IMMUNOLOGICAL RESPONSES IN FERRETS INOCULATED WITH INFECTIOUS BOVINE RHINOTRACHEITIS VIRUS VACCINES

by

ELSETHE JANE LEE

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Approved by:

[Signature]

Major Professor
DEDICATION

To my parents for their love and encouragement, which is never ending.
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## ABBREVIATIONS

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
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<tbody>
<tr>
<td>ATCC</td>
<td>American Type Culture Collection</td>
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<tr>
<td>BUN</td>
<td>Blood Urea Nitrogen</td>
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<tr>
<td>BVDV</td>
<td>Bovine Virus Diarrhea Virus</td>
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<tr>
<td>°C</td>
<td>Degrees Celsius</td>
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<tr>
<td>Ca</td>
<td>Calcium</td>
</tr>
<tr>
<td>CBC</td>
<td>Complete Blood Count</td>
</tr>
<tr>
<td>cc</td>
<td>Cubic Centimeter</td>
</tr>
<tr>
<td>Ci</td>
<td>Curie</td>
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<tr>
<td>Cl</td>
<td>Chloride</td>
</tr>
<tr>
<td>cm</td>
<td>Castrated Male</td>
</tr>
<tr>
<td>CO₂</td>
<td>Carbon Dioxide</td>
</tr>
<tr>
<td>CPE</td>
<td>Cytopathic Effect</td>
</tr>
<tr>
<td>DMSO</td>
<td>Dimethyl Sulfoxide</td>
</tr>
<tr>
<td>DNA</td>
<td>Deoxyribonucleic Acid</td>
</tr>
<tr>
<td>DNase</td>
<td>Deoxyribonuclease</td>
</tr>
<tr>
<td>DPM</td>
<td>Disintegrations per Minute</td>
</tr>
<tr>
<td>EDTA</td>
<td>Ethylene Diamine Tetraacetic Acid</td>
</tr>
<tr>
<td>F°</td>
<td>Fahrenheit</td>
</tr>
<tr>
<td>f</td>
<td>Female</td>
</tr>
<tr>
<td>FCS</td>
<td>Fetal Calf Serum</td>
</tr>
<tr>
<td>g</td>
<td>Gravity</td>
</tr>
<tr>
<td>H₂O</td>
<td>Water</td>
</tr>
<tr>
<td>Mb</td>
<td>Hemoglobin</td>
</tr>
<tr>
<td>HS</td>
<td>Horse Serum</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
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<tr>
<td>--------------</td>
<td>-------------</td>
</tr>
<tr>
<td>$^3$H-TdR</td>
<td>Tritiated Thymidine</td>
</tr>
<tr>
<td>IBRV</td>
<td>Infectious Bovine Rhinotracheitis Virus</td>
</tr>
<tr>
<td>IM</td>
<td>Intramuscular</td>
</tr>
<tr>
<td>IN</td>
<td>Intranasal</td>
</tr>
<tr>
<td>K</td>
<td>Potassium</td>
</tr>
<tr>
<td>LA Strain</td>
<td>Los Angeles Strain</td>
</tr>
<tr>
<td>m</td>
<td>Male</td>
</tr>
<tr>
<td>MDBK</td>
<td>Madin Darby Bovine Kidney</td>
</tr>
<tr>
<td>MEM</td>
<td>Minimal Essential Media</td>
</tr>
<tr>
<td>mg</td>
<td>Milligram</td>
</tr>
<tr>
<td>ml</td>
<td>Milliliter</td>
</tr>
<tr>
<td>MLV</td>
<td>Modified Live Virus</td>
</tr>
<tr>
<td>mmole</td>
<td>Millimole</td>
</tr>
<tr>
<td>Na</td>
<td>Sodium</td>
</tr>
<tr>
<td>NaCl</td>
<td>Sodium Chloride</td>
</tr>
<tr>
<td>NADL</td>
<td>National Animal Disease Laboratories</td>
</tr>
<tr>
<td>ng</td>
<td>Nanogram</td>
</tr>
<tr>
<td>n</td>
<td>Sample Size</td>
</tr>
<tr>
<td>NS</td>
<td>Nonsignificant</td>
</tr>
<tr>
<td>P</td>
<td>Phosphorus</td>
</tr>
<tr>
<td>PBS</td>
<td>Phosphate Buffered Saline</td>
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<tr>
<td>PCV</td>
<td>Packed Cell Volume</td>
</tr>
<tr>
<td>PFU</td>
<td>Plaque Forming Unit</td>
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<tr>
<td>PHA</td>
<td>Phytohemagglutinin</td>
</tr>
<tr>
<td>PSK</td>
<td>Penicillin-Streptomycin-Kanamycin</td>
</tr>
<tr>
<td>RNA</td>
<td>Ribonucleic Acid</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Full Form</td>
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<tr>
<td>--------------</td>
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<tr>
<td>RPMI</td>
<td>Roswell Park Memorial Institute</td>
</tr>
<tr>
<td>SDH</td>
<td>Sorbital Dehydrogenase</td>
</tr>
<tr>
<td>SGPT</td>
<td>Serum Glutamate Pyruvate Transaminase</td>
</tr>
<tr>
<td>SMA-12</td>
<td>Sequential Multiple Analysis</td>
</tr>
<tr>
<td>TCA</td>
<td>Trichloroacetic Acid</td>
</tr>
<tr>
<td>T.P.</td>
<td>Total Protein</td>
</tr>
<tr>
<td>uCi</td>
<td>Microcurie</td>
</tr>
<tr>
<td>ug</td>
<td>Microgram</td>
</tr>
<tr>
<td>UV</td>
<td>Ultraviolet</td>
</tr>
<tr>
<td>WBC</td>
<td>White Blood Cells</td>
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SECTION I
INTRODUCTION
INTRODUCTION

Infectious bovine rhinotracheitis (IBR), a herpes virus infection in cattle, is capable of producing a latent infection and a variety of clinical manifestations ranging from respiratory to reproductive syndromes (Kahrs, 1977; Owens, Chow and Mollelo, 1964; French, 1962; Abinati and Plumer, 1961; McKercher, et al., 1959; Madin, York and McKercher, 1956). Recrudescence or activation of latent IBRV infections can occur spontaneously during periods of stress when physiological hormone levels are altered or with the administration of corticosteroids (Davies and Carmichael, 1973). These characteristic syndromes in addition to the ubiquitous nature of IBRV in the United States and other parts of the world have attributed to an increasing concern to manage this disease. The detrimental effects of IBRV infections are manifested in cattle of all ages and cause economic losses in milk and meat production (Kahrs, 1977).

Presently, the prevention and control of IBRV is accomplished through proper animal management and most importantly through vaccination schemes. The availability of a variety of vaccines, such as modified live virus (MLV) or inactivated virus vaccines and the different routes of inoculation have left the practitioner and cattle owner with a seemingly difficult choice. Investigations evaluating these vaccines have elucidated some of the mechanisms by
which immunity to IBRV infections is achieved. It has been reported that the modified live IN vaccine provides protection for pregnant dams and elicits local antibody and interferon production (Todd, Volenc and Paton, 1972) but has no distinct advantage over ML-IM vaccines in humoral antibody production (McKercher and Crenshaw, 1971). Inactivated vaccines with and without adjuvant have been found to produce adequate levels of humoral antibody (Haralambiev, 1976; Karadjov, 1976; Juhasz, Palatka and Toth, 1973).

In influenza virus studies, the ferret, Mustela putorius furo, has been used to evaluate the efficacy of potential human vaccines, since ferrets closely simulate a similar course of disease observed in humans. These studies have provided much information on the local and humoral responses to various vaccines (Potter, et al., 1972a, 1972b). The National Animal Disease Center (1977) and Porter, Larsen and Cox, (1975) have studied the humoral response of IBRV infected ferrets, and have found that ferrets produce serum antibodies to IBRV.

The immune response is complex and it has been demonstrated that IBRV latency and spontaneous recrudescence occurred in the presence of humoral antibodies (Sheffy and Rodman, 1973). As a result, investigators are trying to determine the importance of the cell mediated immune responses to IBRV. The primary factors of the cellular immune system are the thymus (T) dependent lymphocytes. There are various techniques for measuring T cell stimulation or
blastogenesis to a mitogen or specific antigen. The method reported by Sorenson, Anderson and Giese, (1969) measures the incorporation of radioactive labelled DNA precursors as an index of T-lymphocyte blastogenesis and is commonly used as an indication of CMI responses.

It has been found that adrenal cortical hormones by causing cytotoxic effects and altering the metabolism and activity in the cellular components of the immune system, suppress the CMI responses to either mitogen or specific antigen. In vitro studies have illustrated the inhibition of blastogenesis in lymphocyte cultures supplemented with cortisol before or simultaneously with the stimulus (Fauci and Dale, 1974; Tormey, Fudenberg and Karmin, 1967; Stevens and Dougherty, 1964; Nowell, 1961).

A better understanding of the complex mechanisms of the immune response in an animal to an antigen and the suppression observed in lymphocytes with cortisol, would provide insight for establishing adequate resistance through vaccination and controlling latency and the recrudescence of IBRV infections. The objectives of this investigation are to (1) study the hematological and serum profiles of ferrets, (2) determine whether the ferret can be used as a suitable laboratory animal model for evaluating immunity to IBRV infections by studying the humoral and cell mediated immune responses of control and IBRV vaccine inoculated ferrets and (3) study the in vitro effect of cortisol on ferret lymphocytes in whole blood cultures.
SECTION II

REVIEW OF LITERATURE
The Ferret as a Laboratory Animal.

The domestic ferret, *Mustela putorius furo* L. in the United States includes the brownish-black and yellowish colored fitch ferret and the yellow-cream colored, ruby-eyed albino or English ferret (Willis and Barrow, 1971). As members of the weasel family, Mustelidae, these carnivores are small in size and are easily manageable in colonies for research purposes. Contrary to myths, the ferret has an agreeable disposition and can be handled without an animal restrainer and gloves, for procedures such as rectal temperatures, inoculations and weighings. The breeding, management, growth and development has been studied (Hammond and Chesterman, 1972; Willis and Barrow, 1971; Hammond, 1969; Pockson, 1956; Grinham, 1952; and Pyle, 1940), and the maintenance of laboratory colonies is conducive to ferrets.

Anesthesia for the Ferret.

Procedures causing prolonged discomfort and pain in ferrets require the use of anesthesia. In general, the mustelids are easily anesthetised with injectable agents and have similar responses as cats, except that adverse effects to morphine-like drugs are not encountered. Chemical restraint by alphoxolone-alphadalone, ketamine hydrochloride and fentanyl citrate-fluanisone have been found to provide safe and effective sedation (Green, 1978).
Increased doses of the above agents provide light anesthesia for surgical procedures. Fentanyl citrate and fluanisone supplemented with a halothane-oxygen mixture followed by nalorphine hydrobromide also produced satisfactory results (Harpur, 1978). Light ether induction provides short periods of immobilization and anesthesia, but should not be used for restraint purposes due to the difficulty in handling as a result of the animals sneezing and paddling reflexes (Pyle, 1940). Pentobarbitone sodium, although having a narrow safety margin given intraperitoneally (IP) for prolonged anesthesia or in conjunction with ether has been found to be satisfactory (Lumb, 1963). The use of chloroform is not recommended near the animal housing facilities, since low concentrations of vapor cause nursing ferrets to scatter the young about the cage (Hammond and Chesterman, 1971).

**Blood Collection Procedures in Mustelids.**

Small quantities of blood for studies such as packed cell volume and blood smears can be obtained by deeply pricking the skin at the tip of the ear (Pyle, 1940) or by overcutting the toenails (Bergman, Lodmell and Hadlow, 1972). Larger samples up to 12 ml can be drawn with relative ease by cardiac puncture when animals are anesthetised (Baker and Gorham, 1951). Safe multiple bleedings in minks *Mustela vison*, have been reported by Fetch and Wobeser, (1970) using a jugular venipuncture technique with a 20 or
22 gauge, 1 inch needle. Bergman, et al. (1972) describes the surgical implantation of a polyethylene catheter into the jugular vein for studies which require numerous blood samplings over short time intervals. The coccygeal, saphenous and cephalic veins are small and not routinely used (Pyle, 1940).

Hematological and Serum Studies.

The majority of information on mustelid hematology and serology has been provided by studies on the mink. The blood parameters studied were packed cell volume (PCV), hemoglobin (Hb), total erythrocyte count (RBC), plasma and blood volume, total and differential white blood cell count (WBC), and mean corpuscular volume and density (Schalm, Jain and Carroll, 1975; Ringer, et al., 1974; Fletch and Karstad, 1972; Kennedy, 1935a, 1935b, 1935c). Various constituents in serum that have been studied in minks include total protein, albumin, globulin fractions, total lipids, 3-lipoproteins, phospholipids, total, free and esterified cholesterols, glucose, calcium, nitrogen and carbon dioxide levels (Rotenberg and Jorgensen, 1971; Kubin and Mason, 1948). In the ferret, however, the information is limited to total RBC and WBC, and differential WBC counts (Hammond and Chesterman, 1972; Pyle, 1940; Fox, 1923). Preliminary blood chemistry studies on glucose, calcium and nonprotein nitrogen have been done by Pyle, (1940). Physiological parameters such as average normal rectal temperatures
respiration rate ranges (Pyle, 1940), cardiac output (Kempf and Chang, 1949) and growth and development rates (Shump and Shump, 1978) have been studied.

**Studies Using Ferrets.**

An invaluable laboratory animal model, the ferret, has been used in a diversity of studies, including endocrinology (Wu and Chang, 1972), reproductive physiology (Chang, Casas and Hunt, 1971), pharmacology and toxicology (Steinberg, 1973) and microbiology (Rowlands, 1957). The ferret, because of its susceptibility to the canine distemper (Carpenter, 1976; Farrell, 1971) and human influenza viruses (Toms, *et al*., 1977, 1975; Small, 1976) have been used extensively for *in vivo* investigations and for the evaluation of vaccines for these viruses. *In vitro* cultures of respiratory and other ferret tissues have been developed (Kingsman, Toms and Smith, 1977a, 1977b; Toms, Rosztoczy and Smith, 1974; Gould *et al*., 1972) and are used for the localization of influenza viruses.

**The Clinical and Immune Responses to Human Influenza Viruses in Ferrets.**

The ferret is most frequently used in influenza virus studies, since the influenza virus, although highly transmissible to a variety of animal hosts, simulates a pathological response in ferrets most similar to that found in man. This is of particular interest for studying the pathogenesis and effects of human influenza virus vaccines.

Previous studies (Potter, Jennings and McLaren, 1973;
Potter, McLaren and Shore, 1973; Potter, et al., 1972a; Potter, et al., 1972b) have compared the clinical (febrile) and immunological responses of ferrets to live, attenuated and inactivated influenza virus vaccines given either IN, IM or IM with adjuvant. Comparisons between Freund's incomplete adjuvant and adjuvant 65 were also made. The concentration of virus and the protein constituents collected in the nasal washes were correlated with immunity to the virus (Toms, et al., 1977; Potter, Shore, McLaren, 1972; Shore, Potter and McLaren, 1972). Hemagglutinin inhibiting (HI) and neutralizing antibodies detected in the nasal washes and HI antibodies detected in the serum were titered and found at high levels in live virus infected (IN) animals, as compared to animals infected (IM) with attenuated or inactivated influenza virus. Ferrets infected with live influenza virus and challenged with the same virus were found to be resistant to infection, as manifested by the lack of a febrile response and the absence of an elevated immune response. Live vaccines administered IN, produced the most effective immunity, followed by attenuated virus vaccines given intranasally. Inactivated virus administered with adjuvant, IM, produced humoral immunity but to a lesser degree. Inactivated virus given IM or IN with saline produced no immunity. Challenge with an influenza virus of a different strain, elicited elevated antibody and febrile responses in live virus infected (IN) ferrets, similar to nonimmunized control animals (Potter, Jennings and McLaren,

Isolation and Studies of IBR in Mustelidae.

Porter, Larsen and Cox (1975) reported the isolation of bovine herpesvirus I, commonly known as infectious bovine rhinotracheitis virus (IBRV) from minks and ferrets. An isolate of IBRV was obtained from a clinically normal, healthy ferret while a second was removed from the intestine of a mink displaying signs of clinical mink viral enteritis. Electron microscopic examinations of the cell cultures infected with these isolates showed viral particles with typical morphological characteristics similar to a herpes virus. Serum neutralization tests, using rabbit antisera prepared against the mink and ferret isolates prevented the formation of infectious foci. Indirect immunofluorescence tests using acetone-fixed ferret kidney cells, showed no reaction of isolates with antisera to a variety of herpes type viruses (human cytomegalovirus Ab 169, monkey cytomegalovirus GR 2757, varicellazoster virus, Epstein-Barr virus, equine herpes virus (type 1 and 2), feline herpes virus C27, herpes tamarinus virus, pseudorabies virus or Marek's disease virus). A weak cross-immunoflourescence reaction occurred between human herpes simplex type one serum and the mustelid isolates, but no cross-neutralization was observed. An extensive cross-immunoflourescence reaction occurred between human herpes simplex type one serum and the mustelid isolates, but no cross-neutralization was observed.
An extensive cross-immunofluorescence reaction was observed between the Cooper strain of IBR and the mink and ferret isolates, using antisera to the three viruses. Serum from the animals contained no detectable neutralizing antibodies, although one had weak reactions with IBRV by immunofluorescence. Porter, Larsen and Cox, (1975) also indicate that minks inoculated intraperitoneally (IP) with IBRV did not develop clinical signs of illness, but did produce antibodies three weeks post infection.

In more recent studies (NADC, 1977) young adult English ferrets were inoculated with a virulent field strain of IBRV, IN and IM in high and low doses. The ferrets were challenged six weeks post-inoculation with high doses of the virulent field strain and immunity was evaluated on the basis of antibody development and virus recovered from the nasal wash. It was found that only high doses of virulent field strains produced full immunity and protection against the challenge. Only varying degrees of low level protection were achieved for animals inoculated with low doses of the virus.

**Infectious Bovine Rhinotracheitis Virus.**

**Properties of the virion.** The IBRV, classified in the herpes group, is a double stranded DNA virus that replicates in the nucleus. The IBR virion morphologically typifies the herpes virus family in having an icosahedral capsid with an envelope and is approximately 100-150 millimicrons in diameter (Bruner and Gillespie, 1973). Characteristic of the herpes viruses,
IBRV tends to remain cell associated after replication and contains both host and viral constituents in the envelope (Wagner, 1974).

**Historical Appearance of Infectious Bovine Rhinotracheitis in Cattle.**

Although known for many years in Europe as the cause of Blaschenausschlag, a mild genital tract infection in cattle, (Saxegaard, 1970; McKercher, 1963; Zwick and Gminder, 1913), IBR has been recognized as a clinical respiratory tract disease afflicting cattle in the U.S. only since 1950 (Miller, 1955; Schroeder and Mays, 1954). Isolation from nasal secretions of cattle and the subsequent culture in bovine embryonic kidney cells was first accomplished by Madin, York and McKercher, (1956). Later it was found that the causative agent of IBR was also implicated in infectious pustular vulvovaginitis (Kendrick, Gillespie and McEntee, 1958; McKercher, et al., 1959). Less than a decade after IBR was isolated, Armstrong, Pereira and Andrews, (1961) classified IBR into the herpes virus family. Subsequently, studies characterizing the morphology and biochemical features of the virus followed (Tousimis, et al., 1958; Hahnfeld and Hahnfeld, 1964; Jasty, et al., 1969). Although ubiquitous to most of the United States, (Kahrs, 1974), IBRV has also been reported in other areas of the world (St. George and Philpott, 1972). Reports indicate that IBR has been isolated from goats (Mohanty, et al., 1972), swine, (Nelson, Mare and Glock, 1972), water buffalo (St. George and Philpot, 1972) and mustelids (Porter,
Larsen and Cox, 1975).

Pathological Manifestations of Infectious Bovine Rhinotracheitis Virus.

The disease syndromes currently recognized as being caused by IBRV are rhinotracheitis, (Madin, York and McKercher, 1956), vaginitis, (Gillespie, et al., 1959; McKercher, et al., 1959), conjunctivitis, (Abinate and Plumer, 1961; McKercher, Wada and Straub, 1958), meningoencephalitis, (French, 1962), septicemia (Galloway, 1972), and abortion, (McKercher and Wada, 1964; Owens, Chow and Mollelo, 1964). Highly contagious, all forms of the disease are transmissible by contact (Bruner and Gillespie, 1973; Chow, Mollelo and Owen, 1964). The usual course of the disease extends from two to seven days (Runnels, Monlux and Monlux, 1967). Although most infections are rarely fatal for the adult bovine, the effects can be profoundly manifested by a decrease in feed efficiency, milk production and reproduction. Mortality can occur if the disease is complicated by secondary bacterial infections or viral infections (Kahrs, 1977).

Primarily a respiratory disease, IBRV commonly causes such clinical signs as pyrexia, anorexia, lethargy, dypsnea, and upper and lower respiratory tract congestion. Profuse ocular and nasal discharges, initially clear, often become mucopurulent (Kahrs, 1977). Necrotic lesions around the nostril and muzzle as well as in the pharynx, larynx, trachea and bronchi, (McKercher, et al., 1959) are found in severe cases.
Encephalomyelitis occasionally occurs in young cattle with chronic IBRV infections and results in incoordination instability, and occasional circling or licking at the hind quarters (Beck, 1975), followed by recumbency and death (McKercher, Bilbrack and Richards, 1970). Neonates infected in utero during late gestation or shortly after birth develop an acute systemic infection which is characterized by respiratory distress and necrotic lesions from the oral cavity down to and including the four stomach compartments. Peritonitis, causing diarrhea, may occur. In general, IBRV infections in newborn calves are fatal (Kahrs, 1977).

Infectious pustular vulvovaginitis (IPV), an IBRV infection of the vaginal and vulvar mucosa is characterized by pustule formation and a mucopurulent discharge. In mild cases, these conditions are not blatant (McKercher, 1963). This form of IBRV infection can be transmitted by natural breeding or other means involving direct contact (Kahrs and Smith, 1965). Careful diagnosis of IPV is warranted, since it is often confused with parturition or sadistic injuries, granular vaginitis and mycoplasma infections. Bulls that are bred to cows with IPV can develop severe balanoposthitis, which is characterized by lesions similar to those found in cows with IPV infections (Kendrick and McEntee, 1967).

Abortion caused by IBRV infections in pregnant and susceptible females occurs in approximately 25% of the cases under field conditions (Kahrs, 1977). The incidence increases in experimental situations when animals are given larger
doses of virus (Owen, Chow and Mollelo, 1964). Abortion may occur as long as 100 days after IBRV is clinically apparent in the herd (Kahrs, 1977).

**Histopathology of Infectious Bovine Rhinotracheitis Lesions.**

The whitish necrotic lesions and pustules found in severe IBRV infections consist mainly of leucocytes, fibrin and necrotic epithelial cells (Kahrs, 1977). Inclusion bodies within cell nuclei have been reported by Crandell, Cheatham and Maurer, (1959).

**Diagnosis and Confirmation of Infectious Bovine Rhinotracheitis.**

Serological tests, detecting the presence of IBRV antibodies, virus isolation or fluorescent antibody tests, coupled with a description of the appropriate clinical signs and history are commonly employed in the laboratory diagnosis of IBRV infections.

**Types of Vaccines Developed Against Infectious Bovine Rhinotracheitis.**

a) Modified live IBRV vaccines

Modified live IBRV for IM administration was the first IBRV vaccine developed (Swartz, *et al.*, 1957). The vaccine is widely used but is restricted from pregnant dams as the virus can induce abortion (Kelling, *et al.*, 1973). Bartha, (1973) reports on the production and use of heat attenuated IBRV vaccine for IM use, which does not cause abortion in pregnant cows and can be inoculated in calves. His study
indicate that repeated inoculations of this vaccine result in increased serum neutralizing antibody levels. In general, vaccinated calves, less than six weeks old, should be re-vaccinated, since colostrally acquired maternal antibodies may interfere with the immune response against the initial inoculation of the antigen. Vaccination of calves suckling pregnant cows is contraindicated, since virus may be shed and cause the dam to abort (Kelling, et al., 1973).

A single successful immunization usually results in the development of humoral antibodies lasting three to six years, (Chow, 1972; Rosner, 1968), and provides partial protection against challenge with virulent virus and abortion, (Kahrs, 1974). Annual vaccinations may be needed to insure cell mediated immune defenses (Studer, 1973).

There are relatively minimal undesirable effects resulting from MLV given IM. Aside from causing abortion in pregnant dams (Kelling, et al., 1973), Frank, Marshall and Smith, (1973) report the occurrence of fever and leukopenia without clinical signs or virus shedding. Straub, (1972) found that heifers vaccinated IM, shed virus when challenged with virulent IBRV.

Field trials by Schell, et al., (1971) have demonstrated the high level of antigenicity in multivalent vaccines for IM use (IBR/bovine virus diarrhea (BVD)/para influenza-3 (PI-3) with Pasteurella and IBR-PI-3 with Pasteurella) and the production of antibodies to all components.

Todd, Volenc and Paton, (1971) developed the first
MLV vaccine against IBR for IN administration. The relative safety of this vaccine for inoculation into all cattle, including pregnant dams has attributed to its widespread acceptance. Utilizing a natural route of infection, IN vaccines provide rapid protection (within 40-72 hours post vaccination) against virulent IBRV challenge (Todd, 1974). Resistance was attributed to the production of interferon and secretory antibodies at the mucosal surfaces (Todd, Volenec and Paton, 1972; Todd, Volenec and Paton, 1971). McKercher and Crenshaw, (1971); however, demonstrated the production of serum neutralizing antibodies in cattle vaccinated with IN vaccine comparable to levels produced in animals inoculated with IM vaccine and report no distinct advantage of IN vaccine over the vaccine produced for IM use. Although not completely evaluated, annual inoculations of IN vaccines are recommended for restimulating humoral and cellular immune responses (Kahrs, 1977). Adverse side effects which are rare and often mild, include fever, nasal discharge, virus shedding (Hyland, Easterday and Pawlisch, 1975) and lesions in the nasal mucosa (Kahrs, Hillman and Todd, 1973). Laborious restraint of the bovine head while inoculating may be an undesirable task (Kahrs, 1977).

Multivalent vaccines for IN use containing IBRV/PI-3/bovine adenovirus-3 and IBRV/PI-3 have been found to induce antibody production in calves and adults, to all components of the vaccine (Zygraich, Vascoboinic and Huygelen, 1976; Todd, 1976).
b) Inactivated IBRV vaccines

The efficacy of inactivated IBR vaccines remains controversial and animals usually require annual vaccination to achieve adequate levels of antibodies. It has been intermittently available as a product containing two *Pasteurella* spp and the inactivated PI-3 virus (Kahrs, 1977). Schipper and Kelling, (1974) tested a commercially available inactivated vaccine for subcutaneous (SC) use and found no detectable serum neutralizing antibodies. Kendrick and Osburn, (1973) found commercially inactivated aluminum hydroxide adjuvated IBR vaccine produced levels of serum antibody in bovine fetuses comparable to the levels in similarly vaccinated adults. Judhasz, Palatka and Toth, (1972) report formalin inactivated IBRV adjuvated with aluminum hydroxide gel absorbent administered SC, resulted in high titers of serum neutralizing antibodies. Only mild symptoms were observed when animals were simultaneously challenged IM and intratracheally or IN and by scarification into the vaginal mucosa with large quantities of virulent virus. Karadjov, (1976) describes the high level of immunogenicity elicited by an inactivated vaccine administered with an oil adjuvant, as determined by the resistance to challenge with virulent IBRV. Haralambiev, (1976) reports of the production of virus neutralizing factors and subsequent resistance to challenge with virulent virus in calves vaccinated with an ethanol and saponin inactivated IBRV given in the nasal submucosa. The efficacy of the IN route was found superior compared to the
SC inoculation of the vaccine.

Possible adverse effects include hypersensitivity reactions such as anaphylaxis from adjuvated vaccines and other less severe, non-fatal allergic syndromes (Kahrs, 1977).

Cell Mediated Immunity and Blastogenesis.

The host response to a viral infection involves both humoral and CMI responses. In the past the main focus of viral immunity studies, centered around the humoral component of the immune system. It is becoming increasingly apparent that resistance to and recovery from certain viral infections involves the CMI responses (W.H.O., 1973). There is mounting evidence that infections caused by herpes and parainfluenza viruses are more severe in individuals with deficiency syndromes of the CMI responses but essentially normal in those incapable of producing humoral antibody (Allison, 1972; Merigan and Stevens, 1971).

Until recently, the lack of suitable quantitative in vitro assays has hampered investigations in CMI. One method for testing CMI to study the blastogenic transformation of lymphocytes by the mitogen, phytohemagglutinin (PHA), a glycoprotein extract from the red kidney bean, was first reported by Nowell, (1960). The stimulation of lymphocytes, resulting in their transformation or blastogenesis (Stites, 1976), is characterized by the synthesis of deoxyribonucleic acid in thymus-dependent (T dependent) lymphocytes (Alhaji, Muscoplat and Johnson, 1974; Oates, et al., 1972; Piquet and Vasalli, 1972; Blomgren and Svedmyr, 1971; Janossy and
Greaves, 1971).

In earlier studies, the lymphocyte response to PHA was evaluated microscopically on the basis of cellular morphology (Pospisil et al., 1975) and by autoradiographic techniques. Sorenson, et al., (1971) developed a method for measuring the incorporation of $^3$H-thymidine into PHA-stimulated cells. Since then, there have been a number of methods for quantitative assessment of the lymphocyte response to PHA (Lazary, et al., 1974; Faquet, 1974; Fernald and Metzgar, 1971; Phillips and Zweinmann, 1970).

It has been found that responsiveness of lymphocytes to PHA is diminished by in vivo or in vitro infection with viruses such as Newcastle disease virus (Montgomery, et al., 1967), herpes virus (Willems, et al., 1969), bovine viral diarrhea virus (Johnson and Muscoplat, 1973) and rubella virus (McMorrow, et al., 1973).

**Infectious Bovine Rhinotracheitis Virus and Blastogenesis.**

Rouse and Babiuk, (1974), demonstrated the blastogenic effect of specific antigen (UV irradiated IBRV) on the peripheral blood lymphocytes from cattle immunized or infected with IBRV by quantitating the cellular uptake of $^3$H-thymidine ($^3$H-TdR). The failure of lymphocytes from these animals to incorporate $^3$H-TdR when cultured with two other herpes virus antigens, herpes simplex and equine rhinopneumonitis, demonstrates the specificity of lymphocyte transformation. Blastogenesis was prevented by reacting the IBRV antigen with IBRV specific antibody before addition to the cultures.
An assay (Rouse and Babiuk, 1975) utilizing peripheral blood lymphocytes from IBRV immunized cattle, confirms the cell-mediated specificity of these lymphocytes to inhibit cytopathic effects by IBRV in cell cultures.

**Suppression of Cell Mediated Immunity and Adrenal Corticoids.**

Herpes viruses can be present as latent infections without clinical manifestations (Sprinkle, Vettri and Wade, 1976) and result in spontaneous recrudescence despite the presence of neutralizing antibodies (Snowden, 1965). The recrudescence of IBRV infections in cattle has been experimentally demonstrated by the administration of synthetic corticoids, as long as four years after the initial infection (Sheffy and Davies, 1972).

Corticosteroids have been found to suppress CMI responses to bacterial antigens (North, 1972); however, suppression of CMI as measured by lymphocyte transformation was not detected when cattle infected with IBRV were treated three months post infection with synthetic corticosteroid to induce recrudescence (Davies and Carmichael, 1973).

The adverse effects of certain adrenal cortical hormones on lymphocytes has been attributed to the actual cytotoxic activity of the hormones (Claman, 1972). It has been demonstrated that glucocorticoids cause destruction of small and medium-sized lymphocytes and the depression of mitosis in lymphatic tissues (Stevens, Colossides and Dougherty, 1966; Moore, 1966; Bellamy, Hanssens and Leonard, 1966). Claman,
(1972) further reported that cortisol produced an inhibitory effect on cellular metabolism by altering glucose uptake and nucleic acid synthesis. Studies measuring the incorporation of radioactive labelled precursors of the DNA molecule have demonstrated the inhibitory effect of cortisol on DNA synthesis in rat lymphatic tissues (Hofert and White, 1968a, 1968b) and the depletion of nucleic acid in mouse lymphatic tissues (Brinch-Johnsen and Dougherty, 1959). Stevens and Dougherty, (1964) reported the decrease of nucleic acid when measured by the incorporation of thymidine-2-14C into lymphatic tissue nucleic acid after cortisol addition.

Fauci and Dale, (1974) and Webel, et al., (1974) observed that corticosteroid administration significantly altered the absolute number of circulatory lymphocytes, monocytes and eosinophils, while increasing the number of neutrophils. The absolute numbers of T lymphocytes were proportionately more affected than the B lymphocytes. Craddock, et al., (1967) showed that cortisone treatment impaired leukocyte or macrophage migration in certain CMI responses. This was supported by the studies of Weston, et al., (1973) demonstrating the inhibitory effect of cortisol on the ability of macrophages to respond to the sensitized small (T) lymphocyte in cellular immunity.

Corticosteroids have been shown to inhibit T cell responses to mitogens and specific antigens when added to lymphocyte cultures (Heilman and Leicher, 1972). In other studies (McIntyre, et al., 1969; Stefani and Oester, 1967;
Tormey, *et al.*, 1967; Nowell, 1961), found that glucocorticoids added to lymphocyte cultures inhibited PHA induced transformation. The most effective inhibition occurred when corticosteroids were added before or simultaneously with a specific antigen or mitogen whereas no inhibition was observed when steroids were added after the lymphocytes had been stimulated (Bach, 1975). There is evidence that some cell populations or animal species as well as functions of T cells may be more resistant than others to corticosteroids and may require higher doses for inhibition to occur (Fauci, 1975; Heilman, 1972; Esteban, 1968).
SECTION III

HEMATOLOGICAL AND SERUM PROFILES OF FERRETS
Abstract

Normal hematological and serum profiles were determined in ferrets (Mustela putorius furo). The arithmetic mean, standard deviation, 90% confidence limits, 90% tolerance limits and simple linear correlation coefficients with respect to age or body weight were calculated on these variables.
Introduction. The recent development of technical and automated laboratory analysis of hematological and serum chemistry profiles has been invaluable in many animal investigations. Comparisons of these parameters between normal and disease suspected cases may provide insight or rapid confirmation of the clinical diagnosis. Some hematological and serum profiles (Rotenberg and Jorgensen, 1971) have been established for the mink (*Mustela vison*); however values for ferret parameters are few (Pyle, 1940). The main objective of this investigation was to determine hematological and serum chemistry profiles in clinically normal ferrets (*Mustela putorius furo*) and to determine the relationship of these parameters with respect to age or body weight.

Materials and Methods.

Animals. Thirteen fitch ferrets, *Mustela putorius furo* (Marshall Research Animals, North Rose, NY), four to eight months of age were used in these studies. All ferrets were vaccinated against canine distemper prior to shipment from Marshall Research Laboratories. The ferrets were housed in individual stainless steel cages (67 x 61 x 67 cm, Harford Metal Products, Aberdeen, MD) under artificial lighting (12 hours photoperiod), at 25° C, and provided with feed (Purina Main Meal Cat Chow, Checkerboard Square, MO) and water *ad libitum*. All animals were acclimatized for at least four weeks prior to initiation of the experiment. Ketamine
hydrochloride, (Bristol Laboratories, Syracuse, NY) administered intramuscularly (IM) at a dose of 12-15 mg/lb of body weight was used to induce anesthesia. Blood was drawn by cardiac or jugular vein puncture, using a 22 gauge, one inch needle. A maximum of eight ml of blood was drawn from the animals at approximately weekly intervals. (This corresponds to 20% of the total blood volume as reported by Hahn and Wester, 1969).

**Serum and hematological profiles.** Serum profiles were determined in the Sequential Multiple Analyzer (SMA-12, Technicon Instruments Corporation, Tarrytown, NY) using one ml serum samples. Complete blood counts (CBC's) were determined using one ml blood samples collected in one mg of K<sub>2</sub>EDTA. Both analyses were performed at the Clinical Pathology Laboratory, Department of Infectious Diseases, Kansas State University, Manhattan, KS.

**Hemolysis test.** Serum containing varying degrees of hemolysis (0-4+ and 10+) was analyzed for the serum chemistry profiles. Degrees of hemolysis in serum were simulated with serum prepared from frozen blood (hemolysis +10) and serum containing slight hemolysis (1+) as follows: hemolysis designated 2+, contained one part 10+ hemolyzed serum to 99 parts 1+ hemolyzed serum and hemolysis designated 4+, contained one part 10+ hemolyzed serum to 19 parts 1+ hemolyzed serum.

**Statistical analysis.** The arithmetic means, standard
deviations, 90% confidence limits, limits on 90% of the individual measurements in the designated population stated with 90% confidence, and simple linear correlation coefficients with respect to age or body weight were determined by standard statistical techniques.

**Results.** In samples with varying degrees of hemolysis, appreciable differences were observed only in total protein, albumin, sorbital dehydrogenase (SDH) and phosphorus (Table 1). Serum from frozen blood demonstrated increased levels of hemoglobin (Hb), potassium (K), serum glutamate pyruvate transaminase (SGPT), total protein and albumin; however chloride (Cl), sodium, (Na), carbon dioxide (CO₂), calcium (Ca), phosphorus (P) and glucose levels decreased.

Tables 2 and 3 present the arithmetic mean, standard deviation and 90% confidence intervals on the population mean of the hematological and serum chemistry profiles for female, male and castrated males. The mean value for white blood cells (WBC's) in females was observed to be less as compared to the males and castrated males; however, the mean values determined for the other parameters in the three groups of animals did not differ appreciably.

Table 4 lists the 90% tolerance limits on 90% of the population of the various hematological and serum parameters for the same groups of animals, assuming an essentially normal distribution of values.

There was a significant age related correlation with
neutrophils (+0.409, p = 0.07) and lymphocytes (-0.448, p = 0.04). Other hematological variables did not change significantly with respect to age or body weight (Table 5).

The serum enzymes, alkaline phosphatase, SGPT and SDH were significantly decreased, while glucose, blood urea nitrogen (BUN) and albumin were significantly elevated with age.

Sodium and chloride were positively correlated with body weight; however, phosphorus, CO₂ and glucose were negatively correlated. Significant correlations were not found between body weight and other serum parameters that were measured.

Discussion. The degree of hemolysis in the blood was directly proportional to the amount of hemoglobin present in the serum sample (Table 1). The hemolysis associated changes in the serum chemistry was a direct result of the damage to blood cell constituents, especially the erythrocytes. These results can be useful for interpreting serum profiles, since the degree of hemolysis and differences in the profiles may depend on the sample collection rather than on abnormal condition in the animal.

Pyle (1940) reported a neutrophil:lymphocyte ratio of 65:33 as compared to a ratio of 34:54 observed in our studies. In studies of minks a 38:53 ratio was reported by Rotenberg and Jorgensen (1971). Higher glucose and lower calcium levels were also demonstrated as compared to the mean values observed in the present study; however, there
were no appreciable differences in the other blood cell constituents. There appeared to be distinct similarities in the hematological and serum profile values of mink reported by Kennedy (1935a), Kubin and Mason (1948) and Rotenberg and Jorgensen (1971) and the ferrets in this study.

This data provides information on the normal hematological and serum profiles of ferrets which have not been studied in detail previously. This may be extremely useful with regard to the undertaking of scientific investigations in these species.
<table>
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<th>2+</th>
<th>4+</th>
<th>10+ (Frozen)</th>
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<td>120</td>
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<td>Serum Glutamate Pyruvate Transaminase (SGPT) mu/ml</td>
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<td>8.8</td>
<td>8.9</td>
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<td>110</td>
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<td>85</td>
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<td>Creatinine mg %</td>
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<tr>
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<td>( f(n_1) )</td>
<td>( m(n_1) )</td>
<td>( cm(n_1) )</td>
<td>( f )</td>
<td>( m )</td>
<td>( cm )</td>
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<td>49.12 (8)</td>
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<td>11.32 (5)</td>
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<td>1.95</td>
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<td>5.00 (8)</td>
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<td>2.39</td>
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<td>4.48</td>
<td>3.32</td>
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<td>Cl</td>
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<td>112.50 (3)</td>
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<td>1.90</td>
<td>9.10</td>
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<td>Na</td>
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<td>K</td>
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TABLE 2 (continued)

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<td>BUN</td>
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<td>Alk. P.</td>
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<td>SGPT</td>
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<td>T. P.</td>
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<td>Albumin</td>
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<tr>
<td>Ca</td>
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</tr>
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<td>P</td>
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<tr>
<td>Creatinine</td>
<td>0.36 (10)</td>
<td>0.40 (5)</td>
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f: females  
m: male  
cm: castrated male  
n₁: sample size
TABLE 3
HEMATOLOGICAL AND SERUM PROFILES IN FERrets

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<thead>
<tr>
<th>Variable</th>
<th>90% Confidence Intervals on Population Means</th>
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<td></td>
<td>f</td>
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<tr>
<td>PCV</td>
<td>47.20 - 49.56</td>
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<td>Hb</td>
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<td>WBC</td>
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<tr>
<td>Bands</td>
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<tr>
<td>Neutrophils</td>
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<td>Lymphocytes</td>
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<td>Monocytes</td>
<td>3.70 - 5.30</td>
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<tr>
<td>Eosinophils</td>
<td>2.26 - 4.74</td>
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<td>Basophils</td>
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<td>SDH</td>
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<td>T. P.</td>
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<td>Ca</td>
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<td>Glucose</td>
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<tr>
<td>Creatinine</td>
<td>0.33 - 0.39</td>
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</tbody>
</table>

f: females  
m: male  
cm: castrated male  
ni: sample size
<table>
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<th>m</th>
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If n=8, K=2.743 in $\bar{x} \pm Ks$; if n=10, K=2.535; if n=5, K=3.494; if n=3, K=5.847; and if n=9, K=2.626.
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<td>or</td>
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<sup>†</sup>, <sup>*</sup>, <sup>**</sup> indicate statistical significance at the 10, 5, 1% levels, respectively.

P = significance probability.

NS = Nonsignificant.
SECTION IV
IMMUNOLOGICAL STUDIES IN FERRETS INOCULATED WITH
INFECTIOUS BOVINE RHINOTRACHEITIS VIRUS
I. Humoral and Cell Mediated Immune Responses
Abstract

Ferrets inoculated with inactivated or modified live (ML) infectious bovine rhinotracheitis virus (IBRV) did not manifest clinical signs of IBRV infection. Statistical analyses were performed on the hematological and serum chemistry profiles for ferrets inoculated with formalin inactivated and modified live IBRV. Three days post intranasal (IN) vaccination with modified live virus (MLV), IBRV was isolated from the nasal washings of one of the three ferrets; however, it was not detected in the subsequent washings. Specific and nonspecific virus neutralizing factors were detected in the nasal washings only at the fourth week post-IN inoculation with inactivated IBRV. Serum neutralizing antibodies were detected three weeks following intramuscular (IM) injection of inactivated IBRV with Freund's incomplete adjuvant. Humoral antibodies increased following a second inoculation given IM. Maximum virus neutralization was observed between the 12th - 14th week. The MLV elicited low levels of serum antibodies that persisted from the third to seventh week post inoculation. The degree of cell mediated immunity was low and variable as observed by specific lymphocyte stimulation (³H-TdR incorporation) in whole blood cultures from ferrets inoculated with inactivated IBRV. Control and MLV inoculated ferrets did not demonstrate cell mediated immune responses to the IBRV antigen.
Introduction. The ubiquitous nature and the variety of clinical manifestations observed in infectious bovine rhinotracheitis, such as conjunctivitis (Abinanti and Plumer, 1961), rhinotracheitis, (Madin, York and McKercher, 1956), meningoencephalitis (French, 1962), vaginitis (McKercher, et al., 1959), abortion (Owens, Chows and Mollelo, 1964) and balanoposthitis (Kahrs, 1977), have resulted in the increasing economic concern to manage this disease in the United States and other parts of the world. The detrimental effects of IBRV infections have a direct bearing on the reproductive health and the milk and meat production of cattle. Inevitably this disease affects cattle of all ages.

Investigations evaluating the various types of IBRV vaccines available and the different routes of inoculation have provided much information on the levels of immunity conferred; however, the complexity of the immune system has also led to considerable obscurities. It has been reported that IBRV infections occurred in the presence of humoral antibodies (Snowden, 1965) implicating the importance of the cell mediated immune (CMI) system.

The ferret, *Mustela putorius furo*, has been found to simulate a pathological response with influenza virus similar to man and has been used to evaluate the efficacy of the virus vaccines. Porter, Larsen and Cox (1975) reported the isolation of IBRV in ferrets. In more recent studies, (NADC, 1977) ferrets have been under investigation to
evaluate the immune response to IBRV inoculations.

The main objective of this investigation is to study the immune response of ferrets inoculated with IBRV vaccines.

**Materials and Methods.**

**Animals.** Ferrets four to eight months old were used in these studies. Rectal temperatures and body weights were recorded prior to and at specific intervals post inoculation with IBRV. Anesthesia was induced with ketamine hydrochloride, (Bristol Laboratories, Syracuse, NY), administered intramuscularly (IM) at a dose of 12-15 mg/lb. of body weight. Blood was drawn by cardiac or jugular vein puncture using a 22 guage, one inch needle.

**Cell cultures.** Monolayer cultures of Madin Darbin bovine kidney cells, obtained from the American Type Culture Collection (ATCC) Laboratories (Rockville, MD) were grown in prescription bottles (32 ounces) or tissue culture dishes (60 x 15 mm; Falcon, Oxnard, CA) and incubated at 37°C in a humidified incubator. The cells were maintained in minimal essential medium (MEM), Eagles (Grand Island Biological Co., Grand Island, NY), supplemented with 10% heat inactivated horse serum (KC Biological Inc., Lenexa, KS), 0.1% sodium bicarbonate (NaHCO₃) and an antibiotic mixture of penicillin, 100 units/ml., streptomycin, 1 ng/ml and kanamycin, 5 ng/ml (PSK).

**Propagation and assay of viruses.** Infectious bovine
rhinotracheitis virus, the Los Angeles (LA) strain and bovine
virus diarrhea virus (BVDV), the National Animal Disease
Laboratory (NADL) strain used in these studies were obtained
from ATCC Laboratories. Monolayer cultures (70-90% confluent)
of MDBK cells were infected with IBRV or BVDV virus with a
multiplicity of 3-5 plaque forming units (PFU)/cell and main-
tained in Medium 199, (Grand Island Biological Co.) at 37°C.
The cultures were harvested by scraping with a rubber police-
man when maximum cytopathic effect (CPE) was observed (ap-
proximately 48-72 hours). The cell-virus suspension was
frozen and thawed twice to release virus from the cells, then
centrifuged at 2,000 g for 30 minutes, at 5°C. The super-
natant (IBRV) was stored in aliquots at -80°C. The super-
natant containing BVDV was ultracentrifuged (Beckman L2-65B,
Beckman Instruments, Inc., Palo Alto, CA), at 100,000 g for
five hours at 5°C. The pellet was resuspended in Medium 199
and stored in aliquots at -80°C. All viruses were assayed
by the procedure for animal viruses, described by Dulbecco,
(1952) with some modifications. Briefly, serial log dilutions
of virus were adsorbed for one hour onto MDBK cell mono-
layers in dishes (0.2 ml/dish). Seven mls of an agar (2%
purified) and media (2 X MEM) mixture supplemented with 5%
HS were overlaid onto the cultures and incubated (three days
for IBRV and four days for BVDV). Plaques were observed 24
hours after a second overlay of agar and media mixture con-
taining 0.015% neutral red stain.
Ultraviolet inactivation of viruses. One ml of IERV or BVDV (1 x 10⁷ PFU/ml) was UV irradiated for 4.5 and five minutes, respectively, in a petri dish (60 x 15 mm) at a distance of 10 cm from a UV germicidal sterilamp (630T8, 30 watt, Westinghouse Electric Co., Bloomfield, NJ). The virus was completely inactivated as determined by the plaque assay method.

Preparation of inactivated IERV vaccine. One ml of IERV (5 x 10⁸ PFU/ml) was treated with formalin at a final concentration of 1:1000 (40% formaldehyde solution, Fisher Scientific Co., Fairlawn, NJ) for six hours at 25⁰ C, then overnight at 4⁰ C. The virus was dialyzed in 1/4" dialyzing tubing against three changes of cold PBS at 4⁰ C over a 24 hour period. Inactivated virus as determined by the plaque assay method was stored in aliquots at -80⁰ C.

Serum and hematological profiles. Serum profiles were determined in the Sequential Multiple Analyzer (SMA-12, Technicon Instruments Corp., Tarrytown, NY) using one ml serum samples. Complete blood counts (CBC) were determined using one ml of blood. Both analyses were performed prior to and at specific intervals post inoculation of IERV at the Clinical Pathology Laboratory, Department of Infectious Diseases, Kansas State University, Manhattan, KS.

Inoculations of ferrets. The absence of IERV serum neutralizing antibodies was confirmed by the plaque reduction method prior to inoculation with IERV vaccines.
Group A: Five ferrets were inoculated intranasally (IN) with $2.5 \times 10^7$ PFU/0.1 ml of formalin inactivated IBRV. Five control ferrets were sham inoculated with formalized cell lysate (from MDEK cells). Formalin inactivated IBRV, $5 \times 10^7$ PFU/0.2 ml was homogenized (Multi-Churn Homogenizer) with an equal volume of Freund's Incomplete Adjuvant (Grand Island Biological Co.) and administered IM into animals three weeks post IN inoculation. A booster of the homogenate was given into the opposite quarter three weeks later. Controls were inoculated with a homogenous mixture of the cell lysate and Freund's incomplete adjuvant.

Group B: Three ferrets were inoculated IM with $1.6 \times 10^4$ PFU/0.2 ml of modified live IBRV (Delvine®, Dellen Laboratories, Omaha, NB). One control animal was inoculated with the diluent provided for rehydration of the antigen.

Group C: Three ferrets were etherized and inoculated intranasally with 0.1 of ML IBRV vaccine containing $2 \times 10^5$ PFU, (Nasalgen I, Monovalent, Serial 511) kindly provided by Dr. J. D. Todd, Jensen-Salsbery Laboratories, Kansas City, MO. One control animal was inoculated with the diluent provided for rehydration of the antigen.

Collection of nasal washings. Ferrets were tranquilized with Ketamine hydrochloride (5 mg/lb of body weight). Three mls of PBS were introduced intranasally with a 3 cc. syringe (without a needle). Sneezing reflexes resulted and the contents of each nostril were collected in a sterile petri dish.
(100 x 20 mm, Falcon). Approximately 2.5 ml was recovered and immediately transferred into tubes in an ice bath containing antibiotics supplemented with 0.5 µg/ml of Fungizone® (E. R. Squibb and Sons, NY).

The nasal washes, 0.3 ml from animals in group C were passaged thrice on MDBK monolayers and observed for CPE.

Detection of serum neutralizing antibodies by the plaque reduction method. The presence of serum neutralizing antibodies against IBRV in heat inactivated ferret serum (56°C for 30 minutes) was determined by the plaque reduction method. Briefly, the serum samples were mixed with equal volumes (0.3 ml) of IBRV, 5 x 10²-1 x 10³ PFU and incubated for one hour at 25°C. The samples were plaque assayed to determine the percent recovery of the virus.

Detection of specific and nonspecific factors against viruses in nasal washings. The absence of specific and nonspecific neutralizing factors in ferret nasal washings was confirmed prior to inoculations with IBRV vaccines. The presence of specific and nonspecific virus neutralizing factors was determined by the plaque reduction method post virus inoculation. The samples were mixed with equal volumes (0.3 ml) of either IBRV or BVDV, 5 x 10² - 1 x 10³ PFU and incubated for one hour at 25°C. The samples were plaque assayed to determine the percent recovery of the virus.
**Blastogenesis studies in whole blood cultures from control and vaccinated ferrets.** The method of whole blood cultures used by Gerber and Brown (1974) was adopted with modifications. Heparinized whole blood, 0.1 ml (5 units/ml of blood, Panheprin, Abbott Laboratories, North Chicago, IL) was added to duplicate tubes (17 x 100 mm, Falcon) containing RPMI 1640 (Roswell Park Memorial Institute, Grand Island Biological Co., Grand Island, NY) and 10% heat inactivated fetal calf serum, free of IBRV and BVDV antibodies (KC Biological Co.). UV inactivated virus (2.0 x 10^5 PFU/0.2 ml) was added to the cultures. The cultures were incubated for 108 hours then exposed to 0.5 uCi/ml of ^3^H-TdR (specific activity 5 Ci/mmole, New England Nuclear). Cultures were removed from the CO\_2 incubator after 12 hours of exposure to label and placed in an ice bath to terminate cellular activity. The tubes were centrifuged at 500 g for 15 minutes in a refrigerated centrifuge. The supernatant was carefully aspirated, cultures gently vortexed and treated two to three times with six ml of cold distilled (not deionized) water for 30 seconds each time to lyse erythrocytes. Isotonicity was restored with six ml of cold 1.7% NaCl solution. The remaining cells (mixture of WBC's in a one ml volume) were precipitated with five mls of cold 5% trichloroacetic acid (TCA) and kept in an ice bath for 30 minutes. Acid insoluble, precipitated macromolecules were collected on glass filters (GF/C, 2.4 cm. Whatman, England) using a sampling manifold (Millipore Corporation, Bedford, MA). The tubes were washed twice with
five ml of cold 5% TCA and the contents poured onto the filters. The filters were finally placed into glass scintillation vials containing 10 ml of scintillation fluid. Samples were counted for 20 minutes by a Scintillation Spectrometer (Packard Tri-Carb, Model 3002, Downers Grove, IL).

Results.

Clinical observations. Clinical signs of IBRV infection were not observed in ferrets inoculated with inactivated or modified live IBRV; however, local inflammation persisted for approximately two weeks at the site of inoculation of the adjuvated vaccine. The mean rectal temperatures of each group remained within the normal range (101.0-104.0°F) as reported for ferrets by Pyle (1940). The behavior and appetites appeared normal as compared to pre-inoculation observations. Control and inactivated IBRV inoculated ferrets demonstrated a plateau in weight between 28 and 44 weeks of age, followed by a decrease in weight for four to five weeks (Figure 1). Although the females experienced a greater weight loss than the males, both reached a plateau by 50 weeks of age. The maximum weight observed for females and males was 1.2 Kg and 1.55 Kg, respectively.

Hematological and serum profiles. Table 1 presents the statistical significance of changes in the hematological and serum chemistry profiles between pre- and post-inoculation values for four groups of ferrets. Ferrets inoculated with formalin inactivated IBRV with Freund's Incomplete Adjuvant
demonstrated significant changes \( (P = 0.05 \text{ or } 0.01) \) in chloride (Cl), potassium (K), carbon dioxide (\( \text{CO}_2 \)), albumin, calcium (Ca), phosphorus (P), and bands. Animals inoculated with modified-live IBRV, given IM had significant differences \( (P = 0.05 \text{ or } 0.01) \) between pre- and post-inoculation means in Cl, sodium (Na), blood urea nitrogen (BUN), creatinine, packed cell volume (PCV) and hemoglobin (Hb). Statistically significant differences \( (P = 0.05 \text{ or } 0.01) \) in pre- and post-inoculation means from ferrets inoculated with modified live IBRV given IN were detected in Cl, BUN, alkaline phosphatase, glucose, and creatinine. The control ferret group, inoculated with incomplete adjuvant only, demonstrated significant pre- and post-inoculation changes \( (P = 0.05, 0.01) \) in total protein, P, white blood cell (WBC) and monocytes. Albumin, BUN, SDH, creatinine, PCV and bands were significant among the treatments (Table 2).

**Virus isolation from nasal washings.** Two of the three ferrets inoculated with modified live IN IBRV vaccine had a red hue in the nasal washings three days post-inoculation; however, subsequent samples were similar to the control washings. Virus was not detected in the nasal washings from ferrets inoculated IN with inactivated IBRV.

Virus \( (3 \times 10^3 \text{ PFU/ml}) \) was recovered on day three post-inoculation in one of the three nasal washings from ferrets inoculated IN with MLV. This sample produced CPE in the first passage onto MDBK cells. Virus was not demonstrated
in subsequent samples.

Detection of nasal antibodies. The presence of specific and nonspecific virus neutralizing factors was detected four weeks post IN inoculation in samples taken from ferrets vaccinated with inactivated IBRV (Figure 2). These factors were not found in following samples.

Detection of humoral antibodies. Serum neutralizing antibodies were detected three weeks following IM injection of inactivated IBRV vaccine (six weeks post IN inoculation). Humoral antibodies were detected as indicated by a 40% recovery of IBRV and increased following a second IM booster (Figure 3). Maximum antibody production (0% virus recovery) occurred between the 12th and 14th week and declined thereafter to an 80% virus recovery rate by the 20th week at which time the experiment was terminated.

Low and variable levels of serum neutralizing antibodies were detected in ferrets following the inoculation of IN and IM modified live vaccines (Figure 4). The IM vaccine elicited antibodies two weeks post inoculation and persisted at low levels until week seven. The IN vaccine produced a weak immune response at week one and three post-inoculation.

The stimulation of lymphocytes by specific antigen. The synthesis of DNA as measured by $^3$H-TdR incorporation in lymphocytes from ferrets vaccinated with inactivated adjuvated IBR vaccine was detected at variable levels three weeks
post-inoculation (Table 3). A maximum stimulation index of 27.7 was observed 11 weeks post-inoculation.

Ferrets inoculated with modified live virus did not show the incorporation of $^3$H-TdR in lymphocyte cultures (Table 4).

Lymphocytes from the control animals did not incorporate appreciable amounts of $^3$H-TdR (SI = 0-4.5). Relatively low levels of non-specific stimulation by lymphocytes from both control and vaccinated animals was observed (S.I. = 1.3-6.2) in cultures supplemented with UV inactivated IBRV beginning the eighth week post IN vaccination of inactivated IBRV. This was observed till the termination of the experiment. Whole blood cultures from ferrets inoculated with MLV vaccine were not tested for non-specific stimulation with UV-BVDV.

**Discussion.** In this study, the absence of clinical IBRV infection in ferrets, confirms the observations made by Porter, Larsen and Cox, (1975). They isolated IBRV from apparently healthy ferrets that were not manifesting signs of IBRV infection. It is not clear from their studies, however, that IBRV once introduced can infect and replicate in ferret tissues. In studies by Shump and Shump (1978), the weights of ferrets plateaued at approximately 20 weeks of age as opposed to 28 weeks observed in our studies. The decrease in weight observed at 44 weeks (after inoculations) may have been due to numerous blood samplings with greater stress and weight loss occurring in the females. Ferrets
reached higher weight levels, 1.55 Kg in males and 1.2 Kg in females as compared to 1.5 Kg in males and 0.9 Kg in females reported by Shump and Shump (1978).

The pre- and post-inoculation mean differences observed in the hematological serum chemistry profiles between the ferrets inoculated with formalin inactivated IBRV with Freund's Incomplete Adjuvant and the controls inoculated only with incomplete adjuvant are significantly different (P = 0.10) for albumin, creatinine and bands (Table 1 and 2). Ferrets inoculated with MLV were significantly different (P = 0.10) for albumin and PCV with no significant changes detected in the other 21 profiles, suggesting little differences between the routes of inoculation.

The absence of significance for the other 17 variables among the treatments (Table 1 and 2), indicates that the treatments A,B,C and D did not have appreciable effects on those variables during the pre-inoculation to the post-inoculation period.

The isolation of virus three days post IN vaccination with MLV from one of the three ferrets was probably due to the presence of residual virus from the inoculum, rather than replicated virus particles in the nasal mucosa. Subsequent nasal washings did not yield any virus. The red coloration in two of the three nasal washings may have been caused by a local inflammatory reaction to the antigen.
Inflammation resulting in the production of local secretory antibodies and interferon in the nasal mucosa, as demonstrated in cattle vaccinated IN with MLV (Todd, Volenec and Paton, 1972) would prevent virus replication. The reason CPE was not observed beyond the first passage of the sample on MDBK cells is obscure. It is likely that the ferret nasal washings containing the products of an inflammatory response were cytotoxic to MDBK cells in the initial passage and was diluted with subsequent passages. The inability of IBRV to persist in the ferret nasal mucosa may have resulted in the low virus recovery and the subsequent loss of the virus from following samples.

The specific and non-specific virus inactivating factors detected in the nasal washings (Figure 2) from ferrets inoculated IN with inactivated IBRV may demonstrate the presence of local immunity.

The duration and levels of serum neutralizing antibodies in ferrets inoculated with inactivated IBRV with Freund's Incomplete Adjuvant exceeded the duration and levels of antibody detected in ferrets inoculated IN or IM with MLV vaccines (Figures 3 and 4). Since it is likely that the MLV did not replicate in the ferret, the titer of the antigen present in vivo may not have been sufficient to induce a comparable humoral response as observed in the ferrets inoculated with inactivated adjuvated vaccine. In studies by Porter, Larsen and Cox, (1975), the presence of serum antibodies was detected three weeks post-inoculation of IBRV
(10^6 PFU), given intraperitoneally in mink. Inactivated influenza virus vaccines with Freund's Incomplete Adjuvant have been found to elicit serum antibodies in ferrets (Potter, Jennings and McLaren, 1973). It is possible that the activity of the vaccine with adjuvant involved one or a combination of the following phenomena: (1) the establishment of the antigen in a persistent form at the site of injection, allowing the gradual but continuous release of antigen for stimulating antibody production, (2) the provision of a vehicle for the transport of emulsified antigen to the lymphatic system, and (3) the formation of an inflammatory response and the accumulation of mononuclear cells which elicit antibody production at local and distant sites (Hilleman, 1967). The later explains the inflammatory response observed in the ferrets post IM vaccination of inactivated IBRV with adjuvant (Figure 2). It is probable that the presence of adjuvant in the inactivated vaccine was an important factor for eliciting an immune response in ferrets. Hobson, (1973) and Woodhour, et al. (1969), demonstrated that vaccine preparations with adjuvant produced higher levels of antibody than equal doses of aqueous vaccine. They also reported that a single dose of vaccine containing adjuvated virus, produced higher levels of antibody than two doses containing a greater amount of antigen in an aqueous vaccine preparation. In influenza virus studies, only ferrets inoculated with inactivated virus vaccine in Freund's Incomplete Adjuvant produced serum antibodies, as compared to ferrets inoculated with an equal dose
of virus without adjuvant (Potter, Jennings and McLaren, 1973). Future studies using increasing doses of IBRV with a constant amount of adjuvant would provide more conclusive evidence for comparing the degrees of immunity conferred by an adjuvated virus preparation.

Cell mediated immune responses as indicated by lymphocyte transformation and the incorporation of $^{3}$H-TdR in the presence of a specific antigen were only detected in ferrets inoculated with inactivated IBRV with adjuvant (Table 3). The degree of specific stimulation demonstrated by the lymphocytes from inactivated IBRV vaccinated ferrets was higher than the stimulation observed in cultures from control ferrets. However, both control and inactivated IBRV vaccinated ferret cultures demonstrated low degrees of non-specific stimulation. The reasons for this type of stimulation are not clear. It is known that adjuvants can modify the expression of antigenic determinants, alter the ratio of various antibody classes and change the balance between antibody mediated and cell mediated immunity (White, 1967). From these observations the mechanisms resulting in the superior levels of immunity elicited by the inactivated vaccine with adjuvant as compared to the aqueous MLV can be explained.

In this investigation the humoral and cell mediated immune responses of ferrets to inactivated and ML IBRV was studied. Since ferrets do not simulate bovines with regard to IBRV pathogenesis (Kahrs, 1977; Porter, Larsen and Cox, 1975) and immunity to inactivated and ML IBRV vaccines
(present study) it is questionable whether these animals may be satisfactory models for studying IBRV infections or evaluating IBRV vaccines.
Figure 3. Serum neutralizing antibodies in ferrets inoculated with inactivated virus. Two samples, one each from a control and an experimental ferret, were tested at specified intervals. The serum samples were incubated for one hour at 25°C with an equal volume of IBRV, $5 \times 10^2 - 1 \times 10^3$ PFU/0.3 ml. The samples were plaque assayed to determine the percent recovery of the virus.
Figure 2. Specific and nonspecific virus neutralizing factors in nasal washings. Two samples, one each from a control and an experimental ferret, were collected at specified intervals. The nasal washings from ferrets inoculated IN with inactivated virus were incubated for one hour at 25°C with an equal volume of IBR and BVD virus, 5 x 10^2 - 1 x 10^3 PFU/0.3 ml. The samples were plaque assayed to determine the percent recovery of virus: •——• Specific Factors; △——△ Non-specific Factors.
Weights of male and female ferrets. Weights of animals before inoculation from five castrated males and five females were recorded. At 35 weeks of age, four males and one female were inoculated with inactivated IBRV given IN, while one male and two females were sham inoculated. At 38 and 41 weeks of age, the experimental animals were inoculated IM with a homogenous mixture of inactivated IBRV and Freund's Incomplete Adjuvant. The controls received the mixture without virus:

- Castrated Males Pre-inoculation; -...- Castrated Male Control; --- Castrated Male Experimental;

- Female Pre-inoculation; -...- Female Control;

- Female Experimental.
## TABLE 1

STATISTICALLY SIGNIFICANT CHANGES BETWEEN PRE- AND POST-INOCULATION MEANS IN FERRET HEMATOLOGICAL AND SERUM CHEMISTRY PROFILES

<table>
<thead>
<tr>
<th>Variables Measured</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A Inactivated IBRV and Freund's Incomplete Adjuvant n = 10-13</td>
</tr>
<tr>
<td>-----------------------------------------</td>
<td>---------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Packed Cell Volume (PCV) %</td>
<td>2.36 +</td>
</tr>
<tr>
<td>Hemoglobin gm %</td>
<td>-0.0818 NS</td>
</tr>
<tr>
<td>White Blood Cell (WBC) Cu mm</td>
<td>0.755 NS</td>
</tr>
<tr>
<td>Bands/Cu mm</td>
<td>2.55 *</td>
</tr>
<tr>
<td>Neutrophil/Cu mm</td>
<td>0.818 NS</td>
</tr>
<tr>
<td>Lymphocyte/Cu mm</td>
<td>-6.64 NS</td>
</tr>
<tr>
<td>Monocyte/Cu mm</td>
<td>-1.60 +</td>
</tr>
<tr>
<td>Eosinophil/Cu mm</td>
<td>0.455 NS</td>
</tr>
<tr>
<td>Basophil/Cu mm</td>
<td>-0.0909 NS</td>
</tr>
<tr>
<td>Sorbitol Dehydrogenase (SDH) I.U.</td>
<td>0.342 NS</td>
</tr>
<tr>
<td>Chloride (Cl) meq/L</td>
<td>-4.82 **</td>
</tr>
<tr>
<td>Sodium (Na) meq/L</td>
<td>1.29 NS</td>
</tr>
<tr>
<td>Potassium (K) meq/L</td>
<td>0.569 **</td>
</tr>
<tr>
<td>Carbon Dioxide (CO2) meq/L</td>
<td>-4.22 **</td>
</tr>
<tr>
<td>Blood Urea Nitrogen (BUN) mg %</td>
<td>-2.16 NS</td>
</tr>
<tr>
<td>Alkaline Phosphatase mu/ml</td>
<td>7.23 NS</td>
</tr>
<tr>
<td>Serum Glutamate Pyruvate Transaminase (SGPT) mu/ml</td>
<td>16.75 NS</td>
</tr>
<tr>
<td>Total Protein (T.P.) gm %</td>
<td>-0.100 NS</td>
</tr>
<tr>
<td>Albumin gm %</td>
<td>0.315 **</td>
</tr>
</tbody>
</table>
TABLE 1 (continued)

<table>
<thead>
<tr>
<th>Variables Measured</th>
<th>Treatment</th>
<th>A Inactivated IBRV and Freund's Incomplete Adjuvant n = 10-13</th>
<th>B Modified Life IBRV n = 3-5</th>
<th>C Modified Live IBRV n = 3-5</th>
<th>D Incomplete Adjuvant Only n = 3-8</th>
<th>Among Treatments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcium (Ca) mg %</td>
<td>0.923 **</td>
<td>-0.160 NS</td>
<td>0.0857 NS</td>
<td>0.943 NS</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phosphorus (P) mg %</td>
<td>0.854 **</td>
<td>0.460 NS</td>
<td>-0.171 NS</td>
<td>0.871 *</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glucose mg %</td>
<td>-10.92 +</td>
<td>-8.00 NS</td>
<td>-25.29 *</td>
<td>8.86 NS</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Creatinine mg %</td>
<td>-0.0154 NS</td>
<td>-0.260 *</td>
<td>-0.229 **</td>
<td>0.0571 +</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

NS, +, * and ** = statistically significant at nonsignificant, 10%, 5% and 1% levels, respectively.
TABLE 2
LEAST SIGNIFICANT DIFFERENCES (LSD)
BETWEEN PRE- AND POST-TREATMENT MEANS

<table>
<thead>
<tr>
<th>Variable Measured</th>
<th>Significant Treatments</th>
</tr>
</thead>
<tbody>
<tr>
<td>SDH</td>
<td>C&amp;B, B&amp;A, B&amp;D, C&amp;A</td>
</tr>
<tr>
<td></td>
<td>*</td>
</tr>
<tr>
<td></td>
<td>**</td>
</tr>
<tr>
<td></td>
<td>*</td>
</tr>
<tr>
<td></td>
<td>*</td>
</tr>
<tr>
<td>PVC</td>
<td>C&amp;B, D&amp;B, A&amp;B</td>
</tr>
<tr>
<td></td>
<td>*</td>
</tr>
</tbody>
</table>

1. Group A: Inactivated Virus and Adjuvant
   Group B: Modified Virus (IM)
   Group C: Modified Virus (IN)
   Group D: Adjuvant Only

2. +, *, ** = statistically significant at 10%, 5% and 1% levels, respectively.
### Table 3

**Stimulation Indices**

**Ferrets inoculated with formalin inactivated IBRV**

<table>
<thead>
<tr>
<th>WEEKS POST-INOCULATION</th>
<th>CONTROL (^B) S.I.</th>
<th>EXPERIMENTAL (^B) S.I.</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>6</td>
<td>0.0</td>
<td>6.0</td>
</tr>
<tr>
<td>8</td>
<td>0.0</td>
<td>1.1</td>
</tr>
<tr>
<td>10</td>
<td>0.0</td>
<td>1.3</td>
</tr>
<tr>
<td>11</td>
<td>1.3</td>
<td>27.7</td>
</tr>
<tr>
<td>12</td>
<td>1.9</td>
<td>2.9</td>
</tr>
<tr>
<td>13</td>
<td>0.0</td>
<td>10.4</td>
</tr>
<tr>
<td>14</td>
<td>3.6</td>
<td>1.1</td>
</tr>
</tbody>
</table>

\(^{A}\) Stimulation Index (S.I.) = \(\frac{\text{DPM (Cells + Antigen)}}{\text{DPM Cells}}\)

\(^{B}\) Two ferrets, one control and one experimental, were bled each time.
TABLE 4
STIMULATION INDICES

FERRETS INOCULATED WITH MODIFIED LIVE IBRV

<table>
<thead>
<tr>
<th>Modified Live IBRV</th>
<th>Control S.I.</th>
<th>Experimental S.I.</th>
</tr>
</thead>
<tbody>
<tr>
<td>I.N.</td>
<td>4.5</td>
<td>0</td>
</tr>
<tr>
<td>10 Weeks</td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>Post-Inoculation</td>
<td></td>
<td>&lt;1</td>
</tr>
<tr>
<td>I.M.</td>
<td>0.0</td>
<td>0</td>
</tr>
<tr>
<td>8 Weeks</td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>Post-Inoculation</td>
<td></td>
<td>0</td>
</tr>
</tbody>
</table>

\[ ^a \text{Stimulation Index (S.I.)} = \frac{DPM (Cells + Antigen)}{DPM (Cells)} \]
SECTION V
IMMUNOLOGICAL STUDIES IN FERrets INOCULATED WITH
INFECTIONous BOVine RHINOTRACHEITIS VIRUS
II. Effect of Phytohemagglutinin and Cortisol on Leukocytes
Abstract

Phytohemagglutinin (PHA) at a concentration of 5 ug/ml demonstrated maximum stimulation of ferret lymphocytes in whole blood cultures. The optimal effect of PHA on deoxy-ribonucleic acid (DNA) synthesis was observed between 72 and 168 hours after incubation of the cultures. Cultures stimulated with PHA incorporated maximum label when they were exposed to tritiated thymidine ($^3$H-TdR) for 12 hours prior to harvest.

Viability of leukocytes after incubation for 96-144 hours was approximately 10 fold greater in cultures with PHA as compared to cultures without PHA or containing ultraviolet irradiated infectious bovine rhinotracheitis virus (UV-IBRV).

Cortisol in physiological concentrations inhibited the viability of PHA stimulated lymphocytes as indicated by an appreciable decrease in cell count and incorporation of $^3$H-TdR. The cell mediated immune response was also depressed in UV-IBR stimulated cultures. The presence of increasing concentration of cortisol (0-5000 mg/ml) did not affect the adsorption of $^3$H-IBRV to ferret leukocytes.
Introduction. Infectious bovine rhinotracheitis is capable of producing a latent infection and a variety of clinical manifestations (Kahrs, 1977; Owens, Chow and Mollelo, 1964; French, 1962; Abinanti and Plumer, 1961; McKercher, et al., 1959; Madin, York and McKercher, 1956). Recrudescence or activation of latent IBR infections can occur spontaneously during periods of stress when physiological hormone levels are altered or with the administration of corticosteroids (Davies and Duncan, 1974; Davies and Carmicheal, 1973).

It has been demonstrated that IBRV latency and spontaneous recrudescence can occur in the presence of humoral antibodies (Sheffy and Rodman, 1973). As a result there has been increasing interest in the role of the cell mediated immune responses in an attempt to understand and control IBRV infections. The primary factors of the cellular immune system are the thymus (T) dependent lymphocytes. There are various techniques for measuring T cell stimulation or blastogenesis to a mitogen or specific antigen. The method reported by Sorensen, Anderson, and Giese (1969) measures the incorporation of radioactively labelled DNA precursors as an indication of T-lymphocyte blastogenesis and is used as an index for measuring CMI responses.

It has been found that adrenal cortical hormones by causing cytotoxic effects and altering cellular metabolism and activity in the cellular components of the immune system, suppress the CMI responses to either mitogen or specific
antigen. In vitro studies have illustrated the inhibition of blastogenesis in lymphocyte cultures supplemented with cortisol before or simultaneously with the stimulus (Fauci and Dale, 1974; Tormey, Fudenberg and Kamin, 1967; Stevens and Dougherty, 1964; Nowell, 1961).

A better understanding of the complex mechanisms of the immune response in an animal to an antigen or a mitogen and the suppression observed in lymphocytes with cortisol would provide insight for controlling latency and the recrudescence of IBRV infections. The objective of this investigation is to study the effect of cortisol on lymphocytes in phytohemagglutinin and antigen supplemented ferret whole blood cultures.

Materials and Methods.
Animals. Fitch and English ferrets, Mustela putorius furo (Marshall Research Animals, North Rose, NY) were used in these studies. The animals were vaccinated with infectious bovine rhinotracheitis virus (LA strain) as previously described in Section IV.

Preparation of ultraviolet inactivated virus for lymphocyte stimulation. Virus (IBRV) for in vitro stimulation of whole blood cultures was prepared as described in Section IV.

Determining the time course of DNA synthesis. The method of whole blood cultures used by Gerber and Brown (1974) was adopted with modification as previously described. A stock
solution of phytohemagglutinin (PHA-V; Sigma Chemical Co., St. Louis, MO) was prepared in RPMi-1640 (Grand Island Biological Co., Grand Island, NY). Whole blood cultures containing 5 μg/ml of PHA were incubated for varying time periods up to 168 and 240 hours. The cultures were exposed to 0.5 uCi/ml of tridiated thymidine ($^3$H-TdR; specific activity 5 Ci/m mole, New England, Boston, MA) in medium, 12 hours prior to harvest each time. Samples were harvested at various intervals. The cultures were treated with equal volumes of distilled water and a 1.7% NaCl solution, processed and counted on glass filters as previously described. A set of unlabelled samples were treated in a similar manner and the leukocyte viability determined by the trypan blue stain (0.04%, in normal saline, Grand Island Biological Co.) exclusion method. The fold increase over the zero hour control was calculated.

**The effect of PHA concentration on DNA synthesis.** Whole blood cultures supplemented with different concentrations of PHA were incubated and labelled with $^3$H-TdR for 24 hours prior to harvest at 120 hours. The cultures were harvested, processed and DPM's recorded as previously described.

**Determining the time of exposure for maximum $^3$H-TdR incorporation into lymphocytes.** Whole blood cultures containing 5 μg/ml of PHA were incubated. Beginning at 96 hours of incubation, $^3$H-TdR was added to different sets of cultures at four hour intervals. All cultures were harvested at 120
hours, processed and DPM's determined.

**Effect of UV inactivated IBRV and PHA on leukocyte viability.** Whole blood cultures from control and experimental ferrets were supplemented with UV-IBRV \( (2.0 \times 10^5 \text{ PFU}/0.2 \text{ ml}) \), PHA \( (5 \text{ ug/ml or UV-IBRV and PHA} \) and incubated between 0 and 240 hours. Percent cell viability over the zero hour control was determined at harvest by the method as described.

**Preparation of \(^3\text{H}-\text{thymidine labelled IBRV.}** Monolayers of MDBK cells (ATCC Laboratories, Rockville, MD) grown in tissue culture dishes \( (100 \times 20 \text{ mm, Falcon}) \), were infected with IBRV at a multiplicity of 10 PFU/cell and maintained in medium 199. Radioactive medium containing 0.25 uCi/ml of \(^3\text{H}-\text{thymidine (}^{3}\text{H-TdR, specific activity 90 Ci/m mole, New England Nuclear, Boston, MA) was added to monolayers after virus adsorption. Cultures were harvested when 90\% of the cells showed CPE (approximately 40 hours). The cells were scraped from the dishes and the cell-virus suspension was collected and partially purified as follows. After low speed centrifugation, the supernatant containing virus was ultracentrifuged at 100,000 g for five hours. The supernatant was discarded and the virus pellet was resuspended in cold medium 199 and treated with DNase at a final concentration of 10 ug/ml. The mixture was incubated for 30 minutes at 25\( ^\circ \text{ C, and ultracentrifuged for 1.5 hours at 200,000 g. The pellet was washed in cold media, resuspended and dialyzed (1/4" tubing, size one, Fisher Scientific,
Pittsburgh, PA) against MEM in an ice bath for three hours and stored at -80° C.

Effect of Cortisol on Lymphocytes from Control and IBRV Vaccinated Ferrets.

(a) Inhibition of $^3$H-TdR incorporation in ferret lymphocytes.

A stock solution of cortisol (Calbiochem, San Diego, CA) was prepared in an acetone-methanol (1:1) mixture and diluted in RPMI 1640 to various concentrations (0 ng-50,000 ng). Different concentrations of cortisol were added to whole blood cultures containing either 5 ug PHA or UV inactivated IBR (2.0 x $10^5$ PFU/0.2 ml). The cultures were exposed to 0.5 uCi/ml of $^3$HTdR at 108 hours. Samples were removed from a CO$_2$ incubator after 12 hours of exposure to the label and placed in an ice bath. The cultures were processed and counted as previously described. The percent DPM's was calculated from the PHA control.

(b) The effect of cortisol on the viability of lymphocytes.

The viability of lymphocytes isolated from whole blood cultures incubated 120 hours, containing or lacking 5 ug PHA or containing PHA and varying concentrations of cortisol (0 ng-50,000 ng) was determined by the method of trypan blue stain exclusion.

(c) Competition of cortisol and $^3$H-IBR for leukocyte receptor sites.
A suspension of leukocytes (0.1 ml) isolated from heparinized blood, was added to duplicate sets of tubes containing a mixture of varying concentrations of cortisol (0 ng-5,000 ng) and a constant amount of $^3$H-IBRV ($2.0 \times 10^{14}$ PFU containing 3800 DPM. The mixture was incubated for four hours, gently mixing every hour. The cultures were placed in an ice bath and washed thrice with cold sterile PBS to remove the unadsorbed virus. Cells were then pelleted at 900 g for 10 minutes. The final pellet was resuspended in two ml of cold PBS and 0.2 ml of the sample was removed for plaquing. The remaining suspension was collected on glass filters and counted as previously described.

Results.

The effect of PHA concentrations on DNA synthesis. Phytohemagglutinin at a concentration of 5 µg/ml stimulated lymphocytes maximally as indicated by the incorporation of $^3$H-TdR in cultures incubated for 120 hours (Figure 1). Concentrations greater than 5 µg/ml demonstrated relatively decreased incorporation of the label.

Time course of DNA synthesis. Figure 2A and 2B present results of the time course of DNA synthesis in whole blood culture each from two different control and vaccinated ferrets. Cultures lacking PHA did not incorporate $^3$H-TdR, whereas those containing PHA demonstrated maximum DNA synthesis as indicated by the incorporation of the label between 72 and 120 hours.
Time of exposure of cells for maximum $^3$H-TdR incorporation. Incorporation of $^3$H-TdR at variable but high levels was demonstrated in cultures incubated for 120 hours with PHA, when the label was added between four and 24 hours prior to harvest (Figure 3). Maximum incorporation, however, was observed 12 hours prior to harvest.

Leukocyte viability in PHA supplemented whole blood cultures. Figure 4 represents the composite results from five control and four experimental ferrets. Cultures containing PHA showed a decrease in cell viability up to 48 hours, followed by an increase (approximately seven fold) at 144 and 168 hours in the control and experimental ferrets, respectively. A substantial decline in viability was subsequently observed until 240 hours of incubation. The viability of leukocytes from control ferrets containing PHA was relatively greater as compared to leukocytes from vaccinated ferrets.

Viability of leukocytes in the presence of PHA and UV-IBRV. Leukocytes from both the control and experimental animal lacking PHA showed a decline in the percent viability over time (Figure 5A, 5B). Cultures containing PHA demonstrated a decrease in percent viability up to 48 hours, followed by a five to eight fold increase in viable cells at 96 and 144 hours in control (Figure 5A) and vaccinated (Figure 5B) animals, respectively. The number of viable cells from control and vaccinated ferrets was greater in cultures containing either PHA or PHA and UV-IBRV as compared to cultures
lacking them. The cells from the vaccinated ferret, containing both PHA and UV-IBRV demonstrated greater viability at the peak hour (Figure 5B) as compared to control (Figure 5A).

The effect of cortisol on $^3$H-TdR incorporation and viability of PHA-stimulated lymphocytes. Figure 6A illustrates the composite results of DNA synthesis in PHA stimulated cultures from three control and three experimental ferrets. A decrease in the $^3$H-TdR incorporation was evident with increasing concentrations of cortisol. The degree of inhibition of DNA synthesis was greater in the cultures from controls as compared to the vaccinated animals and was minimum at a cortisol concentration of 5,000 ng/ml.

Figure 6B represents the percent viability of lymphocytes from control and experimental ferrets. Viability was greatly reduced as the concentration of cortisol was increased. The highest concentrations (5,000 ng/ml) of cortisol demonstrated only three to six percent viability of the cells.

The effect of cortisol on cell-mediated immunity. Lymphocytes from vaccinated animals were specifically stimulated with UV-IBRV in cultures without cortisol as demonstrated by $^3$H-TdR incorporation (Figure 7). Cortisol (5 ng/ml) inhibited label incorporation 50-75% in cultures from the vaccinated animals. Cells from the control animal did not incorporate label in the presence of UV-IBRV suggesting the
lack of cell mediated immunity to IBRV in the control.

Effect of cortisol on $^3$H-IBRV adsorption. Table 1 demonstrates that the presence of different concentrations of cortisol (0-5,000 ng/ml) did not affect the adsorption of a constant amount of labelled IBRV onto ferret leukocytes. This was indicated by relatively similar radioactive counts and PFU's recovered from the leukocyte suspensions. It appears that the amount of labelled virus adsorbed to experimental ferret leukocytes was somewhat higher as compared to the adsorption to control animals.

Discussion. Studies on the maximum incorporation of $^3$H-TdR in ferret whole blood cultures, supplemented with PHA, was regarded as an important part of these studies, since PHA induced stimulation of lymphocytes provided an index of functional competence of the CMI response (Good, 1972). Based on these results, the effect of cortisol on CMI responses to either specific or non-specific stimulation could be determined adequately. Phytohemagglutinin, in ferret whole blood cultures (incubated 120 hours) at a concentration of 5 ug/ml demonstrated maximum DNA synthesis as measured by the incorporation of $^3$H-TdR (Figure 1). Higher levels of PHA appeared to depress $^3$H-TdR incorporation, while concentrations of three and four ug/ml produced intermediate degrees of stimulation. Concentrations of PHA ranging from 2.5-50 ug/ml have been used in studies with bovine and human lymphocytes (Unsgaard and Lamvik, 1977;
The differences in concentrations reported are probably due to the species differences and the various type of PHA used. In the present study, it is not known that the low $^3$H-TdR incorporation in cultures with concentration of PHA higher than 5 ug/ml was due to the decrease in viability or metabolic activity of the cells.

Maximum DNA synthesis, in PHA stimulated ferret cultures, occurred between 72 and 120 hours of incubation (Figure 2A, 2B). It was reported by Fernald and Metzgar, (1971), that in a given experiment, the cell concentration influenced the timing of optimal DNA synthesis induced by PHA. Furthermore, increased cell concentrations tended to shorten the time for the maximal response to occur. Despite the differences observed for individual animals in these studies, whole blood cultures for subsequent studies were incubated for 120 hours. Ferret cultures containing PHA and exceeding 120 hours of incubation became clumped and the lysis of erythrocytes by the distilled water treatment, to obtain lymphocytes was increasingly difficult over time. Some investigators (Muscoplat, et al., 1974a, 1974b; Pospisil, et al., 1973; Davies and Duncan, 1973; Phillips and Zweiman, 1970) using bovine and guinea pig cultures containing PHA observed optimum incorporation of the label between 48-72 hours of incubation. In other studies, bovine and canine whole blood cultures were similarly incubated for 120 hours (Gerber and Brown, 1974; Rouse and Babiuk, 1974). Incorporation of
$^3$H-TdR decreased rapidly in all cases following the peak of DNA synthesis. This was probably due to the depletion of nutrients in the media and decreased cell viability. Whole blood cultures lacking PHA essentially did not incorporate $^3$H-TdR and the viability counts of these cultures revealed few viable cells (Figure 4). It is known that PHA is a T cell mitogen (Janossy and Greaves, 1971; Sorensen, Anderson and Giese, 1969) suggesting that in these studies, PHA maintained the lymphocytes in whole blood cultures by stimulating mitosis and increasing lymphocyte numbers. Electron microscopic examination of PHA stimulated mink cell cultures demonstrated the blast transformation of lymphocytes at 45 hours of incubation (Marsh, Miller and Hanson, 1973).

Uptake of $^3$H-TdR was optimum in 120 hour PHA stimulated cultures, labelled 12 hours prior to harvest (Figure 3). This indicated that the maximum uptake of the label, a DNA precursor, occurred between 108-120 hours of PHA exposure. Rouse and Babiuk (1974) and Muscoplat, et al. (1974a, 1974b) reported the addition of $^3$H-TdR into bovine lymphocyte cultures 24 hours prior to harvest. In our studies, high levels of $^3$H-TdR incorporation was also observed when label was added four to 24 hours prior to harvest, suggesting relatively active DNA synthesis between these time intervals.

Whole blood cultures from control and vaccinated ferrets (lacking PHA) had low cell viabilities as measured by the trypan blue stain exclusion method (Figure 5A, 5B). It is apparent that the mitotic effects of PHA are not manifested
until 48 hours after supplementation, as illustrated by the decrease in viability (Figure 5A and 5B). In the control animal cultures, the viability of PHA stimulated cultures exceeded the viability of cells containing both PHA and UV-IBRV, indicating that the full mitotic activity of PHA was not realized in the presence of UV-IBRV (Figure 5A). The viability of vaccinated ferret lymphocytes containing both UV-IBRV and PHA or PHA alone, demonstrated similar peaks. This may be due to the added effect by specific antigen (UV-IBRV) on the stimulation of vaccinated ferret lymphocytes (Figure 5B) as compared to the control (Figure 5A). Cell mediated immunity was also demonstrated with specific antigen in cultures of vaccinated animals as indicated by $^3$H-TdR incorporation.

Decreased incorporation of $^3$H-TdR (Figure 6A) and low viability (Figure 6B) in PHA stimulated lymphocytes were demonstrated with increasing concentrations of cortisol. It is probable that cortisol in these studies may have resulted in a cytotoxic effect (Claman, 1972) or the inhibition of PHA stimulation (Zeman, et al., 1973; Roath and Cuppari, 1965). The mechanism of suppression may involve the presence of steroid-receptor complexes in the cell nucleus which are capable of combining with the chromatin. The nuclear complexes then influence DNA transcription, leading to specific changes in cellular function (Feldman, Funder and Edelman, 1972). Lymphocytes from IBRV vaccinated ferrets stimulated with specific antigen (UV-IBRV) required lower concentrations
of cortisol to depress $^{3}\text{H}}$-TdR incorporation (Figure 7) than PHA stimulated lymphocytes (Figure 6A). This effect may have been due to the presence of relatively fewer cells in cultures containing UV-IBRV as illustrated in Figures 5A and 5B. Control ferret lymphocytes did not demonstrate specific stimulation with the IBRV antigen; therefore the effect of cortisol on cell mediated immunity was not demonstrated (Figure 7).

It is apparent from the studies in Table 1 that increasing concentrations of cortisol did not affect the adsorption of $^{3}\text{H}}$-IBRV as indicated by recovery of DPM's and PFU's adsorbed to cells. It is also illustrated that a relatively greater number of virus particles (DPM and PFU) were adsorbed to leukocytes from vaccinated ferrets as compared to controls. This may indicate higher affinity of primed leukocytes for the adsorption of specific antigen. These studies suggest that IBRV and the hormone do not compete for the same receptor sites on ferret leukocytes. Furthermore, it indicates that the mechanism of inhibition of the blastogenic response ($^{3}\text{H}}$-TdR incorporation by UV-IBRV stimulation) by cortisol is possibly independent of the sites of adsorption of IBRV on ferret leukocytes.

The use of both whole blood and short-term ferret lymphocyte cultures in stimulation and immune suppression studies has immense potential. The high levels of response by ferret lymphocytes to PHA has been demonstrated and would be invaluable in studying the mechanisms of immune suppression.
Figure 1. **Effect of PHA concentrations on DNA synthesis:** Whole blood cultures supplemented with varying concentrations of PHA were labelled with $^3$H-TdR 24 hours prior to harvest. At 120 hours the samples were harvested, processed and DPM's recorded.
Figure 2A and 2B. **Time course of DNA synthesis:** Whole blood cultures from control and vaccinated ferrets lacking or containing 5 μg/ml of PHA were incubated for varying time intervals up to 168 hours. Samples were labelled with ³H-TdR 12 hours prior to harvest, processed and counted: ΔΔ Control With PHA; O--O Control Without PHA; ■■ Experimental With PHA; □□□ Experimental Without PHA.
Figure 3. Time of exposure of cells to $^3$H-TdR for maximum label incorporation: Whole blood cultures containing 5 ug/ml of PHA were incubated. Beginning at 96 hours of incubation, $^3$H-TdR was added to sets of cultures at four hour intervals. All cultures were harvested at 120 hours, processed and DPM's determined.
EXPOSURE TO $^3$H-TdR (HOURS)

DPM x $10^3$
Figure 4. Leukocyte viability in PHA supplemented whole blood cultures: Whole blood cultures from control and vaccinated ferrets lacking or containing 5 ug/ml of PHA, were incubated for varying time intervals up to 240 hours. The cultures were harvested every 24 hours. The fold increase of viable cells was determined by the trypan blue stain exclusion method: • — Control Without PHA; o—o Control With PHA; △...△ Experimental Without PHA; ▲—▲ Experimental With PHA.
Figure 5A and 5B. **Effect of UV inactivated IBRV and PHA on leukocyte viability:** Whole blood cultures from control and vaccinated ferrets were supplemented with UV-IBRV (2.0 x 10^5, PFU/0.2 ml), PHA (5 ug/ml) or both. Leukocyte viability was determined by the trypan blue stain exclusion method. The percent viability was calculated from the zero hour control: △—△ Media; O—O PHA; △—△ UV-IBRV; ••• PHA and UV-IBRV.
Figure 6A. Effect of cortisol on PHA stimulated lymphocytes: Whole blood cultures from control and vaccinated ferrets were stimulated with 5 ug/ml of PHA in the presence of varying concentrations of cortisol. \(^3\)H-TdR was added 12 hours prior to harvest. The cultures were harvested at 120 hours, processed and counted. The DPM's were calculated as percent of the control with PHA: •-• Control; △-△ Experimental.

Figure 6B. Effect of cortisol on the viability of PHA stimulated lymphocytes: Whole blood cultures from control and vaccinated ferrets were stimulated with 5 ug/ml of PHA in the presence of varying concentrations of cortisol. The cultures were incubated for 120 hours. Viability of cells was calculated as percent of the control with PHA: •-• Control; △-△ Experimental.
Figure 7. Effect of cortisol on cell mediated immunity:
Lymphocytes from vaccinated and control ferrets were inoculated with UV-IBRV in the presence of varying concentrations of cortisol. \(^3\)H-TdR was added 12 hours prior to harvest. The cultures were harvested and processed at 120 hours and DPM's determined:

- Experimental:
- Control.
CONCENTRATION OF CORTISOL (ng/ml)

DPM x 10^2

0 25 50 75 100 5000
TABLE 1
EFFECT OF CORTISOL ON $^{3}$H-IBRV ADSORPTION TO FERRET LEUKOCYTES

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<thead>
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<th>Concentration of Cortisol</th>
<th>Control DPM</th>
<th>Control Total PFU</th>
<th>Experimental DPM</th>
<th>Experimental Total PFU</th>
<th>Experimental DPM</th>
<th>Experimental Total PFU</th>
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<tr>
<td>0 ng</td>
<td>3028.5</td>
<td>8.0x10$^{3}$</td>
<td>2309.5</td>
<td>8.0x10$^{3}$</td>
<td>3259.1</td>
<td>-</td>
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<tr>
<td>5 ng</td>
<td>2731.3</td>
<td>7.9x10$^{3}$</td>
<td>2776.5</td>
<td>8.4x10$^{3}$</td>
<td>3149.0</td>
<td>1.42x10$^{4}$</td>
</tr>
<tr>
<td>50</td>
<td>2831.3</td>
<td>7.8x10$^{3}$</td>
<td>2144.3</td>
<td>8.3x10$^{3}$</td>
<td>3248.5</td>
<td>1.7x10$^{4}$</td>
</tr>
<tr>
<td>500 ng</td>
<td>3061.1</td>
<td>-</td>
<td>2525.6</td>
<td>1.18x10$^{4}$</td>
<td>3381.4</td>
<td>1.64x10$^{4}$</td>
</tr>
<tr>
<td>5000 ng</td>
<td>2853.9</td>
<td>1.1x10$^{4}$</td>
<td>2835.0</td>
<td>7.6x10$^{3}$</td>
<td>3240.5</td>
<td>-</td>
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</table>
SECTION IV
APPENDIX
Materials and Methods.

UV inactivation of viruses. One ml of BVDV or IBRV (1 x 10^7 PFU/ml) was UV irradiated at one minute intervals for 0-6 minutes, in a petri dish (60 x 15 mm) at a distance of 10 cm from a UV germicidal sterilamp (G30T8, 30 watt, Westinghouse Electric Co., Bloomfield, NJ) (Figure 1). The virus was collected and plaque assayed to determine the percent inactivation for each interval of exposure.

Formalin inactivation of IBRV. For the preparation of inactivated vaccine, 5 x 10^8 PFU/ml of IBRV was treated with formalin at final concentrations of 1:1000, 1:2000, 1:4000, 1:8000, and 1:16000 (40% formaldehyde solution, Fisher Scientific Co., Fairlawn, NJ) for six hours at 25°C, then overnight at 4°C (Figure 2). The virus was dialyzed in 1/4" dialyzing tubing against three changes of cold PBS at 4°C over a 24 hour period. The virus was collected and plaque assayed to determine the percent inactivation for each concentration.

Butylated hydroxytoluene inactivation of IBRV. A stock solution of butylated hydroxytoluene (BHT) (Sigma Chemical Co., St. Louis, MO) 0.5 ug/10 ml ethanol was prepared. One ml of IBRV, 1 x 10^6 PFU/ml was treated with 0, 5, 50, 100, 250 and 500 ug BHT, agitated every five minutes, for 30 minutes at 37°C (Figure 3). The preparation was stored at -80°C then plaque assayed to determine the percent
inactivation of virus.

**Determining the time course of DNA synthesis.** The method of whole cultures used by Gerber and Brown (1974) was adopted with modification as previously described. Whole blood cultures containing 5 µg/ml of PHA were incubated for varying time periods up to 240 hours. The cultures were exposed to 0.5 uCi/ml of tritiated thymidine (\(^{3}\)H-TdR) in medium 24 hours prior to harvest each time. The samples were harvested at various intervals. The cultures were treated with equal volumes of distilled water and 1.7% NaCl solution, processed and counted on glass filters as previously described (Figure 4).
Figure 1. Inactivation of IBRV and BVDV by UV irradiation. One ml of virus ($1 \times 10^7$ PFU) was UV irradiated for 0-6 minutes. The virus was collected and plaque assayed to determine the percent inactivation for each interval of exposure; •—•IBRV; ○—○BVDV.
Figure 2. **Formalin inactivation of IBRV.** One ml of IBRV \(5 \times 10^8\) PFU) was treated with formalin at final concentrations of 1:1000, 1:2000, 1:4000, 1:8000, 1:16,000 and 1:32,000. The virus was plaque assayed to determine the percent inactivation for each concentration.
PERCENT INACTIVATION vs. FORMALIN CONCENTRATION ($\times 10^{-4}$)
Figure 3. **Inactivation of IBV by butylated hydroxytoluene.** One ml of IBV (1 x 10^6 PFU) was treated with 0-500 ug BHT. The virus preparation was plaque assayed to determine the percent inactivation for each concentration.
Figure 4. Time course of DNA synthesis: Whole blood cultures lacking or containing 5 ug/ml of PHA were incubated for varying time intervals up to 240 hours. Samples were labelled with $^3$H-TdR 24 hours prior to harvest, processed and counted: •• With PHA; △△ Without PHA.
HOURS OF INCUBATION

DPM x 10^3

0  48  96  144  192  240
Mean Rectal Temperatures\textsuperscript{A} from Ferrets Inoculated with IBRV

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Experimental</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Sample Size</td>
<td>Mean (+S.D.)</td>
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<tr>
<td>Inactivated Vaccine</td>
<td>Pre: 10</td>
<td>102.1 (+0.2)</td>
</tr>
<tr>
<td></td>
<td>Post: 126</td>
<td>102.1 (+0.5)</td>
</tr>
<tr>
<td>Day 1-65 Post-</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vaccination</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MLV Vaccine (IN)</td>
<td>Pre: 1</td>
<td>102.0</td>
</tr>
<tr>
<td></td>
<td>Post: 7</td>
<td>101.4 (+0.5)</td>
</tr>
<tr>
<td>Day 1-49 Post-</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vaccination</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MLV Vaccine (IM)</td>
<td>Pre: 1</td>
<td>102.4</td>
</tr>
<tr>
<td></td>
<td>Post: 8</td>
<td>101.3 (+0.5)</td>
</tr>
<tr>
<td>Day 1-49 Post-</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vaccination</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\textsuperscript{A}Normal Ferret Temperatures: 101-104\textdegree F (Pyle, 1940).

\textsuperscript{B}Pre-Inoculation
Post-Inoculation
WEIGHTS$^A$ OF CONTROL FERRETS$^B$
INOCULATED WITH FREUNDS INCOMPLETE ADJUVANT

<table>
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<tr>
<th>Age (Weeks)</th>
<th>Weight Mean ± (S.D.)</th>
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</thead>
<tbody>
<tr>
<td>13</td>
<td>0.85 (±0.03)</td>
</tr>
<tr>
<td>14</td>
<td>0.92 (±0.07)</td>
</tr>
<tr>
<td>15</td>
<td>1.00 (±0.15)</td>
</tr>
<tr>
<td>16</td>
<td>1.04 (±0.13)</td>
</tr>
<tr>
<td>17</td>
<td>0.99 (±0.14)</td>
</tr>
<tr>
<td>18</td>
<td>1.07 (±0.13)</td>
</tr>
<tr>
<td>21</td>
<td>1.04 (±0.18)</td>
</tr>
<tr>
<td>22</td>
<td>1.21 (±0.16)</td>
</tr>
<tr>
<td>23</td>
<td>1.15 (±0.13)</td>
</tr>
<tr>
<td>24</td>
<td>1.24 (±0.10)</td>
</tr>
<tr>
<td>28</td>
<td>1.42 (±0.11)</td>
</tr>
<tr>
<td>30</td>
<td>1.45 (±0.09)</td>
</tr>
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</table>

A. Weights (kg)

B. Two castrated male ferrets also served as controls for the inactivated TBRV inoculated ferrets, but were of a different age.
## WEIGHTS\(^A\) OF FERRETS INOCULATED WITH MLV

<table>
<thead>
<tr>
<th>Age (Weeks)</th>
<th>Nasalgen (IN)</th>
<th>Delvine(^R) (IM)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
<td>Experimental Mean (±S.D.)</td>
</tr>
<tr>
<td>16</td>
<td>0.91</td>
<td>0.94 (±0.08)</td>
</tr>
<tr>
<td>17</td>
<td>0.99</td>
<td>0.92 (±0.05)</td>
</tr>
<tr>
<td>18(^B)</td>
<td>0.94</td>
<td>0.91 (±0.0)</td>
</tr>
<tr>
<td>21</td>
<td>0.97</td>
<td>1.17 (±0.09)</td>
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<tr>
<td>22</td>
<td>1.02</td>
<td>1.16 (±0.06)</td>
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<tr>
<td>23</td>
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<td>1.33 (±0.05)</td>
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<tr>
<td>24</td>
<td>1.05</td>
<td>1.36 (±0.05)</td>
</tr>
<tr>
<td>26</td>
<td>1.08</td>
<td>1.49 (±0.08)</td>
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<tr>
<td>28</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>30</td>
<td>1.42</td>
<td>1.57 (±0.13)</td>
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</table>

A. Weight (kg)

B. Ferrets were vaccinated with MLV at 20 weeks of age.
SECTION VII

DETAILED METHODOLOGY AND REAGENTS
Reagents

Agar: 2% purified agar dissolved in deionized water and autoclaved for use. A mixture of 50% agar - 50% (2X) MEM is used for plaquing.

Antibiotics:
Penicillin (1,000,000 units)
Streptomycin (1 g)
Kanamycin (50 mg) is dissolved in 100 ml of sterile deionized water. One ml of PSK stock solution is added to give a final concentration of: 100 units penicillin, 1 mg streptomycin and 5 ng kanamycin/ml media.

Dimethyl Sulfoxide (CH₃)₂SO₄:
For freezing cells, this is made to an appropriate concentration (10%) and autoclaved 10 minutes at 15 minutes.

DNase: 2 mg/ml, stock. A final concentration of 10 μg/ml is used.

DNase: 2 mg/ml, stock. A final concentration of 10 μg/ml is used.

Fungizone: 50 mg (one vial) plus 20 ml sterile deionized water. Add 0.2 ml/l. media.

Neutral Red: A solution (0.015%) for plaquing is prepared from powder with deionized water and dissolved in a 37° C water bath. Whatman #2 filter paper is used for filtration. Sterilize by autoclaving and store at 4° C, protected from light. 1.5 ml stock/100 ml agar and media.
Phytohemagglutinin: 50 ug/ml in RPMI 1640 stock. A final concentration of 5 ug/ml is used.

Phosphate-Buffered Saline (PBS):

Solution 1

\[ \text{Na}_2\text{HPO}_4 \cdot 7\text{H}_2\text{O} \quad 134 \text{ g/l deionized H}_2\text{O} \]

Solution 2

\[ \text{NaH}_2\text{PO}_4 \cdot \text{H}_2\text{O} \quad 69.1 \text{ g/l deionized H}_2\text{O} \]

Buffer Stock

\[ \text{Na}_2\text{HPO}_4 \cdot 7\text{H}_2\text{O} \quad 386 \text{ ml Solution 1} \]
\[ \text{NaH}_2\text{PO}_4 \cdot \text{H}_2\text{O} \quad 114 \text{ ml Solution 2} \]
\[ \text{Deionized Water} \quad 500 \text{ ml} \]
\[ \text{pH 7.15} \]

PBS:

80 ml buffer stock

17 g NaCl

1920 ml deionized water

pH 7.2

Autoclave at 15 lbs. for 15 minutes

Scintillation Fluid:

360 ml Toluene

360 ml Dioxane

80 g Napthalene

216 ml Absolute alcohol

4 g Ominifluor
### Reagents and Addresses

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<tr>
<td>Agar</td>
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<td>Cortisol</td>
<td>Calbiochem, Los Angeles, CA</td>
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<td>Dimethyl Sulfoxide</td>
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<td>Dioxane</td>
<td>Scientific Products, IL</td>
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<td>DNase (pancreatic)</td>
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<tr>
<td>Trypsin</td>
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SECTION VIII

REFERENCES


IMMUNOLOGICAL RESPONSES IN FERRETS INOCULATED WITH INFECTIOUS BOVINE RHINOTRACHEITIS VIRUS VACCINES

by

ELSBETH JANE LEE
B.S., University of Vermont, 1975

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ABSTRACT

The hematological and serum chemistry profiles and the immunological responses of ferrets (*Mustela putorius furo*) to Infectious bovine rhinotracheitis virus (IBRV) vaccines were studied.

Statistical analyses were performed on the hematological and serum profiles of normal female, male and castrated male ferrets. No appreciable differences in the mean values were observed among the groups. Simple linear correlation coefficients were calculated for these parameters with respect to age or body weight. Significant age related changes were found in the neutrophils, lymphocytes, sorbital dehydrogenase, blood urea nitrogen, alkaline phosphatase, serum glutamate pyruvate transaminase, albumin and glucose. Significant weight related changes were associated with sodium, carbon dioxide, calcium, phosphorus and glucose.

Ferrets inoculated with different IBRV vaccines demonstrated significant differences in treatment means for sorbital dehydrogenase, blood urea nitrogen, albumin, creatinine, packed cell volume and bands. Statistical significance was not found among the other hematological and serum chemistry profiles.

Although all ferrets did not manifest clinical signs of infection, IBRV was isolated three days post vaccination in the nasal washings from one of the three ferrets inoculated with modified live virus (MLV); however, virus was not
detected in subsequent samples. Low degrees of specific and non-specific virus neutralizing factors were detected by the plaque reduction method in the nasal washings only at the fourth week post IN inoculation with inactivated IBRV.

Serum neutralizing antibodies were demonstrated in ferrets three weeks following an intramuscular (IM) injection of inactivated IBRV with Freund's Incomplete Adjuvant and increased after a second IM inoculation. Complete virus neutralization was observed between the 12th and 14th week. The MLV vaccines (given IM or IN) elicited low levels of serum antibodies that persisted from the third to the seventh week post inoculation. The degree of cell mediated immune response was variable (stimulation index: 1.1-27.7) as observed by specific (UV-IBRV) lymphocyte stimulation and the incorporation of tritiated thymidine in whole blood cultures from ferrets inoculated with inactivated IBRV. Controls and MLV vaccinated ferrets did not demonstrate cell mediated immune responses to the IBRV antigen.

Phytohemagglutinin (PHA) at a concentration of 5 ug/ml demonstrated maximum stimulation of lymphocytes in ferret whole blood cultures. The optimal effect of PHA on DNA synthesis was observed between 72 and 168 hours after incubation and when the cultures were exposed to the label 12 hours prior to harvest.
Following an initial decline in viability at 48 hours, the number of viable lymphocytes in PHA supplemented cultures increased approximately 10 fold between 96 and 144 hours of incubation. Cultures lacking PHA or containing UV-IBRV demonstrated a time related decline in the viability of lymphocytes.

Cortisol at physiological concentrations inhibited the viability of PHA stimulated lymphocytes as indicated by an appreciable decrease in cell count and the incorporation of $^3$H-TdR. Cortisol also depressed the cell mediated immune response in UV-IBRV stimulated cultures. The presence of increasing concentrations of cortisol (0-5,000 ng/ml) did not affect the adsorption of $^3$H-IBRV to ferret leukocytes. This suggests that cortisol and IBRV did not compete for the same receptor sites.