Avian Encephalomyelitis (Epidemic tremor)
Dr. Rajesh Agrawal, Associate Professor
Division of Veterinary Medicine, FVSc & AH, SKUAST-J, R.S. Pura, Jammu

Avian encephalomyelitis (AE) is a viral disease of young chickens, turkeys, Japanese quail, pheasants, and pigeons. Turkeys are less susceptible to natural infection and generally develop a milder clinical disease than chickens. Ducklings and guinea fowl are susceptible to experimental infection. AE is characterized by neurologic signs that result from infection of the CNS with an RNA virus in the family Picornaviridae. Infection occurs via vertical (egg born disease) and horizontal transmission. If a breeder flock becomes infected during egg production, the virus is vertically transmitted to the offspring and a major outbreak occurs. The disease often appears in a series of flocks hatched from the infected breeder flock. Field strains of the virus are enterotropic and multiply in the intestine. Infected birds shed the virus in their feces for a few days to a few weeks, which serves to spread the infection to hatchmates. There is no convincing evidence that the virus persists in infected birds. AE virus is resistant to environmental conditions and may remain infectious for long periods.

Age: Baby chicks of a few weeks age are most severely affected especially 1-2 weeks old. Adult birds also get infection but with little if any symptoms, except temporary drop in egg production by 20-30%.

Clinical Findings
Vertically infected chicks commonly show clinical signs during the first week after hatching, although signs may be present in a few birds at hatching. Clinical signs appear later in hatchmates that are horizontally infected by the fecal-oral route. Vertical infection followed by horizontal infection causes a characteristic biphasic mortality pattern. The main clinical signs are ataxia and leg weakness that varies from sitting on hocks to paresis that progresses to paralysis and recumbency. Fine tremors of the head and neck are evident in some birds and are characteristic of the disease. They are responsible for the common name, epidemic tremors. Tremors vary in frequency and severity and are best seen after birds are disturbed or excited. This can easily be done by placing the bird on its back and letting it right itself. Cupping the bird in one's hands often results in a buzzing feeling because of rapid, fine tremors. Severely affected birds lay on their side and exhibit intermittent fine tremors of the head, neck, and legs. Horizontally infected chicks usually show clinical signs at 2–4 wk of age; thus, clinical disease progresses through the flock for the first few weeks, and the episode is usually over by the time the flock is ~4 wk old. Morbidity and mortality rates vary and depend on the level of egg transmission and degree of immunity in the flock. In severe outbreaks, both morbidity and mortality may exceed 50%.
After 4 wk of age, chickens are resistant to disease but not infection. An exception occurs occasionally in older chickens after vaccination with egg-embryo–adapted commercial vaccines. Affected birds exhibit typical CNS signs like those seen in younger chicks.

In laying chickens, there is a sudden, 5%–10% drop in egg production, which usually lasts for <2 wk, followed by a return to normal production. There is no deterioration in egg shell quality. Hatchability may drop as much as 5% during the decline in egg production due to late embryonic mortality. Infected eggs are laid during the period of viremia, which usually lasts 1–2 wk.

Lesions:

No gross lesions are seen in the brain of infected birds. Gray to white foci may be visible on cut surfaces of the muscle of the gizzard. Weeks after infection, opacity of eye lenses (cataracts) may occur in a small percentage of chickens that survive the infection. Microscopic lesions in the CNS are found in the brain (cerebral peduncle, cerebellum, brain stem) and spinal cord and consist of degeneration and necrosis of neurons, perivascular lymphocytic cuffing, and gliosis with formation of glial nodules. In the cerebellum, there are areas of necrosis or loss of purkinje cells and replacement by glial nodules that extend into the molecular layer of the gray matter. Neuronal lesions of central chromatolysis, shrinkage and increased basophilia, satellitosis, and neuronophagia are best found in neuron clusters (nuclei) in the brain stem, arbor vitae of the cerebellum, and lateral horn (gray matter) of the spinal cord. Central chromatolysis of neurons is characterized by rounding of the cell contour and displacement of Nissl granules to the periphery. Dorsal root ganglia have multifocal nodular collections of lymphocytes.

A common microscopic lesion outside the CNS is diffuse or nodular lymphocytic infiltrates in the gizzard muscle, muscular layer of the esophagus and proventriculus, myocardium, and pancreas. Except for cataracts, microscopic lesions generally are not found in infected adults unless they are showing nervous signs, in which case they will have CNS lesions similar to those in young birds.

Diagnosis

Diagnosis is based on history, clinical signs, and characteristic histopathologic lesions in the brain and spinal cord. The diagnosis is best confirmed by isolation and identification of the virus. Tissues collected for virus isolation must include the brain and duodenum with the pancreas. Demonstration of AE virus antigen in the brain, spinal cord, and other tissues by immunofluorescent and immunohistochemical staining is a reliable method of diagnosis. The major differential diagnosis for neurologic signs in very young chicks is bacterial or mycotic encephalitis. Rickets and nutritional encephalomalacia are next in the list of differential diagnoses, although the clinical manifestations of these diseases differ from those in AE. In adults, serologic testing of acute and convalescent sera to demonstrate an increase in antibody titers helps establish a diagnosis.

Prevention
Immunization of breeder pullets at 10–15 wk of age with a commercial live vaccine prevents vertical transmission of the virus and provides progeny with maternal immunity. Vaccination of table-egg flocks is also advisable to prevent decreases in egg production. AE vaccine is usually combined with fowlpox vaccine and given by wing-web inoculation. Also available is a live fowlpox-vectored infectious laryngotracheitis and AE combination vaccine. Affected chicks and poults are ordinarily destroyed because they rarely recover.

AE virus does not cause disease in people or other mammals.

TREMVCAC-FP® (Intervet) Avian Encephalomyelitis - Fowl Pox Vaccine Live Virus: it is a lyophilized live vaccine for use in breeder pullets and egg layers to prevent egg production loss and clinical disease due to avian encephalomyelitis and fowl pox. It is approved for wing web inoculation.

TREMVCAC-FP contains: ● The original Calnek strain of avian encephalomyelitis (AE) virus. ● Fowl pox virus.

Indications for Use: Tremvac-FP® is indicated for the immunization of chickens at 8 to 17 weeks of age and turkeys 10 to 22 weeks of age avian encephalomyelitis and fowl pox. Tremvac-FP is approved for use via wing web inoculation.

Proper vaccination should result in local swelling and scab formation at the vaccination site. At 7 to 10 days post-vaccination, a few birds should be examined for takes.

Do not vaccinate prior to 8 weeks of age or within 35 days of the onset of egg production. Do not vaccinate birds during egg production. Do not vaccinate birds within 21 days of slaughter.

VECTORMUNE® FP- LT+AE (AVIAN ENCEPHALOMYELITIS - FOWL POX - LARYNGOTRACHEITIS VACCINE Live Virus, Live Fowl Pox Vector) Manufactured by: Biomune Company, Distributed by Ceva Animal Health, LLC, Lenexa, KS, USA

VECTORMUNE® FP LT & AE is a genetically engineered live virus vaccine for the vaccination of chickens as an aid in the prevention of fowl pox and laryngotracheitis. The vaccine also contains a conventional live virus for vaccination as an aid in the prevention of avian encephalomyelitis. The fowl pox vaccine has been genetically engineered to contain and express key protective laryngotracheitis virus antigens.

Indication: Administer by wing web to healthy, susceptible chickens at least 8 weeks old but at least 4 weeks prior to the start of egg production. Chickens receiving this vaccine must not have been previously vaccinated with a fowl pox vaccine.