



Vaksinering og bivirkninger

Erling Olaf Koppang

Norges veterinærhøgskole

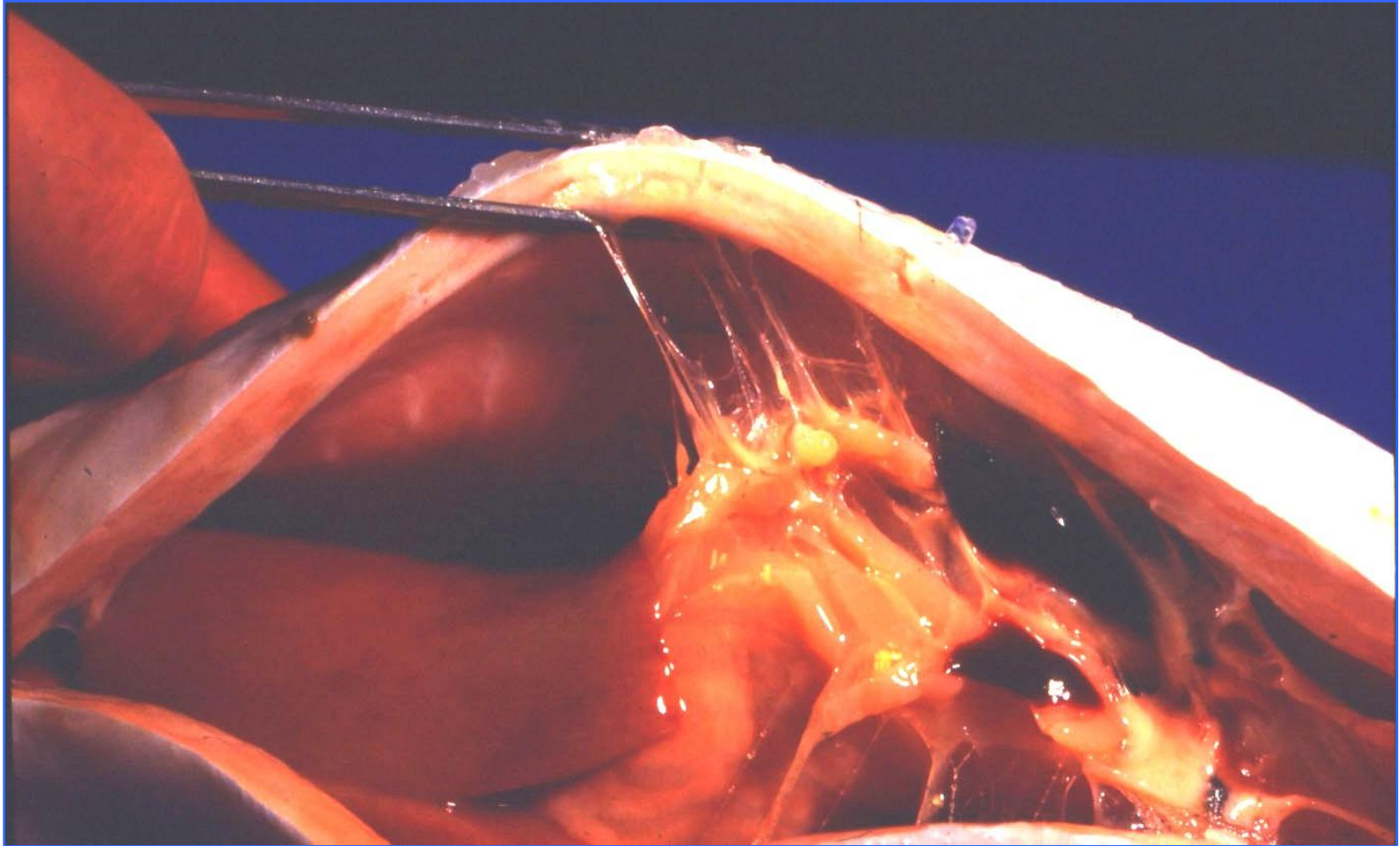
Oslo





0,1 ml / 40 g = 200 ml / 80 kg







Vaccine-associated granulomatous inflammation and melanin accumulation in Atlantic salmon, *Salmo salar* L., white muscle

E O Koppang¹, E Haugarvoll¹, I Hordvik², L Aune¹ and T T Poppe¹

¹ Department of Basic Sciences and Aquatic Medicine, Norwegian School of Veterinary Science, Oslo, Norway
² Department of Fisheries and Marine Biology, HIB, University of Bergen, Norway

Abstract

The purpose of this study was to investigate the nature of variably sized pigmented foci encountered in fillets of farmed Atlantic salmon, *Salmo salar* L. The material was sampled on the fillet production line and on salmon farms from fish with an average size of 3 kg from various producers. The fish had been routinely vaccinated by injection. Gross pathology, histology, immunohistochemistry using antisera against major histocompatibility complex (MHC) class II β chain and transmission electron microscopy (TEM) were used to characterize the changes. Macroscopically, melanized foci were seen penetrating from the peritoneum deep into the abdominal wall, sometimes right through to the skin, and also embedded in the caudal musculature. Histological investigation revealed muscle degeneration and necrosis, fibrosis and granulomatous inflammation containing varying numbers of melano-macrophages. Vacuoles, either empty or containing heterogeneous material, were frequently seen. The presence of abundant MHC class II⁺ cells indicated an active inflammatory condition. TEM showed large extracellular vacuoles and leucocytes containing homogeneous material of lipid-like appearance. The results showed that the melanized foci in Atlantic salmon fillet resulted from an inflammatory condition probably induced by vaccination. The described condition is not known in wild salmon and in farmed salmon where injection vaccination is not applied.

Correspondence: Trygve T. Poppe, Department of Basic Sciences and Aquatic Medicine, Norwegian School of Veterinary Science, Ullevåleiveten 72, Box #148 Cep., 00333 Oslo, Norway. (e-mail: trygve.poppe@vetinst.no)

Keywords: Atlantic salmon, inflammation, melano-macrophage, major histocompatibility complex class II, mineral oil, vaccine.

Introduction

Various pathological conditions may be associated with abnormal pigmentation in tissues and organs. Such pigments may either be of exogenous or endogenous origin. Endogenous pigments include derivatives of lipids, haemoglobin, porphyrins and melanin. The term melanosis is used to describe the presence of melanin in abnormal locations (Thomson 1984). In vertebrates, melanin is synthesized by melanocytes and organized in melanosomes, which are lysosome-related intracellular organelles (Orlov 1995; Raposo, Fevrier, Stoorvogel & Marks 2002). Mammalian melanocytes originate from the embryonic neural tube (Salmón & Kitchell 2003) and it has been observed that such cells can migrate into inflamed tissue (Thomson 1984).

Inflammatory reactions and tissue regeneration in salmonids seem similar to those of mammals (Finn & Nielson 1971), but have in addition been associated with the involvement of so-called melano-macrophages (Roberts 1975; Agius & Roberts 2003). The origin of melanosomes in melanin-containing viscera located cells in fish is not clear (Agius & Roberts 2003), but Siegel, Scalia, Mondio & Corsaro (1997) suggested that melanogenesis in poikilothermic vertebrates may occur in monocyteme-derived cells of the haematopoietic lineage. Although teleost melano-macrophages have been ascribed macrophage-like properties, their functions and significance are

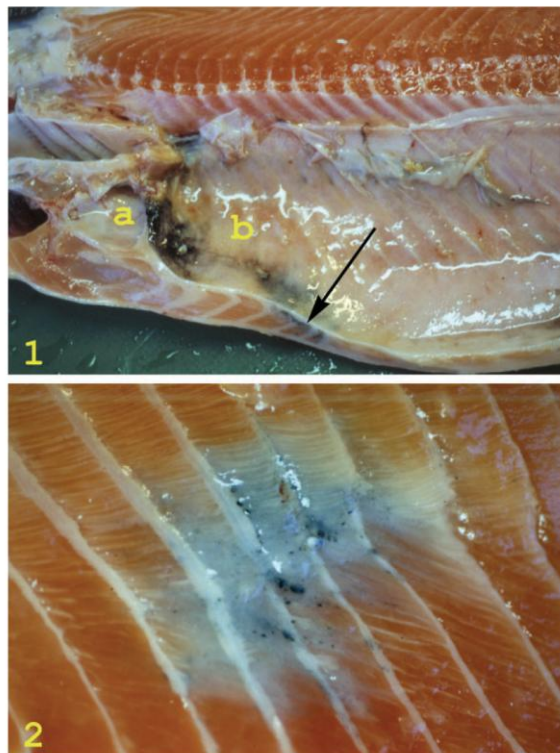


Figure 1 Gross pathological changes in the carcass of an Atlantic salmon. The pericardial cavity (a) is normal, but severe melanization is apparent in the abdominal cavity (b). Melanized musculature subjacent to the peritoneum is seen on the cut surface (arrow).

Figure 2 A melanized area in the musculature of an Atlantic salmon. The peritoneum is removed and darker foci are seen in a dark to grey area involving five myosepta. The lesion is situated laterally in the fish, covering the area of the lateral organ. Note the contraction in the musculature, disrupting the curves of the intramuscular septa.

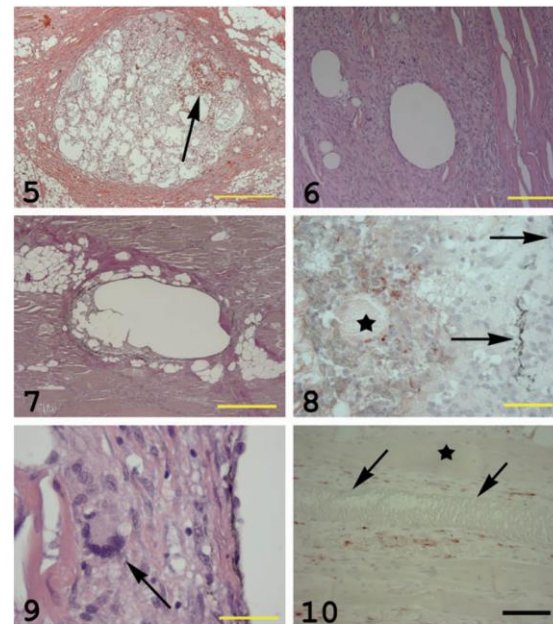


Figure 5 A large vesicle embedded in an intermyotomal septum containing macrophage-like cells, debris and a fresh haemorrhage (arrow) (H&E, bar = 500 μ m).

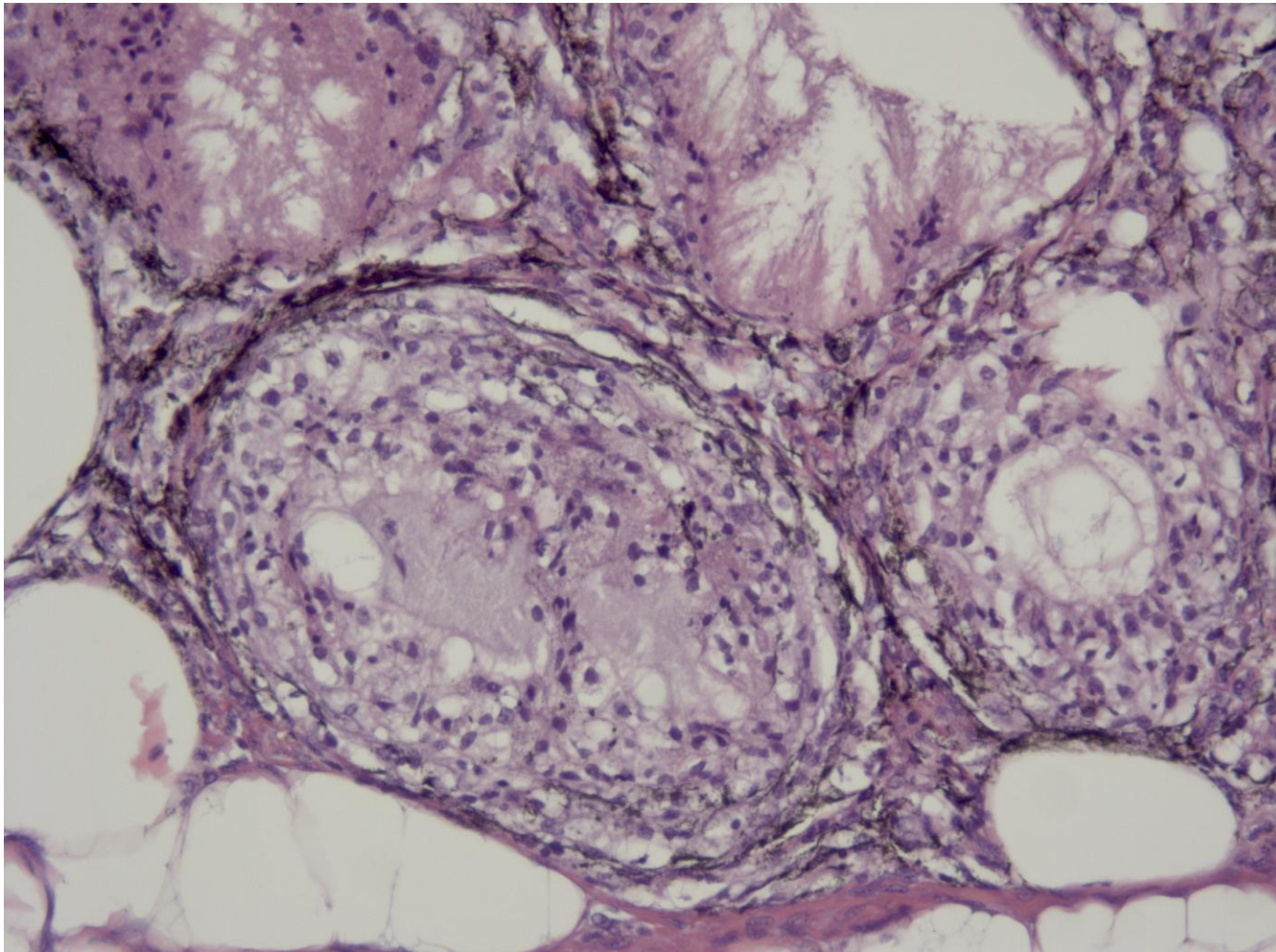
Figure 6 Empty vesicles surrounded by granulomatous tissue embedded in the white musculature. Note adjacent, seemingly unaffected muscle cells (H&E, bar = 200 μ m).

Figure 7 Vesicles embedded in the white musculature surrounded by fibrogranulomatous tissue (red staining) (EVG, bar = 500 μ m).

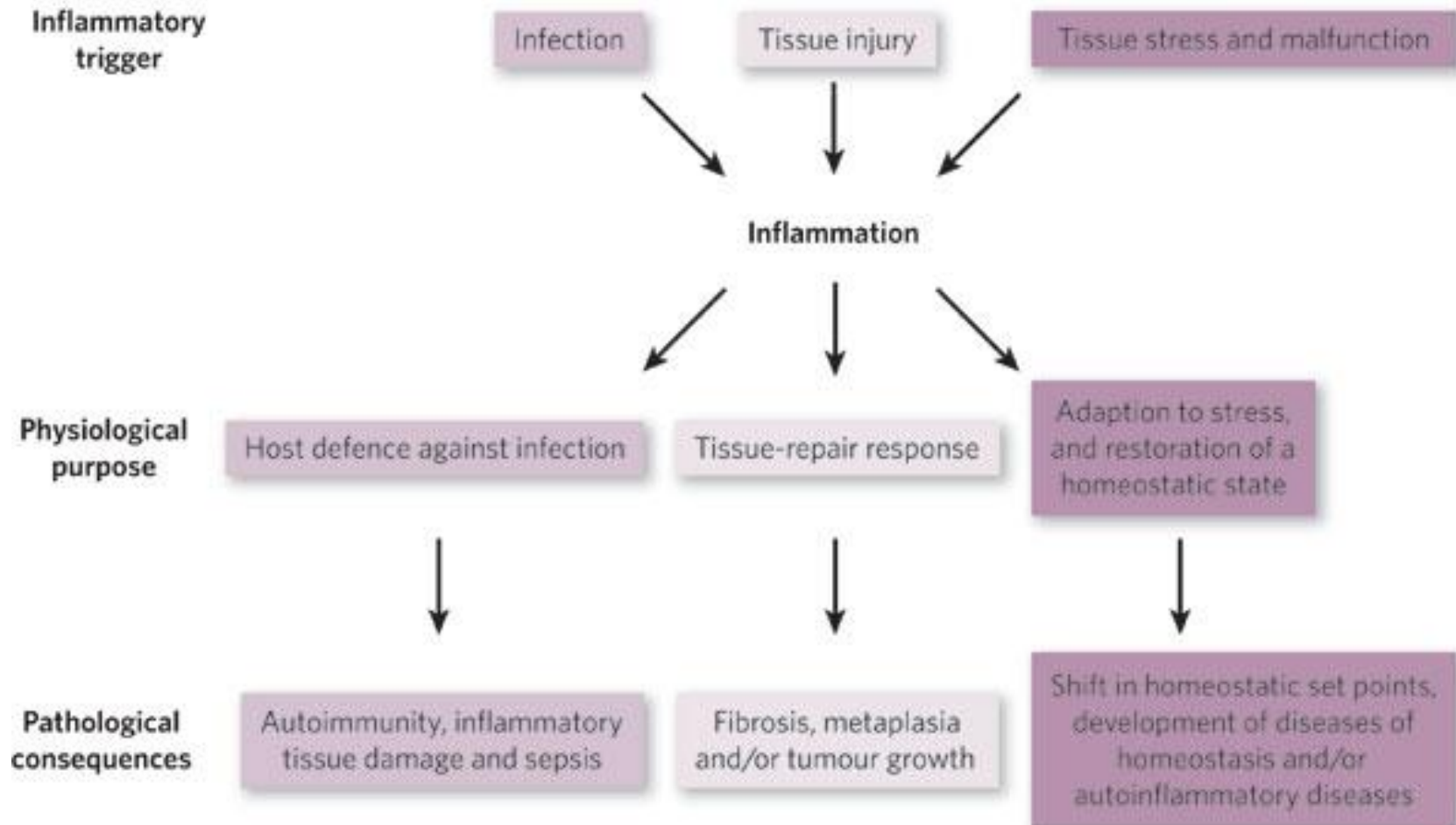
Figure 8 Reaction against oil (red staining) in a vesicle as shown in Fig. 5. Homogeneous masses (asterisk) and macrophage-like cells show positive reactions. Note the melano-macrophages in the vesicle wall (arrows) (oil red O, bar = 50 μ m).

Figure 9 High magnification of the wall of a vesicle as seen in Fig. 6. The wall contains a multinucleated giant cell (MGC) (arrow), epithelioid-like cells, small vacuoles and is lined towards the lumen of the greater vesicle with melanosome-containing cells, probably swollen melano-macrophages (H&E, bar = 40 μ m).

Figure 10 Muscle cells infiltrated with MHC class II⁺ cells. One muscle cell is unaffected (asterisk). One fibre shows severe degeneration (arrowhead), whereas one is invaded by MHC class II⁺ cells (red reaction) (MHC class II immunostain, haematoxylin counterstain, bar = 100 μ m).



Kronisk manifestasjon



Induction of Lupus-associated Autoantibodies in BALB/c Mice by Intraperitoneal Injection of Pristane

By Minoru Satoh and Westley H. Reeves

From the Departments of Medicine and Microbiology/Immunology, Thurston Arthritis Research Center and University of North Carolina Lineberger Comprehensive Cancer Center, University of North Carolina, Chapel Hill, North Carolina 27599-7280

Summary

Intraperitoneal injection of pristane (2,6,10,14 tetramethylpentadecane) is a standard technique for obtaining monoclonal antibody-enriched ascitic fluid. However, pristane also induces plasmacytomas and an erosive arthritis resembling rheumatoid arthritis in BALB/c mice, probably as a consequence of enhanced interleukin 6 production. We report here that the production of autoantibodies characteristic of systemic lupus erythematosus (SLE) is a further consequence of injecting pristane in BALB/c mice. Anti-Su antibodies appeared as early as 1–2 mo after a single injection of 0.5 ml pristane, followed by anti-U1RNP and anti-Sm antibodies after 2–4 mo. Within 6 mo of pristane injection, 9 of 11 BALB/c mice had developed anti-Su, anti-U1RNP, anti-U2RNP, anti-Sm, and possibly anti-U5RNP antibodies. Autoantibodies were not produced by 20 BALB/c mice of the same age and sex that were not injected with pristane. Thus, autoantibodies characteristic of lupus were induced in mice that are not usually considered to be genetically susceptible to the disease. The induction of autoantibodies associated with SLE by pristane may be relevant to understanding the role of abnormal cytokine production in autoantibody production and the pathogenesis of autoimmune disease. Furthermore, the induction of high titer autoantibodies by pristane dictates caution in the use of ascitic fluid as a source of monoclonal antibodies, since the polyclonal autoantibodies induced by pristane may copurify with the monoclonal antibody secreted by an injected hybridoma.

Intraperitoneal administration of pristane (2,6,10,14 tetramethylpentadecane) before the injection of hybridoma cells is a standard technique for obtaining ascitic fluid containing a high concentration of mAbs. In addition to its effects on hybridoma cell growth, pristane-induced alterations in cytokine production have been implicated in the pathogenesis of plasmacytomas (1–3) and erosive arthritis resembling rheumatoid arthritis (4, 5). While characterizing a slowly growing murine hybridoma secreting an IgM mAb, we observed that ascitic fluid from several pristane-primed BALB/c mice injected with hybridoma cells contained polyclonal IgG autoantibodies to Su, U1RNP, U2RNP, and/or Sm. Further investigation revealed that the autoantibodies were a consequence of pristane priming itself, and were unrelated to the hybridoma cells or their secreted monoclonal IgM. Thus, intraperitoneal injection of pristane induced lupus-like autoimmunity in a strain of mouse not usually thought to be prone to autoimmune disease.

Materials and Methods

Cell Lines. The K562 (human erythroleukemia) and L929 (murine fibroblast) cell lines were obtained from the American Type Culture Collection (ATCC; Rockville, MD) and maintained in

RPMI 1640 or MEM, respectively, supplemented with 9% FCS, L-glutamine, and penicillin/streptomycin.

Sera and mAbs. Prototype human autoimmune sera containing anti-Su, anti-U1RNP, anti-Sm, or other specificities, were reported previously (6–8). Additional sera with anti-U1RNP/Sm antibodies were obtained from patients with systemic lupus erythematosus (SLE) followed at the University of North Carolina Hospitals (Chapel Hill, NC) or the Keio University Hospital (Tokyo, Japan). Murine mAbs 2.73 (anti-U1-70K) (9), and 9A9 (anti-U1-A and U2-B⁺) (10) were provided by Dr. Yoshihiko Takeda (Medical College of Georgia, Augusta, GA) and Dr. W.J. van Venrooij (University of Nijmegen, The Netherlands), respectively. mAbs Y2 (anti-Sm B/B and D) (11), 22G12 (anti-Sm B/B) (12), and 2G7 (anti-Sm-D) (13) were provided by Dr. Robert A. Eisenberg (University of North Carolina).

Pristane Priming. 6–8-wk-old female BALB/c ByJ mice were obtained from The Jackson Laboratory (Bar Harbor, ME) and maintained at our animal facility. Eleven mice, ages 4–5 mo, received a single intraperitoneal injection of 0.5 ml of pristane (Sigma Chemical Co., St. Louis, MO). Sera were collected every 4 wk from the tail vein. 20 age- and sex-matched BALB/c ByJ mice that were not injected with pristane served as controls.

Immunoprecipitation. Immunoprecipitation using cell extract from K562 or L929 cells was performed as described previously (7, 8). Briefly, the cells were labeled for 14 h with [³⁵S]methionine/cysteine (25 μ Ci/ml), lysed in 0.5 M NaCl NET/NP-40 buffer

Proc. Natl. Acad. Sci. USA
Vol. 92, pp. 10934–10938, November 1995
Immunology

Anti-nuclear antibody production and immune-complex glomerulonephritis in BALB/c mice treated with pristane

(systemic lupus erythematosus/lupus nephritis/autoantibodies/autoimmunity/small nuclear ribonucleoproteins)

MINORU SATOH*, ANIL KUMAR†, YASHPAL S. KANWAR†, AND WESTLEY H. REEVES*‡

*Departments of Medicine and Microbiology/Immunology, Thurston Arthritis Research Center and University of North Carolina Lineberger Comprehensive Cancer Center, University of North Carolina, Chapel Hill, NC 27599-7280; and †Department of Pathology, Northwestern University Medical School, Chicago, IL 60611

Communicated by Maclyn McCarty, The Rockefeller University, New York, NY, August 10, 1995

ABSTRACT The pathogenesis of systemic lupus erythematosus is thought to be primarily under genetic control, with environmental factors playing a secondary role. However, it has been shown recently that intraperitoneal injection of pristane (2,6,10,14-tetramethylpentadecane) induces autoantibodies typical of lupus in BALB/c mice, a strain not usually considered to be genetically susceptible to the disease. In this study, the induction of autoimmune disease by pristane was investigated. BALB/c mice receiving pristane were tested for autoantibody production and histopathological evidence of glomerulonephritis. Six of 11 mice developed IgM anti-single-stranded DNA antibodies shortly after receiving pristane and 4 developed IgM anti-histone antibodies, but anti-double-stranded DNA antibodies were absent. IgG anti-DNA and anti-histone antibodies were absent. In contrast, the lupus-associated anti-nuclear ribonucleoprotein/Sm and anti-Su autoantibodies produced by these mice were predominantly IgG. In addition to autoantibodies, most of the mice developed significant proteinuria. Light microscopy of the kidney showed segmental or diffuse proliferative glomerulonephritis. Electron microscopy showed subepithelial and mesangial immune-complex deposits and epithelial foot process effacement. Immunofluorescence revealed striking glomerular deposition of IgM, IgG, and C3 with a mesangial or mesangiocapillary distribution. Thus, pristane induces immune-complex glomerulonephritis in association with autoantibodies typical of lupus in BALB/c mice. These data support the idea that lupus is produced by an interplay of genetic and environmental factors and that unlike the MRL or (NZB \times WJ/F) mouse models, in which genetic susceptibility factors are of primary importance, environmental factors are of considerable importance in the autoimmune disease of pristane-treated BALB/c mice.

Systemic lupus erythematosus (SLE) is a multisystem autoimmune disease characterized by anti-nuclear antibodies, immune-complex glomerulonephritis, arthritis, and other manifestations. Anti-double-stranded (ds) DNA autoantibodies are highly specific for SLE and may play a key role in the pathogenesis of immune-complex nephritis in lupus (1, 2). However, autoantibodies to glomerular antigens (3) and/or dysregulated cytokine production (4, 5) may also be involved. Human SLE is influenced strongly by major histocompatibility complex-linked and -nonlinked genes (6–8). Multiple genetic loci that accelerate the onset of autoantibody production and/or nephritis also have been identified in murine lupus models (9, 10). The importance of environmental factors in the pathogenesis of lupus is less clear. However, the role of environmental exposures in autoantibody production is underscored by the recent demonstration that intraperitoneal

(i.p.) injection of pristane (2,6,10,14-tetramethylpentadecane) induces autoantibodies characteristic of SLE, including anti-Su and anti-nuclear ribonucleoprotein (nRNP)/Sm, in BALB/c mice, a strain not usually considered to be predisposed to autoimmunity (11). Titers of these autoantibodies are comparable to those found in MRL/lpr mice (12). The present data show that in addition to IgG anti-Su and anti-nRNP/Sm autoantibodies, pristane induces IgM anti-single-stranded (ss) DNA, anti-histone antibodies, and immune-complex glomerulonephritis in the “nonautoimmune” BALB/c strain.

MATERIALS AND METHODS

Administration of Pristane. Eleven 4- to 5-month-old and 10 2.5-month-old female BALB/c ByJ mice (The Jackson Laboratory) received a single i.p. injection of 0.5 ml of pristane (Sigma) (11). Sera were obtained at 1, 2, and 4 weeks and monthly thereafter. Urine samples were tested monthly for protein concentration by using Albutix reagent strips (Miles).

ELISAs for Anti-nRNP/Sm, Su, ssDNA, and Histone Autoantibodies. Anti-Su and anti-nRNP/Sm antigen-capture ELISAs were performed as described (12) with 1:250 diluted murine serum and alkaline phosphatase-conjugated goat anti-mouse IgG or IgM antibodies. Antibodies to heat-denatured calf thymus DNA (ssDNA, from Sigma) and to total calf thymus histones (United States Biochemical) were detected by ELISAs as described (13, 14) with a 1:500 dilution of murine sera and alkaline phosphatase-conjugated goat anti-mouse IgG or IgM antibodies.

Light and Electron Microscopy. Six months after receiving pristane, BALB/c and control mice not receiving pristane were anesthetized and fixed by perfusion through the left ventricle (15). The inferior vena cava was nicked below the renal veins, and 20 ml of saline was perfused slowly, followed by 10 ml of 2.5% (vol/vol) glutaraldehyde in 0.1 M sodium cacodylate, pH 7.4/4 mM CaCl₂. For light microscopy, 3- μ m sections of aldehyde-fixed renal cortex were stained with hematoxylin and eosin as described (16). For electron microscopy, aldehyde-fixed renal tissue was postfixated with osmium tetroxide, dehydrated in ethanol, and embedded in Epon 812. Thin sections (60 nm) were stained with lead citrate and uranyl acetate and examined by electron microscopy (16).

Immunofluorescence. Kidneys were excised from pristane-primed or control mice and snap-frozen in isopentane chilled in liquid N₂. Cryostat sections (4 μ m) were stained with a 1:40 dilution of fluorescein isothiocyanate (FITC) or rhodamine-conjugated goat anti-mouse IgM, IgG, IgG1, IgG2a, IgG2b, or IgG3 antibodies (Southern Biotechnology Associates) or with FITC-conjugated rabbit anti-mouse C3 antiserum (Organon

Abbreviations: nRNP, nuclear ribonucleoprotein; SLE, systemic lupus erythematosus; ds, double stranded; ss, single stranded; IL, interleukin.

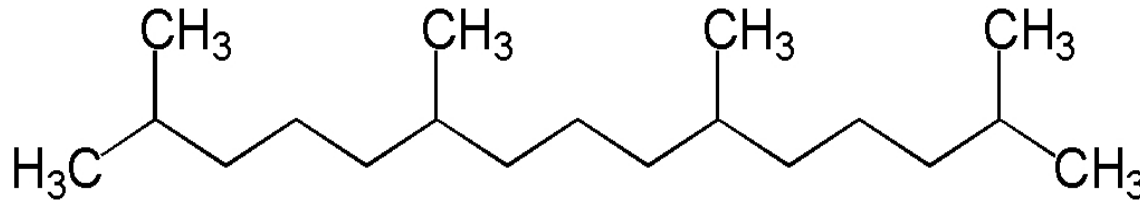
‡To whom reprint requests should be addressed.



Pristane

(2,6,10,14-tetramethylpentadecane, TMPD)

- C19 isoprenoid alkane (C₁₉H₄₀)
- A component of mineral oil



- A single intraperitoneal injection (ip) induces plasmacytoma and chronic destructive arthritis in mice
- **Induction of lupus-like autoimmune syndrome**
 - Antinuclear antibodies, anti-Sm/U1RNP, ribosomal P, dsDNA, immune complex glomerulonephritis
 - (Satoh M and Reeves WH, J Exp Med 1994, Satoh M et al., Proc Natl Acad Sci USA 1995)



Aktuell publisasjon

Vaksinering av oppdrettsfisk – sjukdomsvern med attåttsmak

Vaksinering av oppdrettsfisk er viktig for å unngå tapstringende infeksjonssjukdomer. Utvikling av effektive vaksiner og vaksineringsstrategier har redusert antibiotika-bruken i oppdrettsnæringen i Norge fra et uakseptabelt høyt nivå for 15 år siden til et i dag som er klart høyere enn i liknende animalske produksjoner. Samtundes har produksjonen av oppdrettsfisk mange doble seg. De vanligste nyttåttet vaksinasjoner er formulert som vann-i-olje, eller såkalla "incomplete Freund's adjuvans". Ulike bakterie- og virusantigen vert sette til formuleringa. Vaksinen vert injisert intraperitonealt i måndagane før eller under smoltfiseringa. Her gjev han ein depoteffekt som initerar og oppretheld ein langvarig immunitet gjennom resten av produksjonsytusen.

ORIGINALREFERANSER:

Koppang ED, Bjerkås E, Bjerkås I, Sveier H, Hordvik I. Vaccination induces major histocompatibility complex II expression in the Atlantic salmon eye. *Scand J Immunol* 2003; 58 (1): 9-14.

Koppang ED, Haugravoll E, Hordvik I, Pappe TT, Bjerkås I. Granulomatous uveitis associated with vaccination in the Atlantic salmon. *Vet Pathol* 2004; 41 (2): 123-30.

Koppang ED, Haugravoll E, Hordvik I, Aune L, Pappe TT. Vaccine-associated granulomatous inflammation and melanin accumulation in Atlantic salmon, *Salmo salar* L., white muscle. *J Fish Dis* 2005; 28 (1): 13-22.

Tilhøve kring vaksiner i bruk

Vaksinerne sine immuniserte eigenskapar følgjer av lantemerkingsreguleringane til alle vaksiner og samvandringsvaks og bakterieantigen, som skal gje ei spesifikk verne.

Det er publisert ei rekkje artiklar som tek for seg oppsett og distribusjon av vaksiner i oppdrettsfisk. Dei er særleg vakerne til lipopolysakkarid (LPS) og A-lag fra *Aeromonas salmonicida* subspecies *salmonicida* som har vorte undersøkt. Resultata viser at i tillegg til å persistera ved infeksjonslokus, vert antigen i hovedsak distribuert til leverdysje (betreidskitt permanente proseduro) og milt. Ikkje rekner med at aar-gaav vert presentert og presentert for T-celler i disse lymfatiske organ, slik at ei spesifikk immunrespons kan inntreffe.

Når dei gjeld oppsett og distribusjon av adjuvantkomponentar i vaksinen, er publiserte undersøkingar frå fisk fjerntvernde. Ofte, som vert nytta er i stor utsegn ei blanding kvite mineralolje, som er lipofila til perleprotein (1). Dei er sammensetne av metta hydrokarbonkjeder, arena-

retto- og grønta kjeder eller ringstrukturar, av varierende lengder som er severt essensielle med bakke fisk neddyking. Den relative konsentrasjonen av metta hydrokarbon- (mette- og grønta kjeder) angjer også sine biologiske eigenskapar. Korte hydrokarbon-, med ei kjedelengdeblanding frå 15 til 25 karbonatom, har vist seg å vere mein potente ein-lange hydrokarbon, med ei kjedelengdeblanding frå 25 til 50 karbonatom, til å inkludere betennelsesresponsar og verke som adjuvans. For å få nære informasjon om distribusjonen og metabolismen til vaksiner, er det viktig å konsultere literatur publisert etter undersøkingar på naturlige dyr. I forsking på rotter vert retinomer hydrokarbon injisert og distribusjonen undersøkt. Sidan hydrokarbon er fettløslige, vert dei ikkje verna til dei over tid, vert ikkje absorbert og akkumulert i lever og fettvev (2). Neddykinga var svært langsom. For fisk-oppdrett er det publisert informasjonar om distribusjonen etter vaksineringsart analysert i stor utsegn av gasskromatografi-massaspektrometri. Her vart det sett at dyra kvita seg med hydrokarbon gjennom egg. Dessom høve ikkje verpa, var dei

Presisering om vaksinering av oppdrettsfisk

I NVT nr. 7, 2005 har professor og fagleg medredaktør i NVT Øystein Evensen ein innlegg under emnet Debat, med tittelen "Vaksinering av oppdrettsfisk", der han kjemner ein artikkel vår "Vaksineringsstrategier for oppdrettsfisk – sjukdomsvern med attåttsmak" i NVT nr. 4, 2005. Me takkar for interessa, men seer oss neyt til å oppklare eit par misforståingar.

Me vil fyrst det fyrste påpeike at hovudpoenget vårt i innlegget i NVT nr. 4 er at tilsetningsstoffet (1), (2) til bruk i adjuvans bør undersøkast for om dei kan distribuert systemisk i animalske laks etter intraperitoneal injisering.

Dette spørsmålet er grunn for mange studiar på pattedyr der det er funne patologiske forandringar i ulike vev etter eksponering for ulike hydrokarbon (3). Studiar frå desse eksperimenta vert referert då det er manglande kunnskap på dette området innan fisk, noke Evensen også stadfestar i innlegget sitt i NVT nr. 7, 2005. Det er også kjent frå dyr og menneske at absorberne hydrokarbon primært vert distribuert via lenkelsystemet til lymphknotar og lever, sekundært til fettvev (4).

Undersøkingane våre publisert i *Journal of Fish Diseases (JFD)* og utgitt i NVT nr. 4, varte patologiske funn som kan gje tilslutning på at den primære primære distribusjonen også vert tilfelt til fisk. Men me presiserar i JFD-publikasjonen at kjemiske studiar må utførast for å vere sikre på dette.

Sidan dei publiserte funna kan gje tilslutning og mistanke om at adjuvansen kan vere distribuert i kroppen også på fisk, må det vurderast å utføre farmakokinetiske studiar med omsyn på adjuvansen (olje eller i kombinasjon med antigen) (5).

Poenget er såleis at me ikkje byggja på basert innlegget vårt i NVT nr. 4, 2005 på funn som er publisert i JFD slik Evensen hevda i kommentarane sine. Dette er det også gjalt greie for med si rekkje referansar til andre publikasjonar.

Referansar

1. Hov K. Adjuvans i vaksiner til dyr. *Norsk Veterinærtidsskrift* 1956; 108: 227-33.
2. Eureka Y, Nicholas DG, Anzogi J, Reeves WH, Satch M. Autoimmunity induced by adjuvant hydrocarbon oil components of vaccine. *Bioact Pharmaceut* 2004; 55: 325-37.



Etter Haugravoll og Erlend Olaf Koppang er innlegg de våre om vaksinering av oppdrettsfisk.

3. Haugravoll E, Koppang EO. Vaksinering av oppdrettsfisk – sjukdomsvern med attåttsmak. *Norsk Veterinærtidsskrift* 2005; 117: 286-90.
4. OIEA/FAO/WP/95/FINAL. Committee for veterinary medicinal products mineral hydrocarbons summary report. <http://www.emea.eu.int/pdfs/vet/cdfs/009595enp.pdf>, 1995.
5. EMA/FAO/WP/VEG/17/02/2004/Consultation. Guideline on adjuvants in vaccines. <http://www.emea.eu.int/pdfs/human/vwp/01702cu.pdf>, 2004.

Erlend Haugravoll
Institutt for basalt og akvatisk medisin
Norges veterinærhøgskole

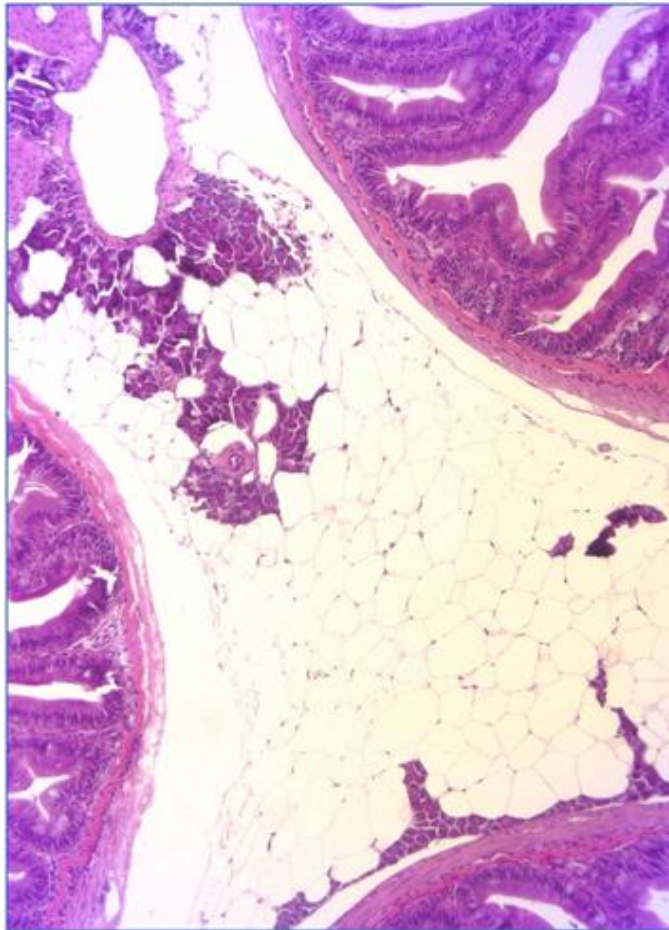
Erling Olaf Koppang
Institutt for basalt og akvatisk medisin
Norges veterinærhøgskole



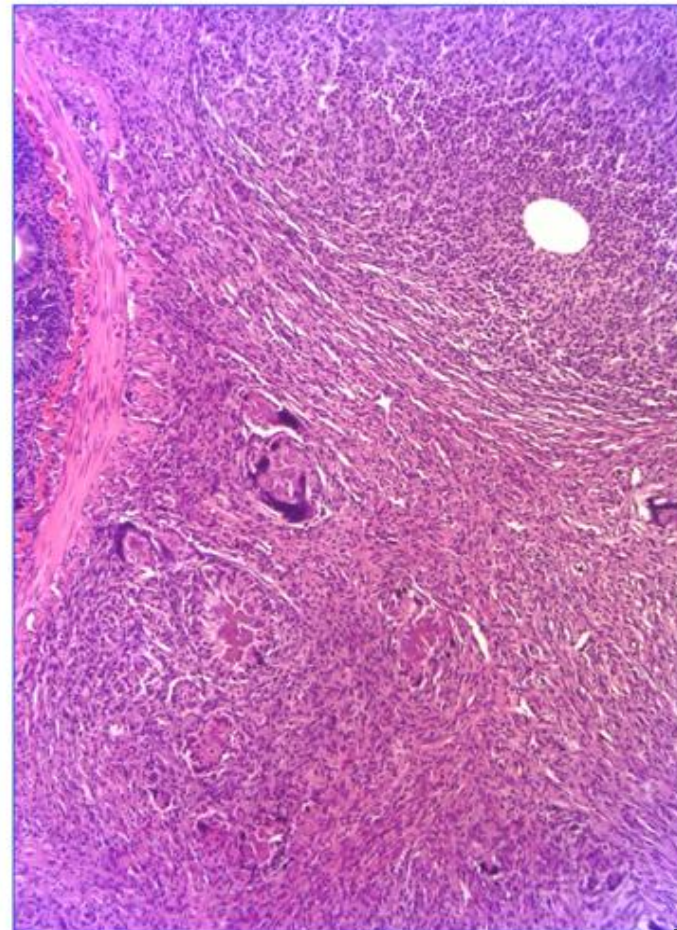


Peritoneal Tissue from Unvaccinated vs. Vaccinated Salmon

Unvaccinated



Vaccinated





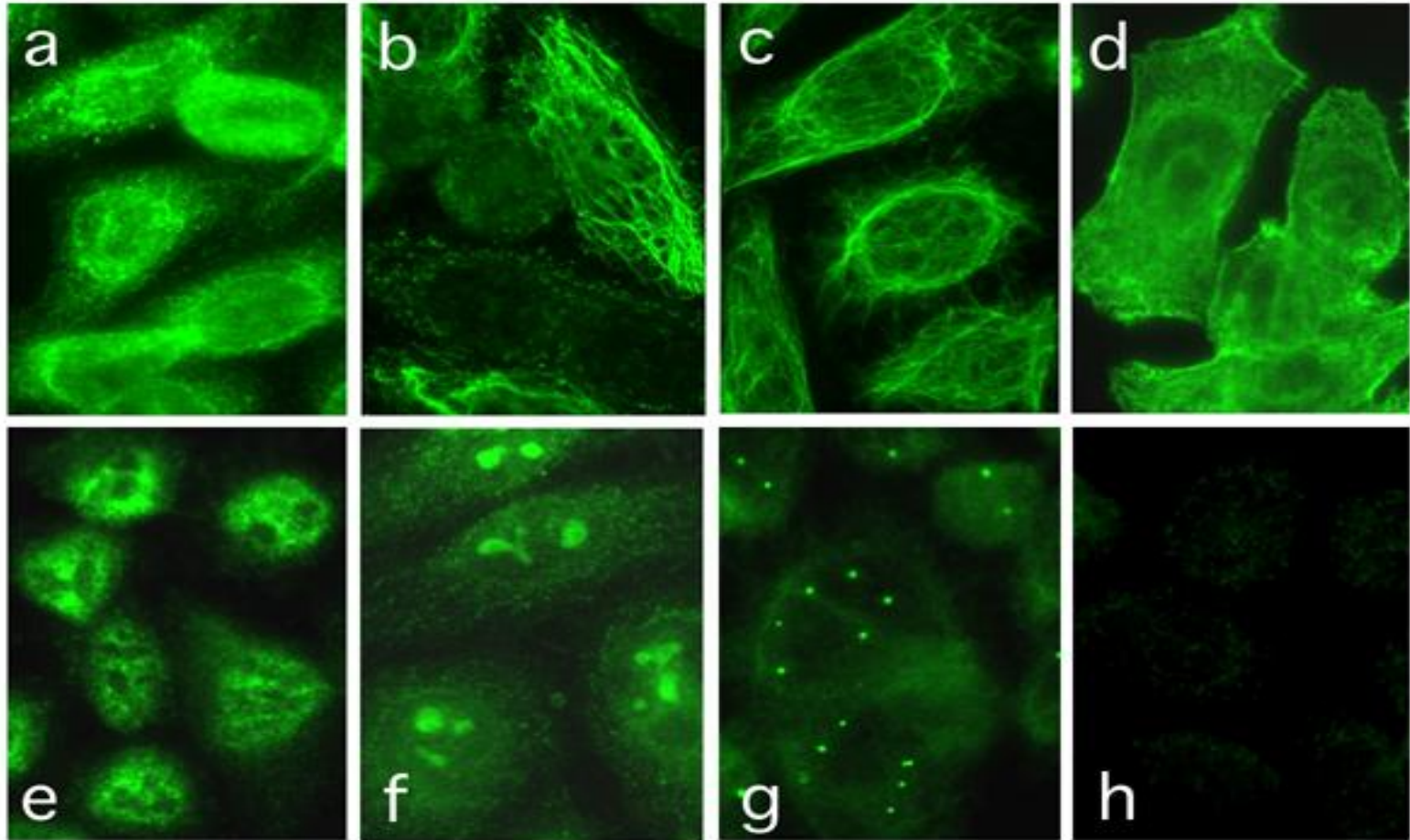


Background of Groups of Salmon

	n		Vaccine	Adjuvant	Vaccination - sample
A	10	Experimental	No	No	NA
B	10	Experimental	Yes	Mineral oil	2 y
C	20	Farmed (cave)	No	No	NA
D	55	Farmed	Yes	Mineral oil	1.5 y
E	20	Farmed	Yes	Mineral oil	2 y
F	19	Farmed	Yes	Mineral oil	2.3 y
G	20	Farmed	Yes	Mineral oil	1.3 y
H	20	Farmed	Yes	Animal/veg oil	1.3 y
I	20	Farmed	Yes	Mineral oil	1.5 y
J	11	Wild	No	No	NA

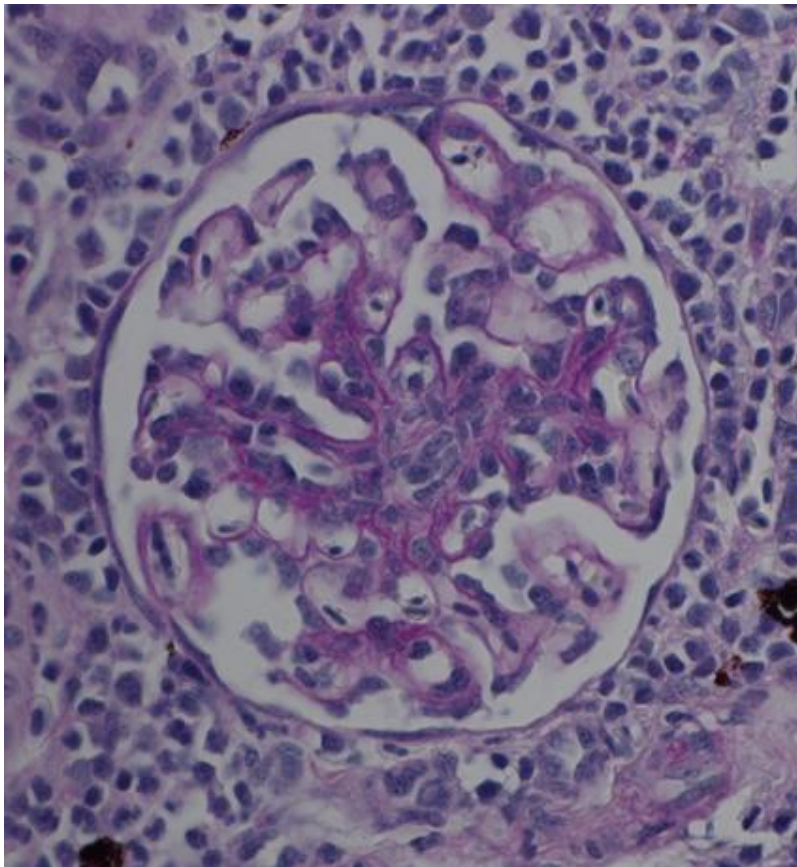


Autoantibodies in Vaccinated Salmon by Immunofluorescence

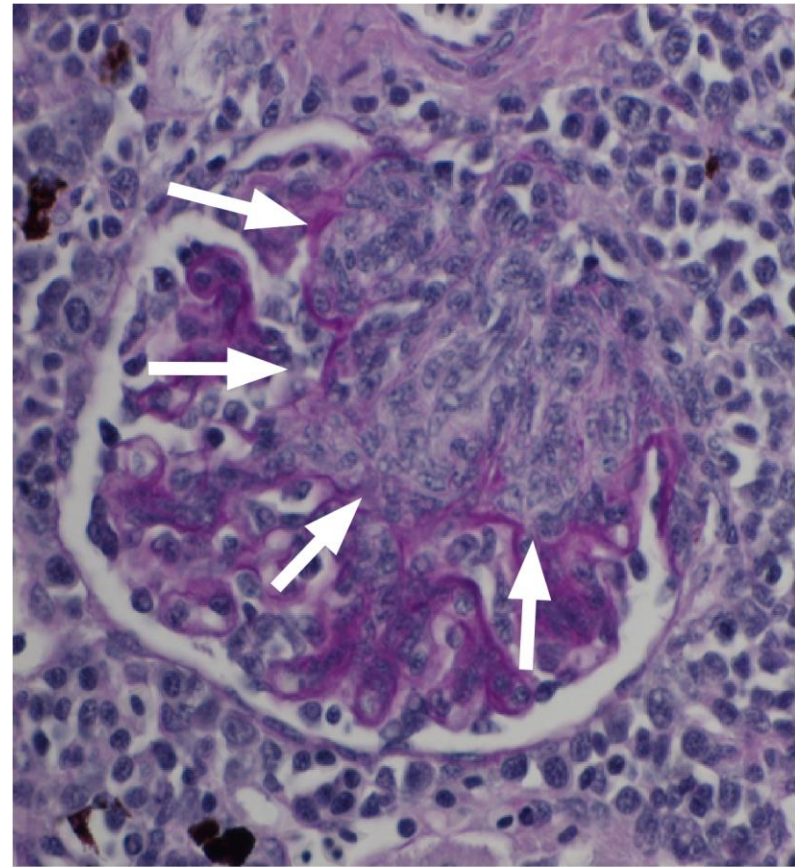




Focal Endocapillary Proliferation in Glomeruli from Vaccinated Salmon (PAS)



Unvaccinated



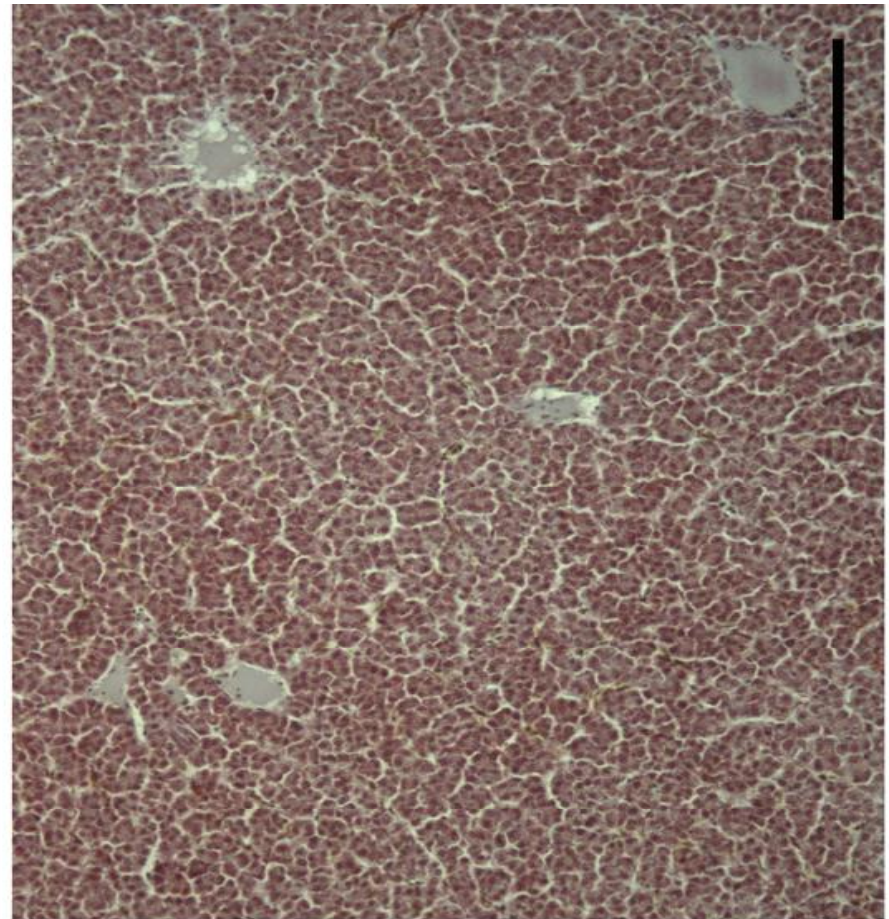
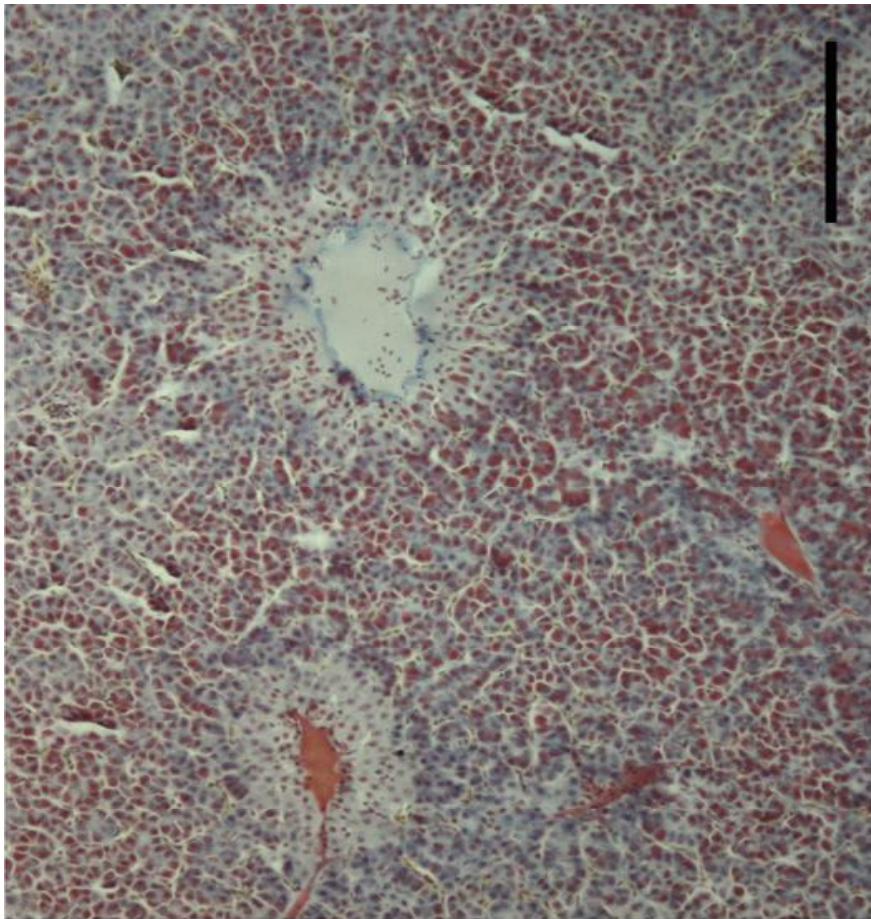
Vaccinated



Levertromber(12 måneder)

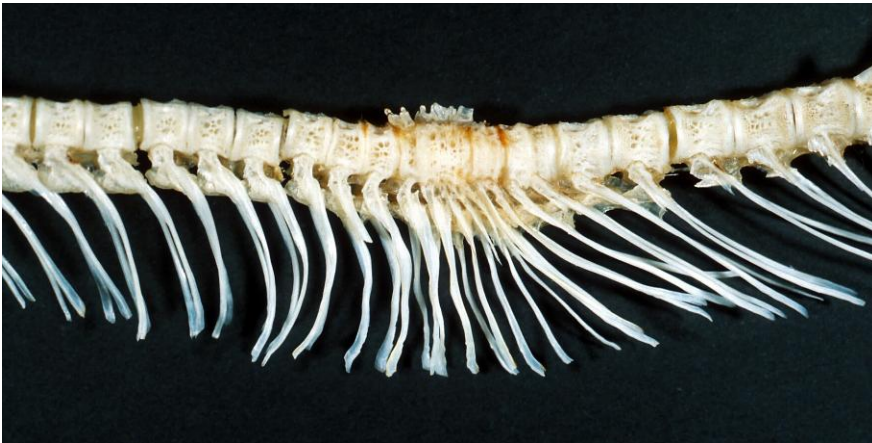
Vaksinert

Uvaksinert



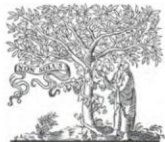
MSB staining (fibrin = red)

Mus 7 måneder etter pristan- injeksjon



Spine deformity in Atlantic salmon





ELSEVIER

Contents lists available at ScienceDirect

Vaccine

journal homepage: www.elsevier.com/locate/vaccine

Manifestations of systemic autoimmunity in vaccinated salmon

Erlend Haugarvoll^{a,1}, Inge Bjerkås^a, Nancy J. Szabo^d, Minoru Satoh^{b,c}, Erling O. Koppang^{a,*}

^a Section of Anatomy and Pathology, Department of Basic Sciences and Aquatic Medicine, Norwegian School of Veterinary Science, Ullevålsveien 72, PO Box 8146 Dep., 0033 Oslo, Norway

^b Division of Rheumatology and Clinical Immunology, Department of Medicine, University of Florida, Gainesville, FL 32610-0221, USA

^c Department of Pathology, Immunology, and Laboratory Medicine, University of Florida, Gainesville, FL 32610-0221, USA

^d Analytical Toxicology Core Laboratory, Department of Physiological Science, University of Florida, Gainesville, FL 32610-0221, USA

ARTICLE INFO

Article history:

Received 19 March 2010

Received in revised form 11 May 2010

Accepted 12 May 2010

Available online 27 May 2010

Keywords:

Systemic autoimmunity

Vaccination

Salmo salar

ABSTRACT

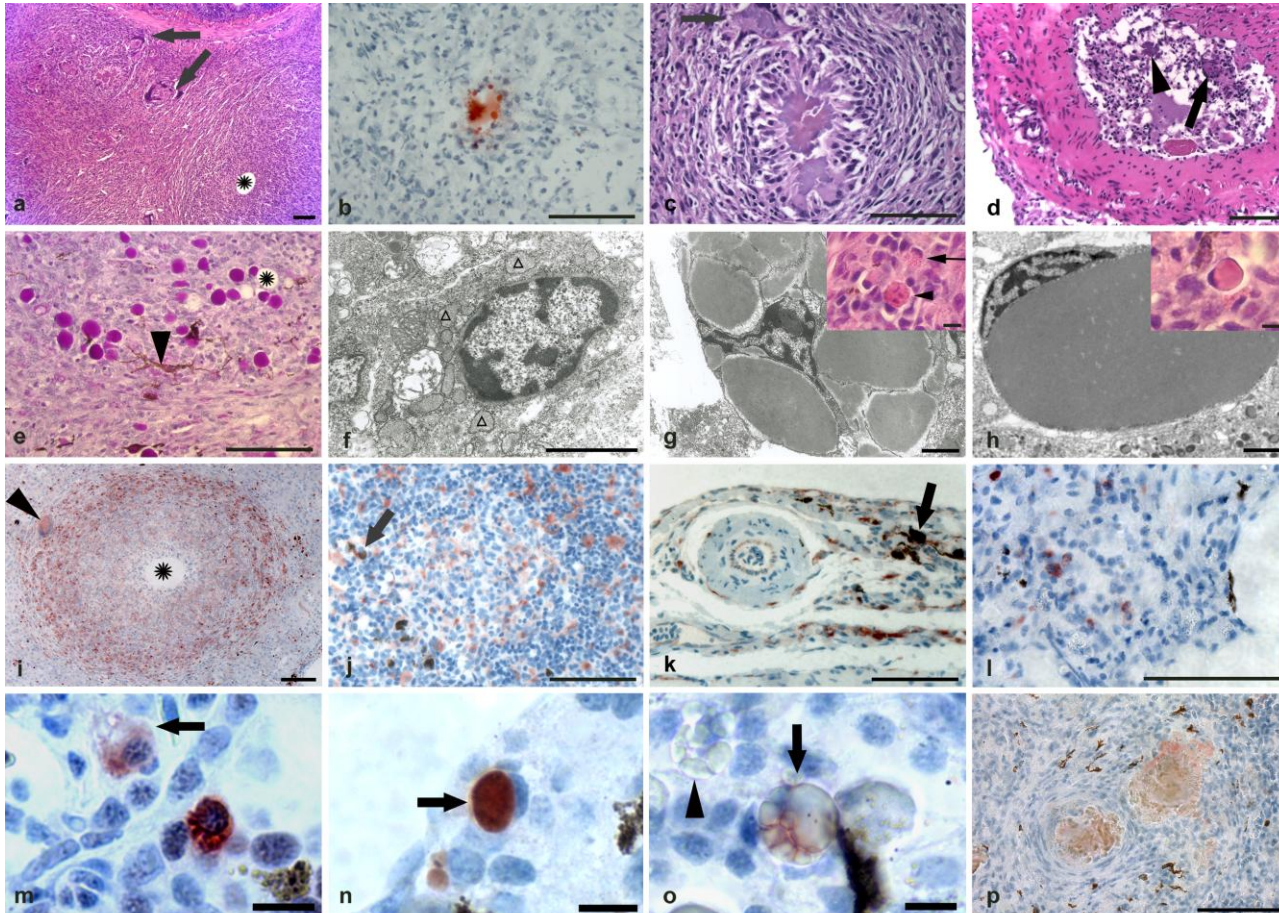
The development of systemic autoimmunity may result as an undesired side-effect following vaccination, and this condition was recently shown to occur in farmed salmon (*Salmo salar*). Several of previously reported side-effects following vaccination of fish should therefore be reviewed in the light of this condition. Here, organs and pathological changes in three separate groups of fish severely affected by vaccination were investigated by different morphological methods ($n = 84$). Granulomas or microgranulomas were observed at the injection site and in several organs. Mott cells were observed in all tissues examined. Pannus-like changes with lymphocyte infiltrates were observed in spines. In conclusion, the reactions following vaccination were of a systemic nature that may be explained by a pathogenetic mechanism caused by systemic autoimmunity.

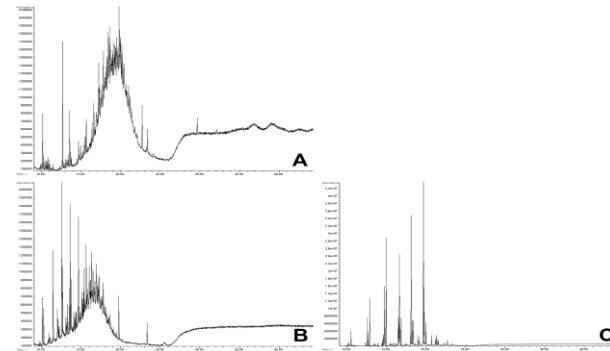
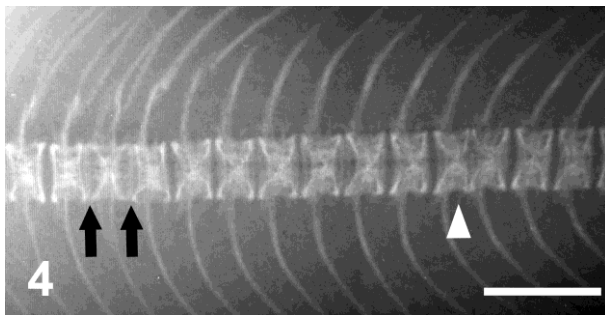
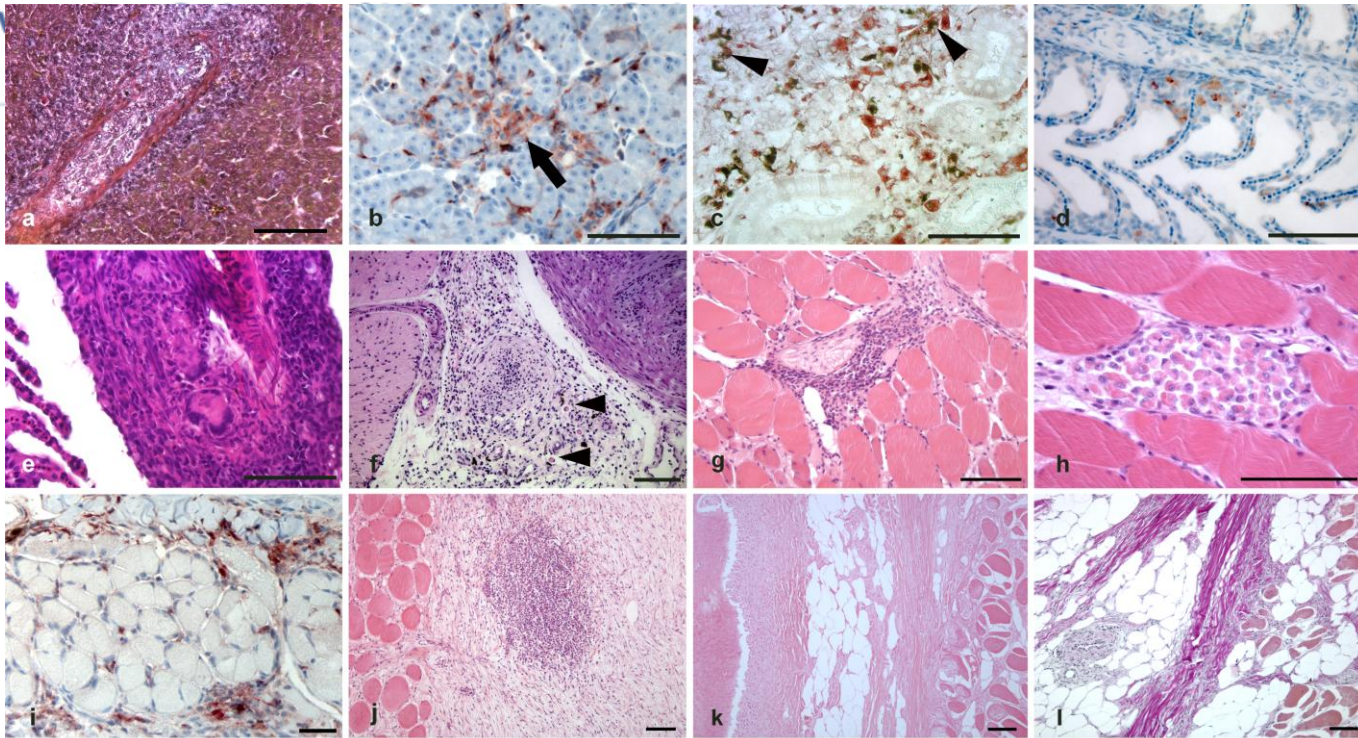
© 2010 Elsevier Ltd. All rights reserved.

1. Introduction

The successful prevention of a series of different infectious diseases in man and animals may be obtained by vaccines, but frequently, concerns are raised regarding their safety. Undesired side-effects following vaccine injections include fever, rash, flu-like symptoms, lymphadenopathy and swelling at the site of inoculation, but long-term effects such as hypersensitivity, induction of infection and autoimmunity are considered rare [1]. In general, the lack of a standardised vaccine safety assessment system is striking [2]. According to The European Agency for the Evaluation of

may develop into autoimmune diseases [1]. In addition, the dosage:body-weight ratio in salmon vaccination is considerably higher than that of mammals [7]. All these factors argue for higher frequency and severity of vaccine-induced side-effects in salmon compared to farmed mammals and humans. IL-1, a cytokine that plays a key role in the pathogenesis of systemic autoimmunity in adjuvant-injected mice [6], is induced in vaccinated salmon [8]. Recipient fish may over time develop mild to severe pathological changes as a consequence of vaccination, and observed adverse reactions include impaired growth rate, decreased carcass quality, spinal deformities, uveitis and inflammatory reactions in the



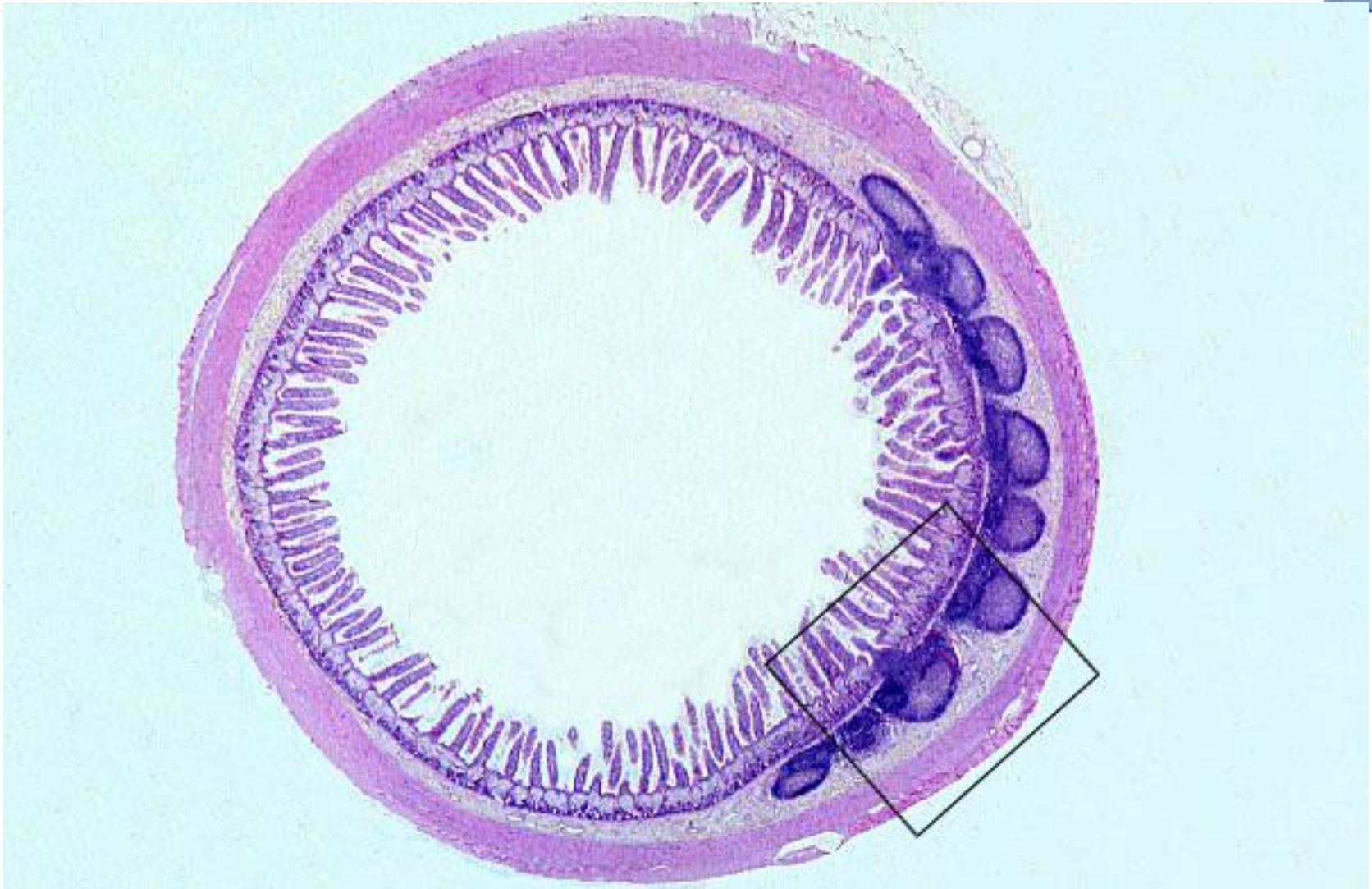


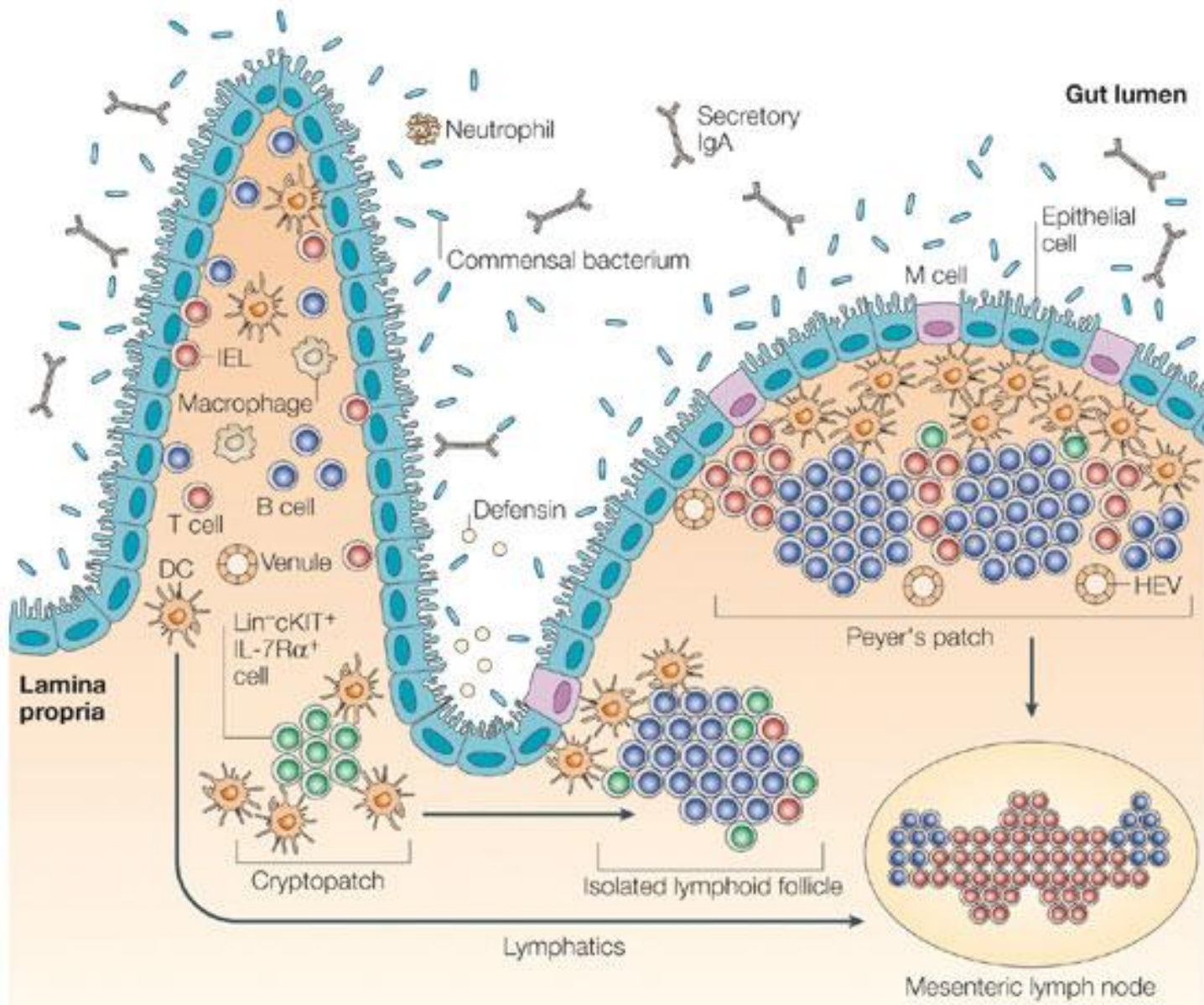
Kan dette forklare observert assosiasjon? (Berg et al. 2006 og Aunsmo et al. 2008)



Fremtidens vaksine

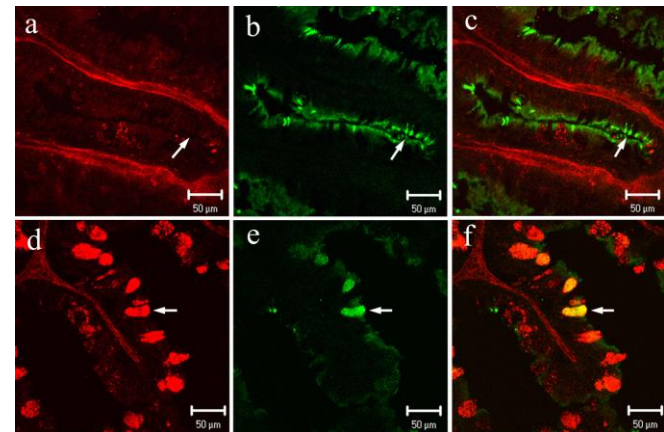
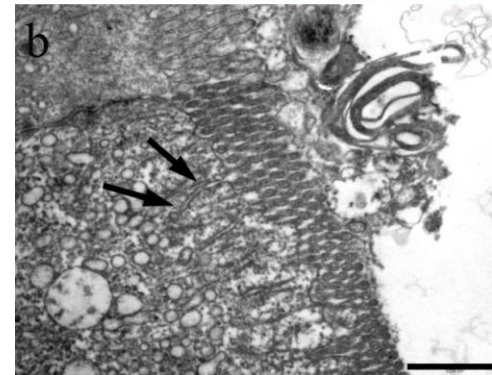
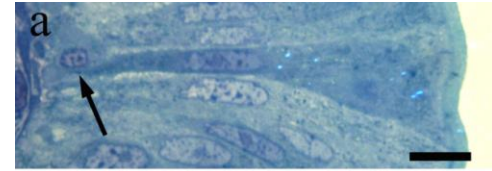
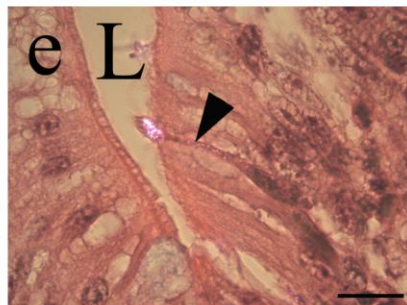
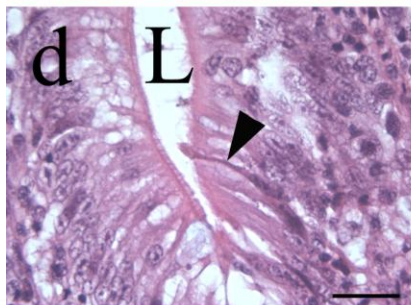
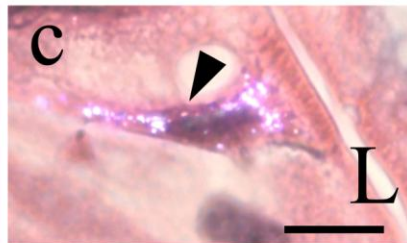
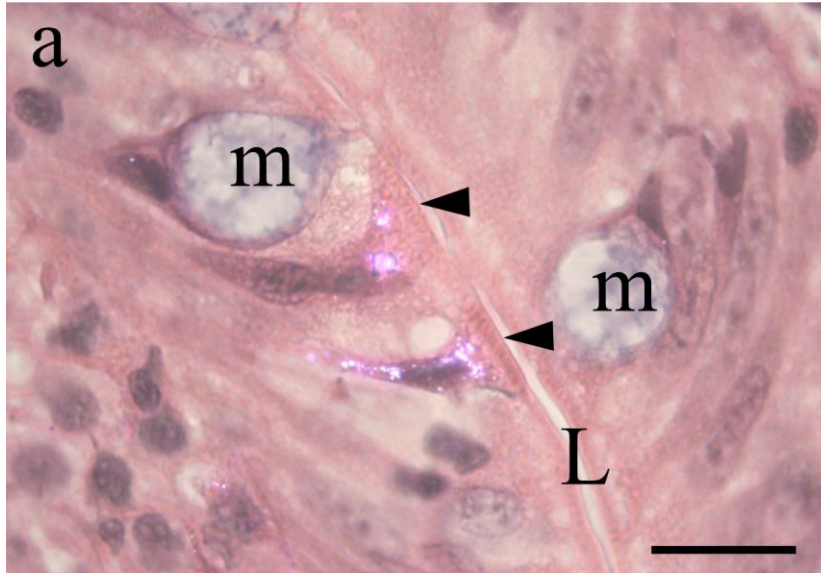
- Billig
- Uten bieffekter
- Lett å administrere
- Sikker – gir god beskyttelse
- Utgjør ingen risiko for konsument
- Kombinasjoner av løsninger?







Professor Hiroshi Kiyono, Universitetet i Tokyo: Vaksiner i ris!



Fuglem et al., *Dev. Comp. Immunol.* 2010; 34: 768-774.

