3–4 year DOI following a single vaccine administered to laboratory kittens. This suggests that while the first rabies vaccine may only be recognized by regulatory agencies as valid for a single year, it is likely that vaccinated cats are protected for much longer. It is recommended that TNR programs offer vaccine booster services for community cats.
Vaccines are biological products that stimulate a series of complex immune reactions that may manifest transient side effects for up to 2 or 3 days following vaccination. It is rare that these self-limiting side effects escalate into serious adverse events. It is advisable to inform clients that their cat may experience reduced appetite or loss of appetite (lasting for two meals), pain at the injection site, lethargy, reluctance to play/walk/run, or mild fever. Treatment is seldom required.

Clients should contact the veterinary practice should any physical or behavioral effects worsen or continue beyond 2–3 days. In the rare event that signs of systemic illness (such as vomiting, diarrhea, seizures, facial swelling, collapse or difficulty breathing) develop, the owner should contact the veterinary practice immediately.

How do I report a vaccine adverse event? The Advisory Panel strongly encourages veterinarians to report all known or suspected vaccine adverse events to the manufacturer and the appropriate regulatory agency responsible for monitoring post-vaccinal adverse events (see details below).

Adverse event FAQs

What constitutes a vaccine adverse event and what should I advise clients to watch for?

Am I legally required to report vaccine adverse events? There is no legal mandate to report post-vaccinal reactions in the USA, Canada or the European Union.

Am I legally liable for using a protocol based on my patient’s risk that differs from the one on the vaccine insert?

Continuous medical decision-making is an inherent aspect of veterinary medicine. There is no reason to believe that decisions regarding vaccine selection and use will carry any greater legal risk than the myriad other medical decisions made in daily practice. Relative risk for utilizing these Guidelines in developing patient vaccination protocols is considered low. Some of the recommendations included in the 2013 AAFP Feline Vaccination Advisory Panel Report will differ from the manufacturer recommendations published in the product package insert/label. However, in most countries, veterinarians in small animal practice have considerable discretion in exercising their judgment relative to the selection and use of licensed veterinary biological products within their professional practice. Rabies vaccination is the obvious exception – veterinarians are required to follow local laws.

(Continued on page 806)
Veterinarians may be held liable for injury or death caused by administration of a vaccine or any other medication. Effective client communication is the best way to avoid legal consequences. Communication of risk and benefit information to clients should be in direct and simple terms. With respect to documentation, practitioners should determine what best suits their practice and their level of risk tolerance. For more information on informed consent (legal considerations), Certificate of Exemption from Rabies Vaccination, or vaccination documentation, please refer to the 2006 Guidelines (see box on page 786).

What do I do if a patient has had a previous vaccine adverse event? Refer to page 796 for recommendations regarding the approach to anaphylaxis and allergic reactions.

References for the FAQs (Appendix 1)

17 Scott F and Geissinger C. Duration of immunity in cats vaccinated with an inactivated feline panleukopenia, herpesvirus, and calicivirus vaccine. Feline Pract 1997; 25: 12–19.
Vaccinations for Your Cat
Pet Owner Guide

WHY DOES MY CAT NEED TO BE VACCINATED?
Vaccines help to protect against specific infectious diseases caused by some viruses and bacteria. They stimulate the body’s immune system to destroy the organism and ‘remember’ it so that it can fight against infection again if necessary in the future. Without vaccination, many cats become seriously ill or may even die from diseases that their immune system is unable to fight effectively on its own. The use of vaccines has prevented death and disease in millions of cats. In addition, vaccines protect people from disease, such as rabies, that could be transmitted from cats.

Some diseases are easier to vaccinate against than others. For example, vaccination is very effective against feline parvovirus infection (panleukopenia) but does not completely protect against respiratory virus infections. However, cats vaccinated against respiratory tract infections generally have milder illness than if they hadn’t been vaccinated and are far less likely to die from the disease.

WHY DOES MY KITTEN NEED A SERIES OF MORE THAN ONE VACCINE?
Newborn kittens depend on their mothers not just for food and warmth, but also for protection against infectious diseases. The first few times they nurse, kittens get antibodies from their mother’s milk that will help to keep them safe for a few weeks to several months. This immunity provided by “maternally derived antibodies” (MDA) is protective while a kitten’s own immune system is immature. However, if the antibody levels decrease before the kitten has developed its own immunity, gaps in protection will occur, leaving the kitten susceptible to disease. Also while the kitten has high levels of MDA, their immune system will not respond optimally to vaccination. Since we cannot predict for each kitten when MDA has decreased adequately to allow an effective response to vaccination, guidelines have been developed to protect as many kittens as possible against disease by giving a series of vaccinations. An incomplete series of kitten vaccinations may leave your kitten vulnerable to infection, so it is important to follow your veterinarian’s recommendations and vaccinate up to at least 16 weeks of age.

You are an important member of your cat’s healthcare team.
You can be instrumental in helping with the success of treatments and improved healthcare for your cat.

The Pet Owner Guide may be downloaded from www.catvets.com/guidelines/client-brochures and is also available as a Supplementary File (see page 799)