Duration of Immunity to Canine and Feline Infectious Disease

- The World Small Animal Veterinary Association states: “We should aim to vaccinate every animal, and to vaccinate each individual less frequently.”
  http://www.wsava.org/SAC.htm

- Age and Long-term Protective Immunity in Dogs and Cats, R.D. Schultz, B. Thiela, E. Mukhtara, P. Sharpa and L.J. Larson, Department of Pathobiological Sciences, School of Veterinary Medicine, University of Wisconsin-Madison:
  “Only one dose of the modified-live canine ‘core’ vaccine (against CDV, CAV-2 and CPV-2) or modified-live feline ‘core’ vaccine (against FPV, FCV and FHV), when administered at 16 weeks or older, will provide long lasting (many years to a lifetime) immunity in a very high percentage of animals ([Schultz, 1998], [Schultz, 2000] and [Schultz, 2006])."

GUIDELINES FOR THE VACCINATION OF DOGS AND ... - WSAVA

1. R.D. Schultz / Veterinary Microbiology 117 (2006) 75–79
5. Schultz RD: Veterinary Vaccines and Diagnostics in Advances in Veterinary Medicine, 41, 1999 pp. 1-853.
Vaccination and Misconceptions

Dr Ronald D Schultz, a leading scientists in this field advises:

A significant number of veterinary practitioners believe:

1. *The annual revaccination recommendation on the vaccine label is evidence the product provides immunity for (only) one year. Not true.*

2. *That they are legally required to vaccinate annually and if they don’t they will not be covered by liability insurance if the animal develops a vaccine preventable disease - Not true. The only vaccine required at all by law [referring to America] is rabies and even that vaccine is not required in some states. There is also a concern that certain companies will not provide assistance if practitioners don’t vaccinate annually with core vaccines. Not true. In fact, all of the major companies have now demonstrated their core products provide at least 3 years of immunity and endorse the “not more often than 3 year” vaccination recommendation made in the AAFP Feline Guidelines and the AAHA Canine Guidelines.*

3. *That not revaccinating will cause the animal to become susceptible soon (days or a few weeks) after the one year. – Not true for the core vaccines. They provide up to a lifetime of immunity or, at the very least, many years of immunity*

4. *If the animal is not revaccinated at or before one year the “whole vaccination program needs to be started again”. – Not true. If the immune response had been stimulated previously, memory cells will persist well beyond a year for the core vaccines.*

5. *If they don’t continue to revaccinate annually, diseases like canine distemper, canine parovirus, feline panleukopenia, and infectious canine hepatitis will “reappear and cause widespread disease similar to what was seen prior to the development of vaccines for these diseases.” – Not true with the core vaccines. It is not how often you revaccinate; it is dependent on how many animals in the population (herd immunity) receive at least one dose of the core vaccines at an age when MDA cannot block active immunity (eg >16 weeks of age).*

6. *That if the revaccination “doesn’t help, it won’t hurt.” – Not true. Vaccines can and do cause adverse reactions, thus don’t administer vaccines if and when they are not needed. Vaccines can cause severe adverse reactions, including death!*

7. *That giving a vaccine annually that has a duration of immunity of 3 or more years provides much better immunity than if the product is given only once during the three or more years. – Not true.*

8. “*It’s much cheaper to revaccinate the pet annually than it is to treat the disease the animal will develop because it didn’t get revaccinated annually." This is the “better safe than sorry” philosophy: it is less expensive to prevent disease. This is why it is necessary to use the core vaccines. However, if the core vaccines are given as a puppy and again at 6 months to a year of age, then annual revaccination is not needed. Furthermore, if a vaccine is given that is not needed and it causes an adverse reaction, this is unacceptable and very expensive.*

9. *They need to revaccinate all new dogs/cats coming to their clinic irrespective of vaccination history even when vaccination records are available from another clinic. Presumably the “other clinic” used the wrong vaccine or didn’t know how to vaccinate. – Not true.*

10. *“Dogs and cats need to be revaccinated annually up to 5 to 7 years of age, then and only then would vaccination every three years be okay.” – Not true. Dogs and cats should be vaccinated as puppies and kittens with the core vaccines, making sure the last dose of vaccine is at 14 to 16 weeks of age. They should be revaccinated again at 6 months to 1 year of age, unless titer were performed; then they need not be revaccinated more often than every 3 years. Also, they do not need to have antibody titers performed more often than every 3 years and only then if you decide not to revaccinate.*
11. “Surgical procedures, including anaesthesia, are immunosuppressive thus dogs should be vaccinated prior to or shortly after surgery.” – Not true. Vaccines should not be given during anaesthesia and animals already vaccinated prior to surgery need not be vaccinated again. If they have never been vaccinated prior to surgery, wait until the animal has recovered from anaesthesia to vaccinate.

12. “Because boarding kennels require annual vaccination, practitioners must continue vaccinating annually with all vaccines.” – Not true. Help change kennel rules through education and just use the vaccines that need to be given (e.g. Kennel Cough.) The kennels need to understand that dogs and cats are up-to-date on their core vaccines when they have been vaccinated within the past 5 to 7 years and no kennel should require core vaccines more often than every 3 years.

**General Scientific References to Vaccine Adverse Effects**

All vaccines come with potential adverse effects. When we vaccinate, we are trading the desire to reduce infectious disease against the known and unknown potential side-effects of vaccines. These side effects include atopy, allergy, autoimmune diseases (including cancer), arthritis, organ failure and neurological effects.

The American Animal Hospital Association (AAHA) Guidelines state:

Do Not Vaccinate Needlessly - Don't revaccinate more often than is needed and only with the vaccines that prevent diseases for which that animal is at risk. They also caution veterinarians: "Do not assume that vaccines cannot harm a patient. Vaccines are potent medically active agents and have the very real potential of producing adverse events."

The American Veterinary Medical Association's (AVMA's) Principles of Vaccination (http://www.avma.org/issues/vaccination/vaccination.asp) states that:

Possible adverse events include failure to immunize, anaphylaxis, immunosuppression, autoimmune disorders, transient infections, and/or long-term infected carrier states. In addition, a causal association in cats between injection sites and the subsequent development of a malignant tumor is the subject of ongoing research.

Dr Ronald D Schultz (2007) provides this list of adverse events known to be induced by vaccines:

**Common Reactions:**

- Lethargy
- Hair Loss, hair color change at injection site
- Fever
- Soreness, stiffness
- Refusal to eat
- Conjunctivitis
- Sneezing
- Oral ulcers

**Moderate Reactions:**

- Immunosuppression
- Behavioral changes
- Vitiligo (skin reactions)
- Weight loss (Cachexia)
- Reduced milk production
- Lameness
- Granulomas/Abscesses
- Hives
- Facial Edema (swelling)
- Atopy (hereditary allergies)
- Respiratory disease
- Allergic Uveitis (Blue Eye)

**Severe Reactions triggered by Vaccines:**

- Vaccine injection site sarcomas (in dogs and ferrets as well as cats)
- Anaphylaxis
- Arthritis, polyarthritis
- HOD hypertrophy osteodystrophy
- Autoimmune Hemolytic Anemia
- Immune Mediated Thrombocytopenia (IMTP)
- Hemolytic disease of the newborn (Neonatal Isoerythrolysis)
- Thyroiditis
- Glomerulonephritis
- Disease or enhanced disease which the vaccine was designed to prevent
- Myocarditis
- Post vaccinal Encephalitis or polyneuritis
- Seizures
- Abortion, congenital anomalies, embryonic/fetal death, failure to conceive

The British Veterinary Medicines Directorate listed the following adverse reactions in its pet vaccine position paper:

- Immune mediated diseases and the association with vaccine reactions have been reviewed by Day (1999, 2006), Pedersen (1999) and the VPC Working Group (2002). Vaccine reactions are generally classified into one of four recognised categories: Type I-IV. The immunological theories behind such reactions would suggest that such adverse events should only occur following an immunological reaction to a previously exposed vaccine antigen, adjuvant, excipient or other production remnants such as bovine serum. However, immune mediated reactions can also follow the administration of a primary dose of vaccine and the exact mechanisms for such a reaction are unknown.

- Type I hypersensitivity reactions involve an immune mediated reaction that releases potent inflammatory mediators and other chemicals that trigger an anaphylactic reaction in the affected animal. The reactions are usually acute, with the clinical signs appearing within minutes or hours of vaccination. Typical signs reported are facial oedema, shock, lethargy, respiratory distress and diarrhoea. Severe anaphylactic reactions may result in death. Urticaria (hives), facial oedema and anaphylactic shock are specific clinical manifestations of Type I hypersensitivities.

- Type II hypersensitivity reactions involve the binding of the animal’s own antibodies to cells or a cell matrix. The formation of auto-immune antibodies is thought to involve a number of complex immunological mechanisms. Secondary immune mediated haemolytic anaemia (IMHA) has been associated with vaccination but may also occur following infection, neoplasia or administration of medications. In this condition, auto-antibodies are produced against the animal’s own red blood cells but the immunological mechanism by which vaccines may produce such a response is not yet established. Confirmation of IMHA is dependent on the demonstration of auto-antibodies and, therefore, not all reports of such adverse events can be recorded accurately if confirmatory diagnostic tests have not been performed. For the period 2005-2010, 57 suspect adverse reaction reports of immune mediated reactions were submitted to the VMD. Subsequently only 25 of these reports were attributed to immune mediated reactions. Other clinical manifestations of Type II disease include immune mediated thrombocytopenia (IMTP) (autoantibodies to blood platelets) myasthenia gravis (autoantibodies to muscle nerve receptors) and pemphigus (foliaceous & vulgaris) (auto-antibodies to epidermal proteins).

- Type III hypersensitivity reactions result from the formation of circulating complexes of antigen and antibody that deposit in certain organs or tissues in the body leading to inflammatory reactions and destruction of cells and associated matrix. The deposition of immune complexes usually results in inflammation of the blood vessels. The reaction is dependent on the continued presence of both antibody and antigen with the latter being derived from infection, vaccination, medication or exposure of ‘self-antigens’ through disease. Infections will inevitably result in the formation of immune-complexes as the body aims to rid itself of a foreign invader. Some examples of well known immune-complexes provide clinical signs such as:
(i) ‘Blue Eye’ is a well documented manifestation of a Type III reaction following infection with CAV-1 and or administration of some of the early CAV-1 vaccines. Replacement of CAV-1 by CAV-2 vaccines have minimised the risk of such adverse events with just six reports of ‘blue eye’ reported to the VMD during the period 2005-2010.

(ii) Reactions involving type III immune mediated mechanisms have also been demonstrated following rabies virus vaccination.

(iii) Systemic lupus erythematosus (SLE), a disease characterised by the development of antinuclear antibodies (ANAs).

(iv) Drug-induced arthritis has been reported as a Type III reaction but the evidence of similar vaccine associated immune-mediated syndrome is sparse. The VMD has received two reports of pemphigus associated with vaccination during the last five years.

- Type IV hypersensitivity reactions or “delayed-type hypersensitivity” are cell, rather than antibody, mediated. These diseases are usually relatively slow to develop and are dependent on the cell-mediated arm of the immune system.

Dr Jean W Dodds, wrote in US Dog World, March, 1995:

Immune–suppressant viruses of the retrovirus and parvovirus classes have recently been implicated as causes of bone marrow failure, immune-mediated blood diseases, haematologic malignancies (lymphoma and leukaemia), dysregulation of humoral and cell-mediated immunity, organ failure (liver, kidney) and autoimmune endocrine disorders — especially of the thyroid gland (thyroiditis), adrenal gland (Addison’s disease) and pancreas (diabetes). Viral disease and recent vaccination with single or combination modified live virus vaccines, especially those containing distemper, adenovirus 1 or 2 and parvovirus, are increasingly recognised contributors to immune-mediated blood diseases, bone marrow failure and organ dysfunction.

http://www.dogs4dogs.com/shots.html:

“Although any vaccine, including those your dog has previously gotten without incident, can cause an adverse reaction, vaccines made with killed virus, however, are the most adversely reactive. This includes the rabies vaccine, Bordetella, Coronavirus, and Leptospirosis. Why? Because killed vaccines contain adjuvants (additives that “boost” the immune reaction). Live viruses don't need boosting. Fortunately, with the exception of rabies, these other shots are not recommended for all dogs. Also, they are seldom effective. Don't give them without good reason.”

Veterinary Research relating to Vaccines, Immunity, and Adverse Effects:


34. J. S. Munday, N. L. Stedman and L. J. Richey. Athens Diagnostic Laboratory and Department of Veterinary Pathology, College of Veterinary Medicine, University of Georgia, Athens, GA. Histology and Immunohistochemistry of Seven Ferret Vaccination-site Fibrosarcomas. Vet Pathol 40:288-293 (2003). www.vetpathology.org

35. Gregory K. Ogilvie, DVM, DACVIM (Internal Medicine, Oncology). Injection Site and Vaccine Associated Sarcomas: New Advances for a New Millennium. 29th World Congress of the World Small Animal Veterinary Association: October 6 – 9, 2004: Rhodes, Greece. www.vin.com


51. Schultz, R.D., 2004. Results presented at 2004 American College of Veterinary Internal Medicine Convention, Minneapolis MN and American Veterinary Medical Association Convention, Philadelphia PA and available in printed material “Canine Distemper and Vaccination” from Merial Limited, Duluth, GA.

52. R.D. Schultz / Veterinary Microbiology 117 (2006) 75–79 79

53. R.D. Schultz / Veterinary Microbiology 117 (2006) 75–79 79


59. Vaccine-Associated Feline Sarcoma Task Force: Roundtable Discussion - The current understanding and management of vaccine-associated sarcomas in cats. JAVMA: June 1, 2005; Vol. 226, No. 11. www.avma.org


62. Veterinary Therapeutics, Vol. 5, No. 3, Fall 2004

Vaccine adjuvants are chemicals, microbial components, or mammalian proteins that enhance the immune response to vaccine antigens. Interest in reducing vaccine-related adverse effects and inducing specific types of immunity has led to the development of numerous new adjuvants. Adjuvants in development or in experimental and commercial vaccines include aluminum salts (alum), oil emulsions, saponins, immune-stimulating complexes (ISCOMs), liposomes, microparticles, nonionic block copolymers, derivatized polysaccharides, cytokines, and a wide variety of bacterial derivatives. The mechanisms of action of these diverse compounds vary, as does their induction of cell-mediated and antibody responses. Factors influencing the selection of an adjuvant include animal species, specific pathogen, vaccine antigen, route of immunization, and type of immunity needed.

Extract:

**Adverse Effects and Potential Hazards of Adjuvants**

When immune responses destroy invading microorganisms, they cause tissue damage and result in some of the clinical signs of illness. Similarly, as agents that enhance immune responses, adjuvants can increase the adverse effects of the vaccine. These adverse effects are influenced by the interactions of the specific adjuvant and antigen.9,20

Systemic, nonspecific adverse effects can include fever, arthritis, uveitis, anorexia, soreness, and lethargy.9,20,21 Theoretically, adjuvants also may increase the probability of autoimmune reactions. Overdoses of IL-2, a cytokine proposed as an adjuvant, have been linked to autoimmune diseases.22 Autoantibodies have been detected after vaccination with typical canine distemper, rabies, and parvovirus vaccines, and a temporal association has been noted between autoimmune hemolytic anemia and vaccination in dogs.21 Adjuvants also may have specific adverse effects related to their chemical nature. For example, some crude saponin adjuvants can result in hemolysis if injected IV.23

W. Jean Dodds DVM on vaccine adverse effects:

“I don’t really believe my training in veterinary college was adequate with regard to animal vaccination. To be fair to current knowledge, I graduated from veterinary college in Canada in 1964. At that time we understood much less about the molecular aspects of immunology, and the long term medical effects of vaccinations, both beneficial and potentially harmful. We also had fewer infectious diseases to treat and prevent, and hence fewer vaccines for them.

“I was always a clinical research scientist, but colleagues and pet owners would tell me about malaise and illness that appeared shortly after pet animals were vaccinated. This was in addition to the rare case of anaphylaxis induced by vaccination. They spoke about irritability, low-grade or even high fever, anorexia, stiffness, occasional seizure-like episodes. These usually occurred from 2-10 days post-vaccination, sometimes longer (up to 45 days).

“The clinical signs associated with vaccine reactions typically include fever, stiffness, sore joints and abdominal tenderness, susceptibility to infections, neurological disorders and
encephalitis, collapse with autoagglutinated red blood cells and icterus (autoimmune hemolytic anemia) (AIHA), or generalized petechiae and ecchymotic hemorrhages (immune-mediated thrombocytopenia) (ITP).  

“Hepatic enzymes may be markedly elevated, and liver or kidney failure may occur by itself or accompany bone marrow suppression. Furthermore, MLV vaccination has been associated with the development of transient seizures in puppies and adult dogs of breeds or cross-breeds susceptible to immune-mediated diseases especially those involving hematologic or endocrine tissues (e.g. AIHA, ITP, autoimmune thyroiditis).”  

Post-vaccinal polyneuropathy is a recognized entity associated occasionally with the use of distemper, parvovirus, rabies and presumably other vaccines. This can result in various clinical signs including muscular atrophy, inhibition or interruption of neuronal control of tissue and organ function, muscular excitation, incoordination and weakness, as well as seizures.  

“Certain breeds or families of dogs appear to be more susceptible to adverse vaccine reactions, particularly post-vaccinal seizures, high fevers, and painful episodes of hypertrophic osteodystrophy (HOD).” Vaccination of pet and research dogs with polyvalent vaccines containing rabies virus or rabies vaccine alone was recently shown to induce production of antithyroglobulin autoantibodies, a provocative and important finding with implications for the subsequent development of hypothyroidism.  

Furthermore, injection site fibrosarcomas have recently been documented in dogs as well as cats, and other cancers such as leukemia have been vaccine-associated.”  

“If I was in private practice today, I would only use a conservative puppy or kitten series (2-3 doses only) of vaccines: one before 12 weeks in puppies and 10 weeks in kittens; and a second between 14-16 weeks in puppies and 12-14 weeks in kittens. All vaccines should be 3-4 weeks apart. I would not vaccinate beyond the puppy and kitten series, and I would not worry about income impact, regardless, as our veterinary oath requires that we “do no harm”. Judicious use of vaccines is paramount.  

“I totally embrace my profession and always have, but the pharmaceutical industry has considerable influence on it; there is a huge marketing effort here – that’s their job. We are the ones that need to ‘sift’ this information appropriately. We, the consumer professionals, have allowed this influence to go unchecked. It’s time for senior members of our profession to step up and place controls on the commercial influence upon relatively naive veterinary students and new graduates. This influence is even stronger in the pet food and supplements industry. The government also needs to be more proactive and keep up to date.”

References


Note:
Dr. Dodds received the D.V.M. degree with honors in 1964 from the Ontario Veterinary College, University of Toronto. In 1965 she accepted a position with the New York State Health Department in Albany and began comparative studies of animals with inherited and acquired bleeding diseases. Her position there began as a Research Scientist and culminated as Chief, Laboratory of Hematology, Wadsworth Center. In 1980 she also became Executive Director, New York State Council on Human Blood and Transfusion Services. This work continued full-time until 1986 when she moved to Southern California to establish Hemopet, the first nonprofit national blood bank program for animals.

From 1965-1986, she was a member of many national and international committees on hematology, animal models of human disease, veterinary medicine, and laboratory animal science. Dr. Dodds was a grantee of the National Heart, Lung, and Blood Institute (NIH) and has over 150 research publications. She was formerly President of the Scientist's Center for Animal Welfare; and Chairman of the Committee on Veterinary Medical Sciences and Vice-Chairman of the Institute of Laboratory Animal Resources, National Academy of Sciences.

In 1974 Dr. Dodds was selected as Outstanding Woman Veterinarian of the Year, AVMA Annual Meeting, Denver, Colorado; in 1977 received the Region I Award for Outstanding Service to the Veterinary Profession from the American Animal Hospital Association, Cherry Hill, New Jersey; in 1978 and 1990 received the Gaines Fido Award as Dogdom's Woman of the Year; and the Award of Merit in 1978 in Recognition of Special Contributions to the Veterinary Profession from the American Animal Hospital Association, Salt Lake City, Utah. In 1984 she was awarded the Centennial Medal from the University of Pennsylvania School of Veterinary Medicine. In 1987 she was elected a distinguished Practitioner of the National Academy of Practice in Veterinary Medicine. In 1994 she was given the Holistic Veterinarian of the Year Award from the American Holistic Veterinary Medical Association. She is an active member of numerous professional societies.
Stephen Blake DVM, holistic veterinarian:

“I emphatically do not believe that I was taught adequately with regard to the vaccine issue. I was taught to believe vaccinations were synonymous with immunization. They are two separate entities.

“I was taught vaccines were safe and it was implied there had been safety studies done on them before they were used on the general public. They are not safe and there have not been any safety studies done on any of them.

“I was taught that if something adverse happens within a few hours after immunization it was related to the vaccines but, if it happened later than that period of time, it had nothing to do with the vaccines. The truth of the matter is vaccines can set up a latent condition that may show up within a few hours or years after immunization.

“I was taught you needed to vaccinate every year to boost animals’ immune systems. There are no studies to show that annual boosters are ever indicated or that there is any science to support annual vaccines to boost an animal’s immune system. I was never taught that mercury and aluminium hydroxide, which are in the vaccines, can cause cancer, are neurotoxins, and can trigger autoimmune disease.

“When I first went into practice, I noticed some animals developed fevers for a few days, became lethargic, lost their appetites, developed ear infections, seizures, pruritus, UTI, musculoskeletal issues, and behaviour issues after vaccination.

“As time went on, I observed that otitis, UTI, autoimmune disease complexes, gingivitis, allergic dermatitis, IBD, asthma, aggression and phobias, convulsions, paralysis, cancer, chronic conjunctivitis, liver disease, kidney disease, cardiac disease, arthritis, anterior cruciate rupture, hip disease, and corneal lesions can be correlated with vaccine damage.

“Pet guardians are being misinformed. There is no scientific evidence that annual vaccines are needed or indicated.

“I feel there is pressure at all levels of veterinary medicine that we are not to say anything negative about vaccines which would alarm the public and make it harder to sell the vaccine concept. There is no informed consent information presented to the pet owner prior to vaccination because they do not want to alarm the owner and make it difficult to promote vaccines. It has been known for over 20 years that rabies vaccines can cause terminal cancer in the feline and I have yet to meet a client who was informed of this scientific fact by their attending veterinarian. This should be made public to all cat owners prior to rabies vaccine so the owner is aware of the low risk of getting rabies compared to the high risk of getting cancer from the rabies vaccine.

“I love the practice of veterinary medicine with an oath to prevent suffering and do no harm. I do not support my profession’s over-use of drugs, chemicals and vaccines as we know them today. I feel my profession needs to be the leader in breaking away from the dangers of these products and show the human medical profession this is not healing. It does cause harm.

“I feel the pharmaceutical industry finances the veterinary schools and the veterinary profession. I feel they are the fox in the hen house that sets policy for the practice of medicine as we know it today in our country.

“The pharmaceutical industry has too much influence in veterinary teaching. Their approach to medicine prevents any other modalities of healing from being available in our veterinary schools. This is done so they have no competition for the pet industry dollars from unpatentable means.”

Duration of Immunity

“Many Veterinary practices still recommend vaccinations annually, yet the recommendation for annual vaccination is a practice that was officially started in 1978. That recommendation was made without scientific validation of the need to booster immunity so frequently. In fact, the presence of good humoral antibody levels can block the anamnestic response to vaccine boosters just as maternal antibody blocks the response in some young animals. In other words, if a body has been given its initial vaccine and first year booster, giving another vaccine does not create a stronger immunity. However, the danger of over-vaccinating is a very real risk. (Schultz, Ronald D. “Current and future canine and feline vaccination programs”, Veterinary Medicine, March 1998, p. 243)
Dr. Ronald Schultz is one of the leading independent researchers and proponents of vaccinating animals less often. Shultz began researching vaccines more than twenty-five years ago when he noticed the two different vaccination approaches for humans and animals—humans were vaccinated as children and then not again...but animals were vaccinated annually. His research confirms that most animal vaccines, like human ones, create long-term immunity. Shultz emphasizes that while it's critical to stimulate initial immunity in animals when they are young, his work has revealed that many vaccines provide lifelong immunity, making repeated vaccinations after the first year of doubtful value. To compound the situation, Shultz has found that indiscriminate vaccination of adult animals can actually trigger adverse physiological reactions.

In studies Dr. Ron Shultz performed at the University of Wisconsin, 106 dogs vaccinated within the previous one to four years were each given a canine parvovirus booster vaccine. Only one of the 106 dogs showed significant increase in serum antibody titer following the booster. These results demonstrated that revaccination does not automatically enhance antibody levels or improve immunity. What happens is that the vaccine virus is neutralized before it can reach the memory T and B cells. The immunity provided by previous vaccination not only protects against the virulent disease but also prevents response to revaccination. (Wolf Alice, Vaccines of the Present and Future, Proceedings of the World Animal Veterinary Congress, Vancouver 2001.)

The term “up-to-date” is only valid if you use the vaccine manufacturers’ protocols, which are reflective of the amount of animal testing the vaccine company has actually done with a particular vaccine.

**Delayed Vaccine Reactions**

Animal studies show delayed reactions to vaccine ingredients. A reaction may occur days, weeks, months or years after an animal or human is exposed to vaccines. (*Food and Drug Administration, 1982* as cited in Kirby, 2005, St. Martins Griffin, New York.)

Viral disease and recent vaccination with single or combination modified live-virus (MLV) vaccines—especially that containing distemper virus, adenovirus 1 or 2, and parvovirus—is an increasingly recognized contributor to immune-mediated blood disease, bone marrow failure, and organ dysfunction. (1-11) Potent adjuvanted killed vaccines, like those for rabies virus, can also trigger immediate and delayed (vaccinosis) adverse vaccine reactions. Beyond immediate hypersensitivity reactions, other acute events tend to occur 24-72 hours afterward, or 7-45 days later in a delayed type immunological response." (1-4, 6-10)

Even more delayed adverse effects include mortality from high-titered measles vaccine in infants, canine distemper antibodies in joint diseases of dogs, and feline and canine injections-site fibrosarcomas.‖ (5, 7)

The increasing antigenic load presented to the host individual by modified-live virus (MLV) vaccines during the period of viremia is presumed to be responsible for the immunological challenge that can result in a delayed hypersensitivity reaction. (2,3,6,7)

The clinical signs associated with vaccine reactions typically include fever, stiffness, sore joints, and abdominal tenderness, susceptibility to infections, neurological disorders and encephalitis, collapse with autoagglutinated red blood cells, and icterus (autoimmune hemolytic anemia—AIHA), or generalized petechiae and ecchymotic hemorrhages (immune-mediated thrombocytopenia—ITP). (1, 2, 4, 7, 8, 12, 13) Hepatic enzymes may be markedly elevated, and liver or kidney failure may occur by itself or accompany bone marrow suppression. Furthermore, MLV vaccination has been associated with the development of transient seizures in puppies and adult dogs of breeds or cross-breeds susceptible to immune-mediated diseases, especially those involving hematologic or endocrine tissues (i.e. AIHA, ITP, autoimmune thyroiditis).‖ (1, 7, 10)

"Post-vaccinated polyneuropathy is a recognized entity associated occasionally with the use of distemper, parvovirus, rabies, and presumably other vaccines." (2, 3, 7) "This can result in various clinical signs including muscular atrophy, inhibition or interruption of neuronal control
of tissue and organ function, muscular excitation, uncoordination and weakness, as well as seizures." (7)

Certain breeds or families of dogs appear to be more susceptible to adverse vaccine reactions, particularly post-vaccinal seizures, high fevers, and painful episodes of hypertrophic osteodystrophy (HOD). (7, 9)

Vaccination of pet and research dogs with polyvalent vaccines containing rabies virus or rabies vaccine alone was recently shown to induce production of antithyroglobulin autoantibodies, a provocative and important finding with implications for the subsequent development of hypothyroidism." (10, 17)

Furthermore, injection site fibrosarcomas have recently been document in dogs as well as cats." (18)

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15. Kyle AHM, Squires RA, Davies PR. Serologic status and response to vaccination against canine distemper (CDV) and canine parvovirus (CPV) of dogs vaccinated at different intervals. J Sm An Pract, June 2002.


Vaccines and Cancer


2. Isolation of an Infectious Endogenous Retrovirus in a Proportion of Live Attenuated Vaccines for Pets, Journal of Virology, April 2010, p. 3690-3694, Vol. 84, No. 7. “It is impossible to rule out chronic effects, especially as we were able to grow RD-114 very efficiently in dog cell lines, confirming older published studies…. A recently identified novel human retrovirus (xenotropic murine leukemia virus-related retrovirus [XMRV]) has been found in some forms of prostate cancers and chronic fatigue syndrome in humans, although causal association has not been proven yet.”


4. “When canine distemper virus was combined with canine adenovirus type 1 or canine adenovirus type 2, significant suppression in lymphocyte responsiveness to mitogen occurred. The results indicate that interactions between canine distemper virus and canine adenovirus type 1 or canine adenovirus type 2 are responsible for the polyvalent vaccine induced suppression of lymphocyte responsiveness”. (Effects of Vaccines on the Canine Immune System, Tom R. Phillips, Jean L. Jensen, Michael J. Rubino, Wen C. Yang and Ronald D. Schultz, Can J Vet Res 1989; 53: 154-160)

5. JFM Series A, August 2003, vol 50, no 6, pp 286-291

6. “Live attenuated rubella vaccine inoculation may cause sustained immunosuppression including defective lymphocyte response to mitogene and impaired cytokine production. The signs of immunosuppression may persist for at least 1 month after vaccination”. (Cytokine profile after rubella vaccine inoculation: evidence of the immunosuppressive effect of vaccination, Mediators of Inflammation, 12(4), 203-207 (August 2003)).

7. Cancer Research 30, October 1970, ‘Spontaneous Development of Mammary Adenocarcinoma following Prolonged Immunosuppression in the Dog’: “Live attenuated rubella vaccine inoculation may cause sustained immunosuppression including defective lymphocyte response to mitogene and impaired cytokine production. The signs of immunosuppression may persist for at least 1 month after vaccination”.

8. “If your vaccine is manufactured in a cell substrate that was derived from a tumor, or that has a tumorigenic phenotype through an unknown mechanism, it might carry a higher theoretical risk of containing oncogenic [tumour forming] substances.” FDA “Guidance for Industry, Characterization and Qualification of Cell
Substrates and Other Biological Materials Used in the Production of Viral Vaccines for Infectious Disease Indications

9. Dyer et al, 2007: “The most significant problems associated with feline vaccines have been injection-associated sarcoma. Previously, this problem seemed most apparent in cats administered adjuvanted rabies virus and feline leukaemia virus vaccines. However, recent information suggests that injection site sarcomas can occur with any type of vaccine. For example, in the United Kingdom in 2005, 23 of 39 injection site sarcomas reported in cats occurred at the site a live vaccine (non-adjuvanted) was administered.”

10. IARC International Agency for Research on Cancer; Summaries and Evaluations Surgical Implants and Other Foreign Bodies 1999 Feb 23; 74:24305-310. “The adjuvant aluminum in vaccines is one culprit in mutating the genome and specifically the P53 oncogene, thereby ruining the individual’s ability to stop tumor genesis.”


Vaccines and Immune-Mediated Disease

1. “There is a real concern that vaccines may predispose certain genetically susceptible individuals to immune-mediated disease. The more antigens we administer, the higher the potential for hypersensitivity. Type I is IgE mediated; type 2, cytotoxic antibody mediated; type 3, immune-complex mediated; and type 4 cellular mediated. All of these hypersensitivities are natural parts of the immune response, but they cause a certain amount of tissue damage. That damage may occur in the kidney, liver, or as was the case with canine adenovirus 1, in the eye. In many cases it is impossible to show a direct connection between the damage and a vaccine, since it is the accumulation of many antigens over many years that results in clinically evident disease.” JAVMA, Vol 207, No 4, August 15, 1995 – Current Concepts, are we vaccinating too much?

2. The seventh edition of the Merck Veterinary Manual (1991) states: “Bone marrow suppression with transient (21 day) or chronic/latent erythroid dysplasia, in the presence or absence of thrombocytopenia and neutropenia, Combs’ positive haemolytic anaemia, and immune-mediated thrombocytopenia have been associated with (i.e., may prove to be caused by) both retroviral and parvoviral infection in man and other species. Also, modified live parvovirus vaccines in dogs, and killed feline leukaemia virus vaccine are suspects as causes (in genetically susceptible animals) of such haematological diseases.”

3. In Practice, Vol 20 No 2, Feb 1998: Michael Day, senior lecturer in Veterinary Pathology at the University of Bristol states that “environmental influences are crucial to the expression of immune mediated disease and that the most important of these is likely to be exposure to microbial antigens following natural infection or vaccination. Mr Day divides immune mediated disease into four main groups – hypersensitivity diseases, autoimmune diseases, immune system neoplasia [tumour formation] and immunodeficiency diseases.”

4. In a letter to Veterinary Times during July 1999, veterinarian Lyn Thomson responded, “This would indicate that veterinarians must consider and report the whole range of immune mediated diseases post vaccination, including flea allergy, atopic dermatitis, dietary hypersensitivity, contact hypersensitivity, asthma, autoimmune diseases, lymphoma, lymphoid leukaemia, multiple myeloma, plasmacytoma, hisiocyctoma, thymoma, and immunodeficiency disease.”


7. In 2000, research showed that polyarthritis and other diseases like amyloidosis in dogs were linked to combined MLV vaccines (Am Coll Vet Intern Med, 2000; 14: 381).


10. Dr Ronald Schultz, Vet Med Today (JAVMA Vol 207, No 4, August 15, 1995): “Immune-mediated disease has developed in human beings following vaccination, as was seen with cases of Guillain-Barre syndrome following swine flu vaccinations, and rheumatoid arthritis following influenza vaccination”.


**Vaccines and Atopy**


2. An augmented immune response to vaccination is seen in dogs with pre-existing inhalant allergies (i.e., atopy) to pollens. Furthermore, the increasing current problems with allergic and immunological diseases have been linked to the introduction of MLV vaccines more than 20 years ago. While other environmental factors no doubt have a contributing role, the introduction of these vaccine antigens and their environmental shedding may provide the final insult that exceeds the immunological tolerance threshold of some individuals in the pet population. (Tizard 1. *Risks associated with use of live vaccines*. J Am Vet Med Assoc 1990;196:1851-1858.) (Dodds WJ. *More bumps on the vaccine road*. Adv Vet Med 1999; 41: 715-732.)


4. Familial predisposition to atopy is reflected in the Merck Manual which advises that a child with, or from a family with, B and/or T cell immunodeficiencies should not receive live virus vaccines due to the risk of fatality. Merck states: “Features of B cell deficiencies include respiratory or food allergies; features of T cell deficiencies include heart disease; and features of combined T and B cell deficiencies include dermatitis, neurological deterioration and eczema.”

5. “*Multiple vaccinations shift this delicate balance [between Th1 and Th2], favoring the development of atopy and, perhaps, autoimmunity through vaccine-induced polyclonal activation leading to autoantibody production.*” An increase in the incidence of childhood atopic diseases may be expected as a result of concurrent vaccination strategies that induce a Th2-biased immune response. What should be discussed is whether the prize of a reduction of common infectious diseases through a policy of mass vaccination from birth is worth the price of a higher prevalence of atopy.” [http://vran.org/health-risks/anaphylaxis-allergies-and-asthma/multiple-vaccination-effects-on-atopy/](http://vran.org/health-risks/anaphylaxis-allergies-and-asthma/multiple-vaccination-effects-on-atopy/)


7. "*Expression and characterization of a low molecular weight recombinant human gelatin: development of a substitute for animal-derived gelatin with superior features.*" "*Gelatin is used as a stabilizer in several vaccines. Allergic reactions to gelatins have been reported, including anaphylaxis.*” These gelatins are derived from animal tissues and thus represent a potential source of contaminants that cause transmissible spongiform encephalopathies.” Olsen D, et al, Protein Expr Purif.; 40(2):346-57 -- 4/1/2005

8. Professor Tara Shirakawa, published the results of a Japanese *study on 867 infants who received BCG vaccine*. Thirty-six percent developed allergies. The number of
TB cases in the province didn’t increase, but the incidence of severe allergy did.
(Science, vol 275, 3 Jan 1997)


10. Pediatric Allergy & Immunology (19 (1): 46-52, February 2008), looked at the potential causal factor in the development of **atopic disease due to the effect of pertussis immunization on specific IgE antibodies**. All associations between vaccination and atopic disorders were positive. The report concluded: “Egg-related allergy is common, particularly in children with asthma or general allergies, and may be as high as 40% in children with atopic dermatitis. The risk of egg-related allergy after vaccination depends on the presence of egg protein in the final product.”

11. Allergy 1978, Jun;33(3):155-9 reported that **aluminium phosphate stimulates the IgE response in guinea pigs to tetanus toxoid**. “It is hypothesized that the regular application of aluminium compound-containing vaccines … could be one of the factors leading to the observed increase of allergic diseases.”

12. In Pediatr Allergy Immunol 1994 May;5(2):118-23, the **role of aluminium for IgG and IgE responses to pertussis toxin, as well as side effects, was investigated in 49 children with known atopy status**. The addition of aluminium to pertussis vaccine was associated with strong IgG antibody response, and a stronger IgE antibody response. The study concluded that the role of immunization in the development of allergy merits further studies.

13. [http://www.ncbi.nlm.nih.gov/pmc/articles/PMC2998561/](http://www.ncbi.nlm.nih.gov/pmc/articles/PMC2998561/) looks at the **role of IgE antibodies in atopic dermatitis and refers to post-vaccination specific IgE responses** to tetanus and diphtheria toxoid, which could result in adverse events to future vaccination or exposure to the diseases.


16. H Odelram, M Granstrom, S Hedenskog, K Duchen, B Bjorkstein, **Immunogloblin E and G responses to pertussis toxin after booster immunization in relation to atopy, local reactions and aluminium content of the vaccines**, Pediatric Allergy and Immunology, 5 (2), 118-123, May 1994


**Neurological effects of vaccines**


2. Veterinary Record 1992 (130, 27-30), AIP McCandlish et al: "*Post-vaccinal encephalitis is a recognised complication of the administration of certain strains of live attenuated canine distemper vaccine* (Hartley 1974, Bestetti and others 1978, Cornwell and others 1988)."

3. Braund’s Clinical Neurology in Small Animals: Localisation, Diagnosis and Treatment:

   “*Post vaccinal canine distemper encephalitis* occurs in young animals, especially those less than six months of age. It has been recognised as a disease entity for a number of years, and is believed to be association with vaccination using live virus. The pathogenesis of this disease is unclear, but may result from insufficient attenuation of the vaccine virus which causes subsequent infections of the CNS; the triggering of a latent distemper infection by vaccination; other vaccine components; or an enhanced susceptibility of the animal (e.g., animals that are immunosuppressed)."


6. Protein glutamate is added to vaccines to preserve the virus in vaccines. Meat, fish eggs, milk and cheese tend to be high in protein glutamate. High levels of glutamic acid have been shown in animal studies to cause damage to parts of the brain unprotected by the blood-brain barrier, leading to a variety of chronic diseases http://www.ncbi.nlm.nih.gov/pubmed/15167034


9. *Myelin basic protein as an encephalitogen in encephalomyelitis and polyneuritis following rabies vaccination*: "Encephalitis and polyneuritis occurring after rabies vaccination are believed to be immunologically mediated. We studied antibody responses

10. Risks of convulsion and aseptic meningitis following measles-mumps-rubella vaccination in the United Kingdom. "Measles-mumps-rubella (MMR) vaccines containing the Urabe strain of mumps were withdrawn in the United Kingdom in 1992 following demonstration of an increased risk of aseptic meningitis 15-35 days after vaccination. Following introduction of a replacement MMR vaccine (Priorix)... an elevated relative incidence of convulsion was found in the 6- to 11-day period after receipt of Priorix." Miller E, et al, Am J Epidemiol.;165(6):704-9. -- 3/15/2007

11. Neurological adverse events associated with vaccination: "These complications include autism (measles vaccine), multiple sclerosis (hepatitis B vaccine), meningoencephalitis (Japanese encephalitis vaccine), Guillain-Barre syndrome and giant cell arteritis (influenza vaccine), and reactions after exposure to animal rabies vaccine. Seizures and hypotonic/hyporesponsive episodes following pertussis vaccination and potential risks associated with varicella vaccination, as well as vaccine-associated paralytic poliomyelitis following oral poliovirus vaccine, are also described." Piyasirisilp, Sucheep a; Hemachudha, Thiravat b, Neurology. 15(3):333-338 -- 6/1/2002


13. Neurologic complications of immunization. "Individual vaccines can produce systemic or neurologic reactions ranging from minor events, such as pain and erythema at the injection site, to major complications, such as seizures, shock, encephalopathy, or death." Bale JF Jr, J Child Neurol.; 19(6): 405-12. -- 6/1/2004

14. Merck: "In acute disseminated encephalomyelitis (post infectious encephalitis), demyelination can occur spontaneously, but usually follows a viral infection or inoculation (or very rarely a bacterial vaccine), suggesting an immunologic cause."

15. "Expression and characterization of a low molecular weight recombinant human gelatin: development of a substitute for animal-derived gelatin with superior features." "Gelatin is used as a stabilizer in several vaccines. Allergic reactions to gelatins have been reported, including anaphylaxis. These gelatins are derived from animal tissues and thus represent a potential source of contaminants that cause transmissible spongiform encephalopathies." Olsen D, et al, Protein Expr Purif.; 40(2):346-57 -- 4/1/2005


17. Toxicol Environ Chem 2008 90(5):997-1008, researchers found a correlation between the Hepatitis B triple series vaccine and developmental disability in US children aged 1-9 years


31. Aluminum Vaccine Adjuvants: Are they Safe? L. Tomljenovic, and C.A. Shaw, Current Medicinal Chemistry, 2011, 18, 2630-2637 “Aluminum presented in this form carries a risk for autoimmunity, long-term brain inflammation and associated neurological complications and may thus have profound and widespread adverse health consequences.”

32. Department of Paediatrics, Tokyo Medical University, Japan, found the measles virus in patients with inflammatory bowel disease and autism. (Dig Dis Sci, 2000, Apr; 45(4) 723-9) . The sequences obtained from the patients with ulcerative colitis and children with autism were consistent with vaccine strains. It should be remembered that the measles virus and canine distemper are very closely related.

33. "Neurologic complications of smallpox vaccine" "A variety of neurologic complications of smallpox vaccination have been reported, including encephalitis, transverse myelitis, meningitis, and polyneuritis" Lanska, D, Meldink.com: Neurology -- 1/1/1900

34. The question of encephalitis following vaccination was investigated by the League of Nations, and on August 27, 1928, the League published a report on the situation. The report stated: "The post-vaccinal encephalitis with which we are dealing has become a problem of itself . . . a new, or at least previously unsuspected or unrecognized risk attaches to vaccination. . . ."
35. The Journal of the American Medical Association on April 2, 1937: "A multiplicity of untoward sequelae have been observed in patients treated with immune serum... The common symptomatology includes fever, urticaria, erythema, oedema, lymphadenoma, arthralgia, smothering sensations, headache, nausea and vomiting. Occasionally there are more serious and lasting manifestations such as peripheral neuritis, epididymitis and orchitis."

36. "The Smallpox Vaccine and Postvaccinal Encephalitis" "Before we become complacent with the idea that we will respond to a bioterrorism attack with a mass immunization program for smallpox, it is important to be reminded of the risk and clinical manifestations of postvaccinal encephalitis... The first case of postvaccinal encephalitis as a complication of the Jennerian cowpox inoculation was observed in 1905. A century later, there is no effective therapy." Karen L. Roos, et al, Semin Neurol 22: 095-098 -- 1/1/2002

37. "Very rarely, yellow fever vaccine-associated neurotropic disease (YEL-AND) has been reported following vaccination, with sequelae or with fatal outcomes in some cases. Clinical features have appeared within one month of vaccination and include high fever with headache that may progress to include one or more of the following: confusion, encephalitis/encephalopathy, meningitis, focal neurological deficits, or Guillain Barre syndrome." Package insert, Sanofi Pasteur, Manufacturer -- 1/1/1900


39. "Relapsing Neuropathy due to tetanus toxoid." "Summary: A unique case history is presented of a 42-year-old patient who has suffered three episodes of a demyelinating neuropathy, each of which followed an injection of tetanus toxoid." Pollard, JD; Selby, G, Journal of the Neurological Sciences, 1978, 37: 113-125 -- 1/1/1900

40. "Optic neuritis and myelitis following rubella vaccination" Kline L, Margulies SL, Arch Neurol 1982;39:443-4 -- 1/1/1900


53. "Murine model for pertussis vaccine encephalopathy: linkage to H−2" "Local, systemic and neurological complications have been observed following pertussis (whooping cough) vaccination in children1,2. These often occur soon after primary or secondary immunization. The neurological syndrome ranges from minor irritability to convulsions, coma, and on rare occasions death." L. Steinman, et al, Nature 299, 738 - 740 -- 10/21/1982

54. "Measles, measles vaccination, and risk of subacute sclerosing panencephalitis (SSPE)" "Occurrence of subacute sclerosing panencephalitis (SSPE) in some children who were vaccinated against measles could be explained by incomplete vaccine efficacy Measles, measles vaccination, and risk of subacute sclerosing panencephalitis (SSPE) " N Zilber, Neurology, Vol 33, Issue 12 1558-1564 -- 12/1/1983


56. "Myelin basic protein as an encephalitogen in encephalomyelitis and polynieuritis following rabies vaccination" "Encephalitis and polynieuritis occurring after rabies vaccination are believed to be immunologically mediated. We studied antibody responses to neural antigens in 36 patients with major neurologic complications, 25 with minor complications, and 39 with no complications after immunization with a brain-derived, Semple rabies vaccine." T Hemachudha, et al, New England Journal of Medicine Volume 316:369-374 , Number 7 -- 2/12/1987


59. "Acute cerebellar ataxia and facial palsy after DPT immunization" "Since the initial report of Beyers & Moll (1948), numerous cases of seizures and encephalopathy after pertussis immunization or DPT immunization have been reported. We report a 1-year-11-month-old girl with acute cerebellar ataxia and facial palsy after DPT immunization." Katafuchi Y, et al, No To Hattatsu. 21(5):465-9. -- 9/1/1989


62. "Neurological adverse events associated with vaccination" "These complications include autism (measles vaccine), multiple sclerosis (hepatitis B vaccine), meningoencephalitis (Japanese encephalitis vaccine), Guillain-Barre syndrome and giant cell arteritis (influenza vaccine), and reactions after exposure to animal rabies vaccine. Seizures and hypotonic/hypo-responsive episodes following pertussis vaccination and potential risks associated with varicella vaccination, as well as vaccine-associated paralytic poliomyelitis following oral poliovirus vaccination, are also described." Piyasirisilp, Sucheep a; Hemachudha, Thiravat b, Neurology. 15(3):333-338 -- 6/1/2002

63. "Development of case definitions for acute encephalopathy, encephalitis, and multiple sclerosis reports to the Vaccine Adverse Event Reporting System" "Acute encephalopathy age <18 months, encephalitis (EI), and multiple sclerosis (MS) after vaccination have been reported to VAERS" Robert Ball, et al, Journal of Clinical Epidemiology Volume 55, Issue 8, Pages 819-824 -- 8/1/2002

64. "Postvaccinal inflammatory neuropathy: peripheral nerve biopsy in 3 cases" "Autoimmune inflammatory polyneuropathy (PN) can be triggered by vaccination. We report 3 such cases. A 36-year-old female nurse presented 15 days after a hepatitis B vaccination (HBV) with acute sensory disturbances in the lower limbs. She had severe ataxia but no weakness." Claude Vital, et al, Journal of the Peripheral Nervous System Volume 7 Page 163 -- 9/1/2002


66. "Neurologic complications of immunization." "Individual vaccines can produce systemic or neurologic reactions ranging from minor events, such as pain and erythema at the injection site, to major complications, such as seizures, shock, encephalopathy, or death." Bale JF Jr, J Child Neurol.; 19(6): 405-12. -- 6/1/2004

Vaccines and Diabetes

1. "We found a large epidemic of diabetes. A 60% increase occurred in New Zealand following this immunization program." Dr. J. Bart Classen MD, former Researcher National Institute of Health, Classen Immunotherapies

2. Four children…developed diabetes mellitus shortly after active mumps vaccination. K. Helme, et al Diabetologia, Vol. 29, Number 1/30-33, 1/1/86

3. Hepatitis B vaccine and Haemophilus influenza type b vaccine have been respectively suspected to be responsible for neurological demyelinating disease and insulin dependent diabetes mellitus. Ovetchkine P., Arch Pediatr, 8(3):316-20 3/101

4. In 1992, 180 European doctors jointly noted that the mumps vaccine can trigger diabetes which only becomes apparent months after vaccination. Albionico et al, JAM 192:9(1)
5. A large ecological study in New Zealand revealed that an epidemic of diabetes followed a campaign to vaccinate children against hepatitis B. This report, published in the New Zealand Medical Journal in 1996 revealed that a 60 percent increase in childhood diabetes occurred in the years following the 1989-1991 vaccination program of children aged 6 to 16.

6. The widespread use of the Haemophilus meningitis vaccine has resulted in diabetes epidemics. Diabetes is an autoimmune disease that has been frequently observed to occur as a consequence of mumps vaccine (Fescharek et al., 1990; Helmke et al., 1986).

7. We believe the effects of vaccines on diabetes are of tremendous clinical importance and that trials need to be started immediately to address the effect of vaccines on diabetes and other autoimmune diseases. (Classen & Classen, 1996).


10. Encephalopathy after vaccination against smallpox with permanent sequel--diabetes insipidus

Vaccines and Arthritis


14. David A Geier, Mark R Geier MD PhD. *Chronic adverse reactions associated with hepatitis B vaccination* The Annals of Pharmacotherapy 2002: Vol. 36, No. 12, pp. 1970–1971. “In conclusion, our study demonstrates that adult HBV is statistically associated not only with acute neuropathy, neuritis, myelitis, vasculitis, thrombocytopenia, gastrointestinal disease, multiple sclerosis, and arthritis, but some of these patients go on to develop chronic adverse reactions that persist for at least 1 year following HBV. These types of chronic adverse reactions following adult HBV should be discussed with patients contemplating being immunized with HBV and should be included in the differential diagnosis of those who develop them following adult HBV.”

15. Geier DA, Geier MR. *A one year follow-up of chronic arthritis following rubella and hepatitis B vaccination based upon analysis of the Vaccine Adverse Events Reporting System (VAERS) database*. Clin Exp Rheumatol 2002 Nov-Dec;20(6):767-71. MedCon, Inc., Silver Spring, Maryland, USA. OBJECTIVES: This analysis examined the incidence rate of chronic arthritis adverse reactions reported following adult rubella and hepatitis B vaccinations. In this analysis, etiologic mechanisms for chronic arthritis following adult rubella and hepatitis B vaccines were also explored. CONCLUSION: This study revealed that adult rubella and adult hepatitis B vaccines were statistically associated with chronic arthritis which persisted for at least one year. The etiology for these adverse reactions may involve autoimmune mechanisms. Furthermore, potential biases in the reporting rates of adverse reactions to VAERS were not observed. PMID: 12508767 [PubMed - in process]


HBV vaccination may induce hypersensitivity and autoimmune reactions in susceptible individuals and healthy subjects. Hepatitis B virus vaccination may induce severe reactions requiring the use of a long term treatment (mean period of time of 32.5 ± 24.8 months) in healthy subjects and in patients who suffer from autoimmune diseases and from ankylosing spondylitis or reactive arthritis, even if a complete remission has been already obtained. Abstract: 1186 November 10, 1998 Poster Session D: Miscellaneous Rheumatic Diseases 12:30-2:00 pm, Hall B1/C


Conclusion: hepatitis B vaccination might be followed by various rheumatic conditions, and might trigger the onset of underlying inflammatory and/or auto-immune rheumatic diseases. However, a causal relation between hepatitis B vaccination and the observed rheumatic manifestations cannot be easily established. Further epidemiological works are needed to establish whether hepatitis B vaccination is associated or not with an incidence of rheumatic disorders higher than normal.


**Vaccines and Immunosuppression**

1. Michael Day notes that “vaccination-induced immunosuppression may on occasion be sufficient to permit the development of severe disease in animals that are carrying subclinical opportunistic pathogens” Day, M.J. 2006. *Vaccine side effects: Fact and fiction.* Veterinary Microbiology. 117, 51-58


