The characterization of two bovine adenoviruses isolated from calves exhibiting clinical and pathological signs of weak calf syndrome
by Patricia Knox Shadoan

A thesis submitted in partial fulfillment of the requirements for the degree of MASTER OF SCIENCE in Veterinary Science
Montana State University
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Abstract:
Of several viruses recovered from calves exhibiting clinical signs of weak calf syndrome, two were selected for characterization studies.

Characterization of isolates 18115 and 1-1504 involved the determination of their nucleic acid type, determination of size and shape, presence of essential lipid, pH and heat sensitivity, growth properties, and serologic analysis.

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By the process of ultrafiltration and transmission electron microscopic examination, the viruses were shown to be spherical particles approximately. 70 nm in diameter.

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Isolates 18115 and 1-1504 were neutralized by antiserum produced against serotype 7 bovine adenovirus and positive fluorescence was observed by the indirect fluorescent antibody test in virus-infected cell monolayers treated with antiserum to bovine adenovirus serotype 7.

Bovine adenoviruses are known to be important in the etiology of bovine respiratory and enteric disease; the isolation of these agents from the tissues of weak calves indicates that additionally they may be of etiologic importance in the pathogenesis of weak calf syndrome.
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THE CHARACTERIZATION OF TWO BOVINE ADENOVIRUSES ISOLATED FROM CALVES EXHIBITING CLINICAL AND PATHOLOGICAL SIGNS OF WEAK CALF SYNDROME

by

PATRICIA KNOX SHADOAN

A thesis submitted in partial fulfillment of the requirements for the degree of MASTER OF SCIENCE in Veterinary Science

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March, 1979
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Bovine adenoviruses are known to be important in the etiology of bovine respiratory and enteric disease; the isolation of these agents from the tissues of weak calves indicates that additionally they may be of etiologic importance in the pathogenesis of weak calf syndrome.
Weak Calf Syndrome

Weak calf syndrome (WCS) is a disease entity of newborn calves that was first recognized in the Bitterroot Valley of western Montana in 1964. Four years later it was seen in Idaho, where veterinary practitioners diagnosed 400 cases around the Salmon-Challis area. Since that time, the condition has been frequently observed in Beaverhead County, Montana and Custer County, Idaho, and has also been seen sporadically in other areas of these two states. Additionally, it has been reported in several other states including California, Colorado, Nevada, Oregon and Nebraska. Calf losses during spring calving of between 6 to 15 percent and up to as many as 50 percent in individual herds implicate it as the cause of great economic loss. This disease entity is known by a variety of terms including neonatal idiopathic polyarthritis, Ward's syndrome and Bitterroot crud, but is most frequently referred to as WCS (12, 25, 31).

Clinical signs attributed to this disease develop within the first ten days of the calf's life, the majority of the afflicted calves showing signs of illness between three and seven days of age. Among the usual presentation of clinical signs accorded this syndrome are severe debilitation and depression; the calf is weak and unable to stand and nurse. Often the animal's muzzle is encrusted and bright
red in color. Many calves are lame and show an inability or unwillingness to stand, or may stand with an arched back and be reluctant to move. Front and rear legs may exhibit edema and swelling of the periarticular tissues of the carpel and tarsal joint sacs. The conjunctiva and third eyelid may show petechial and diffuse hemorrhage. The body temperature is usually normal, dropping below normal as death approaches. Diarrhea and dehydration are not initially observed but are usual sequelae if the calf lives for several days. Mortality rates vary from 60 to 80 percent in untreated cases (12, 25, 31).

Of the lesions noted during necropsy, the most outstanding are subcutaneous hemorrhages and edema in the tissues of the extremities. These are noticeable primarily around the hock and knee joint, and extend distally over the lateral aspects of the respective limbs. The edema is most pronounced around the hock joint and tissues supporting the Achilles tendon. These particular lesions are the ones evident most commonly in calves with WCS and are seen in 95 to 100 percent of the cases.

The second most striking gross lesion is a polyarthritis with hemorrhagic synovial fluid containing varying amounts of fibrin. Of 25 instances of WCS from the Salmon-Challis area of Idaho, this was present in 65 percent of the cases.
In approximately 30 percent of the cases, lesions of the forestomach and abomasum are seen involving diffuse congestion and hemorrhage, with occasional vesiculation, erosion, and ulceration of the epithelial lining. In severe cases, hemorrhage and edema can be observed on the ventral aspect of the thorax and neck (12, 25, 31).

Factors thought to contribute to the severity of clinical illness in WCS outbreaks include the stress of inclement weather and the quality of herd management (12, 25). Studies conducted by Bull and co-workers indicate nutrition levels may also be an important consideration (10).

Attempts to recover infectious agents from weak calves have met with varied results. In 1969 Page and associates isolated a previously undescribed mycoplasmal agent from the placenta of a cow that gave birth to a weak calf (58) and a bacterial agent, *Hemophilus somnus*, was recovered by Waldham from the vaginal mucous of a dam that produced a weak calf (76). Neither of these organisms have been shown to be the causative agent of WCS, although trials with *Hemophilus somnus* produced a polyarthritis similar to that found in field cases (76).

Several viral agents have been isolated from tissues of weak calves over the last several years by investigators in Montana, Idaho, and Iowa. Coria from the National Animal Disease Center (NADC) in Iowa, and Stauber at Idaho State University both isolated bovine viral
diarrhea (BVD) virus and an adenovirus (AV) from weak calf tissues. Characterization studies show these AV isolates to be serotype 7 bovine AV (BAV) (15, 73).

Cutlip and McClurkin from NADC have reported results of their experimental infections of young calves with BAV-7. They were able to reproduce hemorrhagic and necrotic lesions of the subcutaneous tissues of the legs and joints that are commonly associated with WCS, but they also observed hemorrhagic and necrotic lesions of internal organs (kidneys, adrenal gland, and liver) that are not generally noted (17, 45).

Experimental intraamnionic exposure of bovine fetuses to BAV-7 produced calves that were born showing clinical signs resembling those described for WCS. However, the gross and microscopic lesions generated in this experiment were distinctly different from those described for the syndrome and it was concluded that the disease produced was not WCS (72).

**Statement of Purpose**

During the 1975 calving season, Dierks, Smith, and Gollehon of Montana State University, isolated 22 viral agents from the tissues of 10 suspected weak calves, fetuses, and one lamb (26). The purpose of the work reported here was to characterize, by studies determining the
physicochemical and biological properties, two of the virus isolates recovered from the tissues of these weak calves.
Adenoviruses

In 1953, Rowe and associates isolated a cytopathic agent from human adenoid tissue that they had been culturing as a potentially favorable host for an elusive respiratory virus. After a prolonged incubation period, pathologic changes were noted in uninoculated as well as inoculated cell cultures. This cytopathic response was shown to be due to the emergence of a previously unidentified virus from latent infections of the adenoid tissue (69). The designation "adenovirus" was officially given to this group of viruses to indicate their original isolation site (28).

AV are widespread in nature and now comprise a well-defined group of viruses. Members of this family are simple viruses composed of a core of double-stranded DNA and a surrounding protein coat arranged in the form of an icosahedron. Two hundred fifty-two capsomeres comprise this outer protein shell of which 240 are hexon capsomeres and 12 are penton capsomeres. The pentons are apical subunits upon which are found fibers responsible for attachment to the host cell (24).

Mature AV particles range between 65-90 nm in diameter, averaging about 70 nm in diameter. These viruses are non-enveloped and subsequently exhibit resistance to lipid solvents.
(i.e., chloroform or ether) (59). They are acid stable, but are subject to inactivation at 56°C for 10 minutes (63).

AV multiplication and maturation takes place in the nucleus of susceptible cells, producing characteristic cytopathic effect (CPE) and type B intranuclear inclusions. The "cell detaching factor" or "toxin" that is a property of the intact penton is responsible for the typical rounding, clumping and detachment from surfaces that is seen in infected cells (24).

The antigenic character of the AV depends largely on its morphological subunits. The hexon capsomere contains the group-specific or complement-fixing (CF) antigen as well as a type-specific reactive site. The major type-specific antigen as well as minor subgroup antigens are found in the purified fiber, the length of which varies according to serotype. The penton provides some minor antigens of the virion and is also found as a family-reactive soluble antigen (24, 59, 83).

This family of viruses is highly species specific and is arbitrarily divided, according to the natural hosts, into six subgroups represented by human, simian, bovine, canine, murine and avian AV. All except the avian types share a common CF antigen (59).
Bovine Adenoviruses

The first isolations of AV from the bovine species were made in the United States by Klein and co-workers (39). As the result of their search for viruses responsible for the production of poliovirus antibodies in cattle, two AV were isolated in bovine kidney cultures from the feces of clinically normal cattle. These two virus strains, designated bovine no. 10 and bovine no. 19, are now considered the prototypes representing BAV type 1 and type 2, respectively. Neutralization of these bovine viruses by human gammaglobulin shows the possibility of a relationship to human AV (39, 40). Subsequently, BAV-3 was isolated in bovine kidney cultures from the conjunctiva of an apparently healthy cow in Britain (20).

Pathogenesis studies of experimental infection of calves with these BAV have produced varied results. Klein recorded that the intranasal inoculation of two calves with BAV-1 (bovine no. 10) effected a neutralizing antibody response but no concomitant symptoms (40). Using strain 10088 of BAV-1, Mohanty and Lillie demonstrated the production of marked clinical signs of respiratory distress and diarrhea when this virus was inoculated intranasally and intratracheally into 6 to 12-week-old calves. Calves infected via the ocular route displayed no noticeable response (56).

Strain WBR 50 of BAV-1, recovered by Darbyshire from the nasal swabs of a calf suffering from pneumonia, induced type-specific
hemagglutination-inhibiting antibodies and mild clinical signs in experimentally infected calves (18).

In other pathogenesis studies conducted by Darbyshire and associates, BAV-3 (strain WBRI) was shown to invoke a mild clinical response characterized by pyrexia, respiratory distress, nasal and conjunctival discharges. Gross pathologic changes were primarily confined to the lungs which showed areas of consolidation, collapse, and emphysema. The histopathologic changes in the lungs consisted of proliferative bronchitis and necrosis with bronchiolar occlusion and associated alveolar collapse. Diarrhea was evident, but not a prominent feature of the disease (20, 21). Pulmonary reactions to BAV-1 (bovine no. 10) and BAV-2 (bovine no. 19) involved gross and microscopic tissue changes similar but less severe than those observed with type 3. These data indicated that type 3 is the most virulent of the three types of BAV (22). Conversely, Ide and co-workers failed to produce signs of respiratory illness in calves they had inoculated with either BAV-3 (WBRI) or a strain of BAV-1 which was isolated in Canada from a non-fatal case of pneumonia (35).

Additional recorded isolations of these viruses have been reported and include that of BAV-1 from a calf with pneumoenteritis (70), from bovine cell cultures (27), and from normal calves. BAV-2 has been recovered as an adventitious contaminant in primary bovine embryonic kidney cell cultures (57) and Mattson describes the
isolation of BAV-3 (strain 5C) from a naturally occurring pneumo-enteritis of newborn calves in Oregon beef herds (50).

All of these BAV types 1, 2, and 3, have been shown to be associated with field outbreaks of respiratory disease, and neutralizing antibodies to these agents are widely distributed in world cattle populations (9, 17, 19, 21, 30, 43, 50, 55, 68, 84).

In Hungary, three strains of BAV were isolated from calves suffering from diarrhea (4) and 78 strains were recovered from afflicted calves during a severe outbreak of pneumonia associated with enteritis. During recent years this particular condition has been the most widespread viral infectious disease among calves in Hungary, resulting in great economic loss (2). These BAV strains differed serologically from the previously established BAV prototype strains and represented two further types designated serotype 4 (prototype strain THT/62) and serotype 5 (prototype strain B4/65). These viruses were isolated in primary bovine testicle (BT) cell cultures, and failed to grow in any other cell culture (5).

Aldasy and Bartha were able to show experimentally that inoculation with types 4 and 5 produced pneumonia and enteritis in calves (2). In a study conducted by Mohanty, BAV types 4 and 5 caused respiratory illness with pneumonia in infected calves, but did not cause enteritis (55).
Tanaka reported the first recovery of BAV from cattle in Japan. BAV strain Nagano was isolated in primary BT cells from the blood of a 25-month-old bull with pyrexia, soft stools, and reduced appetite. When tested for pathogenicity in calves, this isolate produced generally mild clinical responses including fever, diarrhea, and rhinorrhea. A low-titered viremia was evident in several calves (74).

Studies conducted by Matumoto demonstrated strain Nagano to be serologically related to BAV-4 (52). A conflicting report by Mohanty designates strain Nagano as a new BAV, type 10, unrelated to previously established BAV serotypes (55). More recently BAV-4 isolates have been recovered from Oregon cattle with pyrexia associated with signs of respiratory tract disease (51) and several reports indicate that BAV-4 has been isolated from latent infections of BT cell cultures (6, 61).

In the Netherlands, Rondhuis isolated the prototype strain (671130) of BAV-6 as a latent virus from primary BT cell cultures. As with types 4 and 5, this virus grew only in these cultures. Intratracheal inoculation of type 6 in newborn calves induced mild respiratory illness and catarrhal conjunctivitis. Viremia was associated with the infection, and the virus was found localized in several organs including the CNS (49, 67).
Cole isolated 11 BAV strains in Australia from the lungs of calves with acute pneumonia. On the basis of the two types of CPE produced in monolayers by the isolates, they were divided into two groups designated BIL and RG. The BIL strains were recovered from the lungs of animals with acute exudative pneumonia, and the RG strains from the lungs and/or noses of calves with pneumonia and bronchitis of varying severity. These viruses were isolated only in primary BT cultures and failed to grow in other cell types (13, 14). Strain RG has been serologically identified by Mohanty as being a member of type 6 BAV (55). A mild interstitial pneumonia was produced in calves following intratracheal inoculation with BIL. Here the pathologic reactions were confined to the lung, which evinced areas of hyperemia, consolidation and edema (14).

Wilcox isolated other BAV strains from Australian cattle with conjunctivitis and keratoconjunctivitis. These viruses were isolated in primary BT cultures. Of the two types represented, one strain, KC-6, grew only in primary and secondary BT cultures while the other strain, KC-2, was able to grow in several bovine cell types (81). The KC-2 strain has been identified as a BAV-6 (55).

Another AV closely related to BAV-6 has been isolated from the precapsular lymph node of a cow with leukosis. Experimental inoculation of young calves produced acute rhinitis, undulating fever, swelling of lymph nodes, and in some cases, mild conjunctivitis (54).
A BAV strain was isolated by Inaba and co-workers from cattle in Japan suffering from acute pyrexia accompanied by rhinorrhea and diarrhea (36). This strain, Fukuroi, isolated in primary BT cultures, represents the BAV serotype 7 (36, 52). This BAV serotype has also been reported to have been isolated from newborn calves with weak calf syndrome (15, 26, 73), as well as from latent infections of primary BT cell cultures (61).

BAV type 8, strain Misk/67, was isolated in Hungary from calves with pneumoenteritis. Experimentally, type 8 was non-pathogenic when inoculated into calves (8).

Guenov reported the isolation of strain Sophia-4/67 from cultures of kidney tissue from healthy calves. This isolate showed no serologic relationship in neutralization tests with BAV types 1-8 and Guenov has proposed that this virus represents the new BAV-9 (32). Mohanty has also designated two of the strains isolated in Australia, KC-6 and BIL, as BAV-9 (55). It is not known whether these isolates are of the same serotype as Sophia-4/67, but they cannot be the prototypes for BAV serotype 9, as Guenov's proposal was published prior to that of Mohanty's.

Kretzschmar has reported the isolation of an AV that differs serologically from BAV types 1-9 (42). This may represent either a new BAV serotype 10 or BAV serotype 11, depending upon the classification of isolates BIL and RG.
To date, there are nine recognized BAV serotypes (1). Bartha has proposed the division of these AV into two distinct groups based on differences in their biological and physicochemical properties. He has suggested that strains belonging to serotypes 1, 2, and 3 possess similar properties and should be categorized as subgroup I BAV, and strains belonging to serotypes 4 and 5 as well as the strains designated BIL, Nagano and Fukuroi, should be classified among the subgroup II BAV (3). More recent work includes BAV-9 in the first subgroup (44).

According to Bartha's study, members of subgroup I possess a soluble antigen in common with the human strains of AV, while only traces of the same are detected in BAV of subgroup II. Primary isolations of subgroup I BAV can be made on bovine kidney and testicle cell cultures, generally on the first passage, while subgroup II BAV can only be isolated on bovine testicle and fetal lung cell cultures after several blind passages. Intranuclear inclusions generated by members of the first subgroup are characteristically single and irregular in shape, as opposed to those produced by the members of the second subgroup, which are multiple and regularly shaped. Bartha reports another differentiating criteria to be that of heat sensitivity: Subgroup I BAV are completely inactivated at 56°C after 30 minutes, whereas subgroup II BAV were only reduced in infectivity by such treatment (3). Other studies discounted this as a valid
parameter on the basis that the AV lost their heat sensitivity after several passages in cell culture (7).

All these studies, in addition to serologic surveys of bovine herds which indicate that BAV infection is widespread, have in recent years drawn attention to BAV as important etiologic agents of respiratory and enteric disease.
CHAPTER 3
MATERIALS AND METHODS

Cell Cultures

Several bovine cell types were utilized during this study to ascertain their susceptibility to viral-induced CPE. Primary and low passage kidney, lung, salivary gland, turbinate, testicle, and thyroid cells were obtained from fetuses purchased from the local slaughter house. A bovine kidney (GBK) cell line developed by R. F. Solarzano, Columbia, Missouri, and a BVD-free bovine turbinate (BTU) cell line developed at NADC (46) were also used. The BTU cell line was used almost exclusively for virus propagation and characterization procedures.

The media used for cell cultures was Dulbecco's modified Eagle minimum essential medium (DMEM) supplemented with 10 percent fetal calf serum (FCS) for cell culture growth reduced to two percent FCS for the maintenance of cell culture monolayers. Irradiated bovine serum distributed by Microbiological Associates was used in media for cultures of the BVD-free cell line. All media were buffered with sodium bicarbonate diluted to a final concentration of 0.15 percent. Liquid antibiotics and antimycotics were used for control of contamination and were added to media in the following concentrations: penicillin 100 mcg/ml, streptomycin 100 mcg/ml, and amphotericin B 1-5 mcg/ml. Routinely, cell cultures were grown in 250 ml Costar
plastic tissue culture flasks and incubated at 37°C in a 5 percent CO₂ atmosphere.

The procedures outlined by Younger were used for establishing primary and low passage cell cultures (85). Bovine fetal tissues were taken aseptically and immediately placed in sterile growth medium. After removing superfluous membranes, the tissues were minced with scalpel blades into approximately 1 mm³ pieces. These fragments were washed three times with serum-free DMEM before being transferred to a 250 ml trypsinizing flask containing 100 ml of a .25 percent trypsin solution. The flask was placed on a magnetic stirrer for 15 minutes, after which time the fluid was decanted and replaced with fresh trypsin. This process was repeated twice more prior to straining the suspension through gauze into a conical centrifuge tube for low speed centrifugation (50-90 xg for 10 minutes). The supernatant fluid was then decanted and the packed cells resuspended in nutrient medium containing antibiotics and antifungicals before being added to flasks in concentrations of between 150,000-500,000 cells/ml.

Cells were subcultured regularly to maintain cell viability and as needed for virus work. To subculture cells, the growth medium was first decanted from flasks and the cell monolayers washed with 5 ml of a calcium and magnesium-free saline, Rinaldini enzyme solution (R-saline) (65). The cells were then covered with 2 ml of a solution of .1 percent trypsin and .04 percent EDTA in R-saline and incubated
for about 10 minutes at 37° C until they detached from the surface of the flask. The cells were dispersed by pipetting several times before being passed into a new flask holding 15 ml of growth medium. Antibiotics and antimycotics were omitted from the media used when subculturing cells.

For stock supplies, concentrated cells were suspended in medium containing 20 percent FCS and 10 percent DMSO by volume, and then sealed into freezing vials. After being brought slowly to -70° C, the vials were immersed in liquid nitrogen for storage until needed.

When infecting cells with virus, newly formed cell monolayers were used whenever possible. When the monolayers were to be infected, the growth medium was removed and the cells washed with R-saline before adding the appropriate viral dilutions. A sufficient amount of virus was added to facilitate an even distribution of virus on the cell sheet. The flasks were then incubated for one to two hours to allow virus adsorption to the cells, after which time the inoculum was removed and maintenance medium added. All manipulations involving cell cultures and inoculation of viruses were conducted under a Bioguard laminar-flow hood.
**Virus Titrations**

For titrating virus, cell monolayers were established in wells of Linbro plates of 24. The growth medium was decanted from the wells and the cells washed with R-saline. Two-tenths ml of 10-fold virus dilutions made in DMEM without serum was added to the appropriate wells. All titrations were done in quadruplicate. These plates were then incubated for one to two hours at 37° C in plastic containers designed for retaining a humid environment to avoid drying in the wells. The virus inoculum was then pipetted out and replaced with 1 ml of maintenance medium. The plates were checked daily with an inverted microscope for CPE, and the 50 percent tissue culture infective doses (TCID$_{50}$) were calculated by the method of Reed and Muench (64).

**Source of Virus**

Several viruses were isolated in BTU cell cultures from the tissues of calves showing clinical signs of WCS. Of these, two were selected for characterization studies. Virus isolate 18115 was recovered from the lung tissue of one of the calves after three blind passages in cell culture, and virus isolate 1-1504 was recovered from kidney tissue after four blind passages.

Reference viruses included BAV serotype 3 (strain FS0213) from the Veterinary Medical Research Institute, Ames, Iowa, and BAV
serotype 7, infectious bovine rhinotracheitis (IBR) virus and parainfluenza-3 (PI-3) virus from NADC, Ames, Iowa.

**Characterization Studies**

The nucleic acid type was established using the DNA inhibitor 5-iodo-2'-deoxyuridine (IUDR) and employing the methods described by Hamparian (33) and Coria (15). Monolayers of BTU cell cultures were grown in DMEM maintenance media with IUDR (100 mcg/ml) for 16-18 hours before inoculation with virus. Similar media was put on the cell cultures after viral adsorption. Determination of nucleic acid type was based on suppression of replication of DNA viruses by IUDR. PI-3 and BAV-7 were used as RNA and DNA controls, respectively.

The method of Feldman and Wang was used to determine virus sensitivity to lipid solvents (29). A mixture of .05 ml analytical reagent grade chloroform and 1 ml tissue culture fluid containing virus was shaken for 10 minutes at room temperature. The mixture was then centrifuged at 60 xg for 10 minutes, and the clear supernatant fluid titrated for infectivity. IBR virus served as the enveloped virus control, and BAV-7 as the non-enveloped control.

Virus particle size was estimated by filtration through a series of filters of graded porosity. Purified virus diluted 100-fold in distilled water was passed through sterile 25 mm nucleopore polycarbonate membranes with pore sizes of 200 nm, 100 nm, 80 nm, and
50 nm. Particle adsorption to filters was reduced by utilizing the methods of Ver (75). The viral content of each filtrate, as well as tissue culture fluid obtained prior to filtration, was assayed.

Additionally, virus size as well as shape was determined by examination of negatively-stained particles with the transmission electron microscope (TEM). In preparation for negative staining, a flask of virus-infected cells was frozen and then thawed a total of three times. The cell culture fluid was centrifuged at 60 xg for 10 minutes and the resulting supernatant fluid was centrifuged at 100,328 xg for one hour. The supernatant fluid was discarded and the remaining material resuspended. A mixture of four drops of the virus suspension, four drops of 2 percent PTA (pH 6.8) and one drop of bovine serum albumin was made, and a drop of this material was put on a formvar-coated 300 mesh grid with a tuberculin syringe. The excess fluid was removed by touching a corner of filter paper to the edge of the grid. After drying, the grid was examined with a Zeiss EM95-2 TEM.

Sensitivity to acid was tested employing the technique established by Ketler (38). Tissue culture fluid containing virus was diluted 1:10 in DMEM adjusted to pH 3.0 or pH 7.0 with McIlvaine's buffer. The mixtures of virus and buffer were held at room temperature for 30 minutes, after which time the residual virus content was determined by titration. Reference strain viruses used
were BAV-7 and PI-3.

Heat sensitivity was tested according to the method of Wallis (77, 78). One ml of virus suspension diluted 10-fold in distilled water was delivered into tubes of uniform size and thickness. The tubes were capped and placed in a 56°C water bath that covered the tubes to within one-fourth inch from the cap. At the end of the 30 minute heating period, the tubes were quickly transferred into an ice water bath, and the samples titered for infectivity. BAV-3 and BAV-7 were used as examples of subgroup I and subgroup II controls, respectively.

Cytopathic Effect and Staining Characteristics

Monolayers of BTU cells were grown on chambered tissue culture slides for examination of viral-induced CPE. Infected and non-infected control monolayers were fixed with Bouin's fixative at various intervals before staining with hematoxylin and eosin (H & E). Slides to be used for acridine orange staining were prepared as directed in the technique developed by Dart (23). For the indirect fluorescent antibody test (FAT) the methods of Potgieter were followed (62). The conjugate used for immunofluorescent staining was fluorescein isothiocyanate (FITC), conjugated antibovine antibody of rabbit origin (Miles Laboratory). This was used in dilutions of 1:16.
These stained and mounted preparations were examined and photographed with a binocular Leitz fluorescent microscope equipped with a standard light source. Standard light microscopy was utilized for the H & E stained slides and ultraviolet light for examining acridine orange and indirect FAT preparations.

Serum-virus Neutralization Test

Serum-virus neutralization tests were performed in microtiter plates using standard procedures. Serial two-fold dilutions of the serum were made in microtiter transfer plates. To this was added an equal volume (.05 ml) of 50 TCID_{50} of test virus. The serum-virus mixtures were incubated for one to two hours at 37° C before being transferred to microtiter plates containing monolayers of the appropriate cells. The plates were maintained in a CO_{2} incubator at 37° C until examination with an inverted microscope.
CHAPTER 4

RESULTS

Nucleic Acid Type

As shown in Table 1, treatment of isolates 18115 and 1-1504 with IUDR resulted in a two to three log unit decrease in the titer of these viruses. This sensitivity is presumptive evidence that these are DNA-containing viruses. The RNA virus control, PI-3, was unaffected by similar treatment while the DNA virus control, BAV-7, was reduced in titer.

Table 1. Effect of IUDR on Virus Titer

<table>
<thead>
<tr>
<th>Virus</th>
<th>Titer of Control</th>
<th>Titer after IUDR Treatment</th>
<th>Nucleic Acid Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>18115</td>
<td>5.00</td>
<td>2.20</td>
<td>DNA</td>
</tr>
<tr>
<td>1-1504</td>
<td>4.20</td>
<td>2.20</td>
<td>DNA</td>
</tr>
<tr>
<td>BAV-7</td>
<td>3.00</td>
<td>1.20</td>
<td>DNA</td>
</tr>
<tr>
<td>PI-3</td>
<td>7.20</td>
<td>7.70</td>
<td>RNA</td>
</tr>
</tbody>
</table>

\(^a\)Titers expressed as \(\log_{10} \text{TCID}_{50}\) per/ml.

Further evidence supporting these results was provided by the acridine orange staining of virus infected cell monolayers. Both
isolates generated the intense yellow-green nuclear fluorescence characteristic of known DNA viruses. (See Figure 1.)

**Chloroform Sensitivity**

The infectivity of the viruses was not decreased by treatment with chloroform. A non-enveloped reference strain BAV-7 also remained unaffected by exposure to this lipid solvent while similar treatment affected complete inactivation of the enveloped IBR virus. These results are summarized in Table 2.

Table 2. Sensitivity to Chloroform

<table>
<thead>
<tr>
<th>Virus</th>
<th>Titer of Controls&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Titer after Chloroform Treatment&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>18115</td>
<td>8.30</td>
<td>8.03</td>
</tr>
<tr>
<td>1-1504</td>
<td>7.20</td>
<td>7.36</td>
</tr>
<tr>
<td>BAV-7</td>
<td>8.03</td>
<td>8.03</td>
</tr>
<tr>
<td>IBR</td>
<td>7.36</td>
<td>0</td>
</tr>
</tbody>
</table>

<sup>a</sup>Titers expressed as log<sub>10</sub> TCID<sub>50</sub> per/ml.

**Determination of Size**

As shown in Table 3, the viruses passed through filters with pore sizes of 200 nm, 100 nm, and 80 nm with little reduction in
Figure 1. BTU cell line stained with acridine orange, X 1500.  
A. Non-infected cell culture control after 5 days of incubation.  
B. Cell culture infected with PI-3 after 3 days of incubation. The orange-red fluorescence is characteristic of single-stranded RNA viruses.  
C. Cell culture infected with 18115 after 5 days of incubation. Cells infected with both 18115 and 1-1504 showed the yellow-green fluorescence characteristic of double-stranded DNA viruses.
titer, but were completely retained by filters of 50 nm pore size. This implies that the diameter of the virus particles range somewhere between 50 nm and 80 nm. From TEM micrographs taken of the negatively stained virus particles, the approximate diameter of the virions was calculated to be 70 nm. (See Figure 2.)

Table 3. Determination of Size

<table>
<thead>
<tr>
<th>Virus</th>
<th>Unfiltered Control&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Filtrates&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>200 nm</td>
<td>100 nm</td>
</tr>
<tr>
<td>18115</td>
<td>3.34</td>
<td>3.03</td>
</tr>
<tr>
<td>1-1504</td>
<td>5.70</td>
<td>6.30</td>
</tr>
</tbody>
</table>

<sup>a</sup>Titers expressed as log<sub>10</sub> TCID<sub>50</sub> per/ml.

Acid Sensitivity

The viruses were characterized as being acid stable based upon their retention of infectivity upon exposure to pH 3.0 for 30 minutes. The results of acid sensitivity are summarized in Table 4.
Figure 2. Electron micrograph of 18115 demonstrating the cubic symmetry characteristic of both isolates. Negative stain for this preparation was potassium phosphotungstate pH 5.0, pH 6.8, pH 7.5 and uranyl acetate pH 4.4.
Table 4. Sensitivity to Treatment at pH 3.0

<table>
<thead>
<tr>
<th>Virus</th>
<th>Titer at pH 7.0(^a)</th>
<th>Titer after Treatment at pH 3.0(^a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>18115</td>
<td>4.70</td>
<td>5.70</td>
</tr>
<tr>
<td>1-1504</td>
<td>5.03</td>
<td>4.36</td>
</tr>
<tr>
<td>PI-3</td>
<td>5.70</td>
<td>0</td>
</tr>
<tr>
<td>FSO-213</td>
<td>3.70</td>
<td>3.70</td>
</tr>
</tbody>
</table>

\(^a\)Titers expressed as \(\log_{10} TCID_{50}\) per/ml.

Susceptibility to Heat

As shown in Table 5, the isolates exhibited partial thermal inactivation upon exposure to 56\(^0\) C for 30 minutes. The virus representative of the subgroup I BAV (strain FSO-213, BAV-3), showed complete reduction of titer, while the subgroup II BAV (BAV-7) showed only partial heat susceptibility. In this respect the isolates were more similar to the subgroup II BAV.
Table 5. Thermostability

<table>
<thead>
<tr>
<th>Virus</th>
<th>Titer at Room Temperature&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Titer at 56° C for 30 Minutes&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>18115</td>
<td>6.36</td>
<td>2.20</td>
</tr>
<tr>
<td>1-1504</td>
<td>7.36</td>
<td>2.36</td>
</tr>
<tr>
<td>BAV-7</td>
<td>7.20</td>
<td>3.36</td>
</tr>
<tr>
<td>FSO-213</td>
<td>4.70</td>
<td>0</td>
</tr>
</tbody>
</table>

<sup>a</sup>Titters expressed as $\log_{10} TCID_{50}$ per/ml.

Serologic Analysis

Antiserum to BAV-7 neutralized the infectivity of isolates 18115 and 1-1504. No cross reaction with BAV antisera of any other type was demonstrated. The results of serologic characterization are summarized in Table 6.
Table 6. Serologic Characterization

<table>
<thead>
<tr>
<th>Antisera</th>
<th>18115</th>
<th>1-1504</th>
<th>BAV-3(FSO-213)</th>
<th>BAV-7</th>
</tr>
</thead>
<tbody>
<tr>
<td>BAV-1&lt;sup&gt;a&lt;/sup&gt;</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>BAV-2&lt;sup&gt;a&lt;/sup&gt;</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>BAV-3&lt;sup&gt;a&lt;/sup&gt;</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>BAV-3&lt;sup&gt;c&lt;/sup&gt; (WBRI)</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>BAV-3&lt;sup&gt;c&lt;/sup&gt; (5C)</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>BAV-7&lt;sup&gt;b&lt;/sup&gt;</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>BAV-7&lt;sup&gt;a&lt;/sup&gt;</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>BAV-8&lt;sup&gt;a&lt;/sup&gt;</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

<sup>a</sup>Provided by M. F. Coria, NADC, Ames, Iowa. (Produced in rabbit.)
<sup>b</sup>Provided by M. F. Coria, NADC, Ames, Iowa. (Produced in calf.)
<sup>c</sup>Provided by D. E. Mattson, Oregon State University, Corvallis, Oregon. (Produced in rabbit.)
Characterization of CPE in Cell Cultures

In unstained cell preparations, the first sign of CPE in monolayers infected with the isolates was the increased refractibility of cells that had rounded or become spindle shaped. Formation of cytoplasmic bridges and the gradual detachment of these affected cells would progress to the eventual destruction of the monolayer. (See Figure 3.)

H & E stained preparations of infected cell cultures evinced the production of multiple, eosinophilic, intranuclear inclusion bodies. (See Figure 4.) These were most frequently round in shape, and often appeared to be surrounded by a clear, non-staining halo.

The susceptibility of a number of bovine cell types to viral-induced CPE was examined. Primary and low-passage bovine fetal cells derived from lung, salivary gland, turbinate, testicle and thyroid tissues developed CPE within seven days when inoculated with isolates 18115 and 1-1504. No CPE was demonstrated by either fetal kidney cell cultures or the bovine kidney cell line when similarly infected.

Cells were fixed for indirect immunofluorescent staining while in the early stages of infection. The characteristic fluorescence that was generated in infected cells treated with antiserum to BAV-7 serum appeared to be of greatest intensity along the perimeter of the nucleus. (See Figure 5.)
Figure 3. Unstained BTU monolayers, X 625.
A. Non-infected cell culture control.
B. Cytopathic effect typically produced by 18115 and 1-1504 in BTU cell cultures.
C. Microtumor sometimes seen in cell cultures infected with 18115 and 1-1504.
Figure 4. Examples of various forms of inclusion bodies evident in the nucleus of cell cultures infected with 18115 and 1-1504. Hematoxylin and eosin stain, X 3750.
A. Non-infected cell culture control after 3 days of incubation.
B. Infected cell culture after 3 days of incubation.
C. Infected cell culture after 5 days of incubation. Notice the clear non-staining halo surrounding the inclusion bodies.
D. Infected cell culture after 5 days of incubation.
Figure 4. E-H. A series of micrographs showing progressive degeneration in cells infected with 18115 and 1-1504, 4 and 5 days post inoculation.
Figure 5. Immunofluorescence evident in BTU cell cultures infected with 18115 and 1-1504 18 hours post inoculation and indirectly stained with fluorescein isothiocyanate, X 2000. Notice the intense staining in the area of the nuclear envelope.
CHAPTER 5

DISCUSSION

The condition referred to as WCS has been a recognized problem in cattle herds of Montana and Idaho for over 15 years. Isolates 18115 and 1-1504 were obtained as part of an ongoing research effort in this area to delineate factors involved in the etiology of WCS. As determined in this work, the physicochemical and biological properties of virus isolates 18115 and 1-1504 are consistent with those required for inclusion in the AV group (59).

Replication of these viruses was inhibited when grown in the presence of the thymidine analogue, IUDR. This, and the characteristic apple-green fluorescence produced in acridine orange stained cells infected with the isolates is indicative of their DNA content (83).

The isolates were shown, by way of TEM examination and ultrafiltration to be spherical particles approximately 70 nm in diameter. By virtue of their chloroform resistance, they manifested a lack of essential lipid, thereby demonstrating the absence of a protective envelope (83).

The value of heat sensitivity as a characteristic in BAV classification has been questioned (11). This attribute was nonetheless tested for, and both viruses were found to be only partially sensitive to heat, in that they exhibited a decrease in
titer when exposed to 56°C for 30 minutes, but not complete inactivation. In this respect, according to Bartha's original classification scheme, a closer relationship is evidenced to the subgroup II BAV than to the subgroup I BAV (3). Additionally, the isolates were able to withstand acid conditions, remaining stable after being held at pH 3.0 for 30 minutes.

The behavior of the viruses in cell culture is an important parameter used in the division of BAV into two subgroups. Isolates 18115 and 1-1504 were recovered in BTU cell cultures after several blind passages. They also exhibited growth in primary and low passage bovine fetal lung, salivary gland, turbinate, testicle, and thyroid cell cultures, but failed to produce CPE in bovine kidney cells. The CPE produced in susceptible cells was typical of that seen with AV. In BTU cells stained with H & E, the viruses generated multiple inclusions in the nucleus which were regular, often round, in shape. These attributes are consistent with those of the subgroup II BAV (3).

The result of serologic analysis was the neutralization of isolates 18115 and 1-1504 with hyperimmune serum directed against serotype 7 BAV. BAV appear to be neutralized only by homologous sera, so it is assumed that these viruses are serotype 7 BAV (55). Additionally, positive nuclear immunofluorescence was accomplished when cells infected with these isolates were treated with antiserum to BAV-7.
In general, BAV appear to be agents responsible for a variety of disease conditions that are manifested primarily as disorders of the respiratory and enteric systems in calves. The usual gross lesions include areas of consolidation, collapse, and emphysema in the lungs of afflicted animals, and microscopic examination demonstrates proliferative bronchiolitis, necrosis and bronchiolar occlusion. Although the pneumoenteritis complex has often been noted in weak calves (31), the usual presentation of symptoms accorded WCS is at variance with the general picture of BAV infections. Briefly, calves showing clinical signs of WCS are too debilitated to stand and nurse, they exhibit a lameness and swelling about the joints indicative of a polyarthritic state, and their muzzles are often red and encrusted. The most outstanding gross lesions are subcutaneous hemorrhages and edema in the tissues of the extremities, and hemorrhagic synovial fluid containing varying amounts of fibrin in the swollen joints (12).

It would appear that there is little correlation between the clinical signs and lesions seen in WCS and those present in BAV infections. However, there is another aspect of AV pathogenesis in calves. Investigators have described lesions in calves infected with AV which include many areas of petechial and ecchymotic hemorrhage throughout as well as characteristic AV intranuclear inclusions in the endothelial cells and pericytes of blood vessels (11, 17, 72). This suggests that the calves were viremic at some stage of the infection.
and that the virus was disseminated by the bloodstream. It is of interest to note that among such reports are included experimental infections of calves with AV isolated from calves with WCS. Theoretically, many of the clinical signs and lesions thought to be pathognomonic for WCS can be explained should such a viremic state occur.

The consequences of endothelial damage previously described would be exposure of the collagen present in vessel walls, and the release of tissue factor. Platelet adherence to sites of endothelial damage would occur, and the triggering of the intrinsic and extrinsic hemostatic pathways could culminate with disseminated intravascular coagulation (DIC). Endothelial damage does, in fact, figure as a primary mechanism in the initiation of DIC (48). This condition produces a bleeding diathesis that can be manifested as petechial and ecchymotic hemorrhage into the subcutis and mucous membranes (48, 66). The increased vascular fragility brought on by this state could result in hypovolemia (blood volume failure), and by leakage of intravascular fluids into the tissues, produce the edema so characteristically present in weak calves. Plausibly, this hypovolemia could cause shock sufficient to kill the calf. Using this model of a disease mechanism as a bias, the clinical signs of WCS such as the characteristic pattern of hemorrhage, edema, and fibrin-containing hemorrhagic synovial fluid argue in support of a hemostatic imbalance as a
mechanism for pathogenesis.

Favored sites of implantation for blood borne organisms are the heart valve, meninges, kidneys, and significantly, the joint spaces (66). Vascular damage in this area and the subsequent pressure of fluid accumulation in the joint spaces could produce a painful arthritis in the young animal.

There are several factors which lend credence to this hypothetical model. The proclivity AV show toward wreaking extensive endothelial damage is well documented in the canine species. In the disease infectious canine hepatitis, AV replicates in the vascular endothelium and hepatocytes, initiating DIC. Chief manifestations of the disease are widespread hemorrhage, hepatocellular necrosis, and hemostatic defects (41).

BAV are capable of producing latent infections in animals. Immunosuppression of such an animal can induce the expression of the virus as an active infection (79). In the case of WCS, such immunosuppressive effects might be produced by the secretion of corticosteroids which occur around the time of partuition (47). Additionally, BVD virus, which has been isolated from weak calf tissues, sometimes in conjunction with BAV, has known immunosuppressive capabilities (37, 60). Levels of serum hydrocortisone secreted by both the dam and fetus rise immediately prior to partuition (47). Serum hydrocortisone concentrations also increase as response to
physiologic stress (i.e., severe weather conditions). Studies of IBR-infected calves injected with hydrocortisone showed increased concentrations of serum interferon, and interestingly, calves with the highest levels of interferon also had the highest levels of viremia (16). These facts support the conceivability of a viremic state due to AV.

Several unsuccessful attempts have been made to reproduce WCS by experimental inoculation of young calves with BAV-7 (17, 26). A recent publication by investigators at the University of Idaho describes the experimental intraamnionic exposure of bovine fetuses to BAV-7. Two of the three fetuses infected in this manner were born depressed and unable to stand and nurse. Clinical signs included hyperemia and petechiation of the oral mucosa and conjunctival membrane. Gross postmortem lesions were non-specific and consisted of petechial and ecchymotic hemorrhages and edema of the epithelial surface of the rumen and the mucosal surface of the abomasum and small intestine. Turbid synovial fluid was evident in the hock joints. The most consistent microscopic lesions observed in the tissues from infected calves were vasculitis and perivasculitis of varying severity in most tissues. Intranuclear inclusion bodies were evident in pericytes, hepatocytes, macrophages, and the epithelial cells of the adrenal cortical sinusoids and renal tubular epithelium. Although a great many of these clinical signs and pathologic changes are
consistent with those of WCS, the author concluded that the disease produced was not WCS (73).

It is probable that in the evolution of such a disease process there is a delicate timing in the sequence of events and the interplay of factors involved that must be achieved before it can be experimentally reproduced. Other factors may be shown to be necessary contributors to the production of this disease, but it is likely that BAV will yet prove to be of etiologic importance in the pathogenesis of WCS.
Two virus isolates designated 18115 and 1-1504 were recovered from the tissues of calves showing clinical signs of WCS. As determined by physicochemical and biological characterization, the properties of these two viruses are consistent with those required for inclusion in the AV group. Antigenic relation to the subgroup II serotype 7 BAV was demonstrated by the isolates in serum-virus neutralization tests and indirect fluorescent antibody studies.

Replication of these viruses was inhibited when grown in the presence of IUDR. Characteristic yellow-green fluorescence was generated in infected cell cultures stained with acridine orange.

Ultrafiltration studies and TEM examination of the isolates evinced spherical particles approximately 70 nm in diameter. No decrease in viral infectivity was accomplished upon exposure of the isolates to chloroform, thereby demonstrating the lack of an enveloping lipid membrane.

Partial heat sensitivity was exhibited by isolates 18115 and 1-1504. Additionally, they were able to withstand acid conditions, remaining stable after being held at pH 3.0 for 30 minutes.

The isolates were recovered in BTU cell cultures after several blind passages and produced CPE in several bovine cell types including low passage fetal lung, salivary gland, turbinate, testicle, and
thyroid cell cultures. No CPE was demonstrated in primary kidney or
GBK cell cultures. In the nucleus of BTU cells stained with H & E,
the viruses generated multiple inclusions which were generally round
in shape. The CPE produced in susceptible cells was typical of that
produced by AV.


35. Ide, P. R., R. G. Thompson, and W. J. B. Ditchfield. 1969.


The characterization of two bovine adenoviruses isolated from calves exhibiting clinical and pathological signs of weak calf syndrome.