

1     **OCCURRENCE OF BLUETONGUE VIRUS ANTIBODIES IN CATTLE AND SHEEP**  
2                     **IN OGUN AND OSUN STATES, NIGERIA**

5                                     **ABSTRACT**

6     Bluetongue is an infectious, arthropod-borne viral disease principally affecting ruminants. The  
7     occurrence of bluetongue virus (BTV) antibodies in sheep and cattle from backyard farms, cattle  
8     markets and abattoirs in Ogun and Osun states of Nigeria was investigated. Three hundred and  
9     forty (340) plasma samples comprising 205 from sheep and 135 from cattle were collected  
10    noting the sex, breed and age of the animals. The samples were screened with a commercial  
11    competitive ELISA kit that detects BTV antibodies in ruminant plasma or serum. All cattle  
12    tested from both states were positive for BTV antibodies giving a seroprevalence of 100% while  
13    95% seroprevalence was obtained for sheep. In Ogun state, prevalence rates of 90.5% and 98%  
14    were obtained for male and female sheep respectively while 95.6% and 95% prevalence were  
15    also obtained for male and female sheep respectively in Osun state. Based on breed, 94%, 95%,  
16    95% and 96% prevalence were obtained for Yankasa, Balami, Ouda and West African Dwarf  
17    sheep respectively in Ogun state while 93%, 95.5%, 100% and 93% prevalence were obtained  
18    for Yankasa, Balami, Ouda and WAD sheep respectively in Osun state. Furthermore, prevalence  
19    rates of 92% and 96.7% were obtained for age groups of  $\leq 1$  year and  $> 1$  year respectively in  
20    Ogun state, while prevalence rates of 96% and 94.7% were obtained for age groups of  $\leq 1$  year  
21    and  $> 1$  year respectively in Osun state. Since vaccination against bluetongue disease is not  
22    practiced in Nigeria, the detection of high prevalence of BTV antibodies observed in apparently  
23    healthy animals in this study indicates natural, albeit subclinical, infection with the virus and  
24    sustained activity of the *Culicoides* vector. These findings suggest that bluetongue is widespread  
25    in Nigeria and highlight the need for continuous surveillance of the disease in the country as well  
26    as isolation, identification and characterisation of currently circulating BTV strains in Nigeria.

27    Keywords: Bluetongue virus, prevalence, cattle, sheep, Ogun state, Osun state.

28                     **1. INTRODUCTION**

29    Bluetongue is an infectious, arthropod-borne viral disease principally affecting ruminants. Other  
30    names for this disease include catarrhal fever, sore muzzle, muzzle disease and pseudo-foot-and-

31 mouth disease [1]. It is caused by the pathogenic virus, bluetongue virus (BTV) of the genus  
32 *Orbivirus*, family Reoviridae [2]. It is a non-enveloped virus and the genome is made of 10  
33 segments of double-stranded RNA [3][4].

34 The disease was first described in the Cape colony of Southern Africa after Merino sheep were  
35 introduced into the region in the late 18<sup>th</sup> century, and was subsequently recognized in other parts  
36 of Africa, Europe, the Middle East, Indian subcontinent, the Americas and Asia [5]. Twenty six  
37 serotypes of BTV are recognized globally [6], and the virus has now been isolated on all  
38 continents except Antarctica [5].

39 Orbiviruses are the cause of important and apparently emerging arthropod-borne viral (arboviral)  
40 diseases of livestock, including bluetongue virus, African horse sickness virus, equine  
41 encephalosis virus and epizootic haemorrhagic disease virus that are all transmitted by  
42 haematophagous *Culicoides* insects [5]. Arboviruses are important causes of disease in humans  
43 and animals, and it is conceivable that climate change will increase the distribution and severity  
44 of arboviral diseases. Recent changes in the global distribution and nature of BTV infection have  
45 been especially dramatic, with spread of multiple serotypes of the virus in almost all parts of the  
46 world including Europe and USA with previously exotic virus serotypes [5]. Although climate  
47 change has been incriminated in the emergence of BTV infection of ungulates, the precise role of  
48 anthropogenic factors and the like is less certain [5].

49 The central role of flying insects in bluetongue epidemiology means that the prevalence of the  
50 disease is governed by ecological factors that favour insect survival such as high rainfall,  
51 temperature, humidity and soil characteristics [7] [8]. Bluetongue outbreaks generally occur  
52 seasonally and in warm climates. In the temperate regions, *Culicoides* vector infects most  
53 ruminant species during mid-summer to early fall when it is active [9]. The infection subsides  
54 when temperatures drop and hard frosts kill the adult midge vectors [9]. In the tropical and  
55 subtropical regions, however, infection occurs throughout the year as the vector is present year-  
56 round [10]. In the absence of competent vector populations, animal to animal transmission is not  
57 capable of maintaining an endemic state [11].

58 Bluetongue is not known to be harmful to humans. However, it causes considerable damage to  
59 livestock populations. The virulence of BTV varies quite markedly; even strains with matching

60 serotypes have variable virulence [9]. It is a transboundary disease [12][13], and the  
61 epidemiological situation in one country can affect neighboring countries while national  
62 measures tend not to be sufficient to control its spread. BTV infects many domestic and wild  
63 ruminants including sheep, goats, cattle, buffaloes, deer, antelopes, bighorn sheep and camels  
64 [14][15]. Sheep are the main hosts and exhibit the most clinical disease [16]. Severe disease can  
65 also occur in some wild ruminants including white-tailed deer (*Odocoileus virginianus*),  
66 pronghorn (*Antilocapra Americana*) and desert bighorn sheep (*Ovis Canadensis*) [15]. Other  
67 ruminants like cattle, goats, camels, buffaloes, and some wild ruminants typically have sub-  
68 clinical disease. Cattle are considered amplifiers and maintenance hosts [16].

69 Bluetongue is characterized by BTV-induced vascular injury that results in haemorrhage and  
70 ulceration of the mucous membranes in the upper portion of the gastro-intestinal tract, coronitis  
71 and lameness, facial and intramuscular oedema, pleural and pericardial effusions, pulmonary  
72 oedema and necrosis of the skeletal and cardiac muscle [17] [18][19]. Swelling of the lips and  
73 cyanosis of the tongue give the tongue its typical blue appearance, though this sign is confined to  
74 a minority of the animals. Excessive salivation and nasal discharge are also observed. In sheep  
75 lameness due to coronitis can lead to knee-walking. In cattle, constant changing of position of  
76 the feet gives bluetongue disease the nick-name, the dancing disease [20]. Clinical bluetongue is  
77 mainly seen in certain fine wool and mutton breeds of sheep which, seemingly by chance, are  
78 found in countries at the limits of the virus distribution (Spain, Portugal, Turkey, Cyprus, USA,  
79 South Africa) [21]. Elsewhere, the disease is likely whenever similarly susceptible animals are  
80 transported to countries within the bluetongue enzootic areas e.g. Nigeria [22], Cameroon [23]  
81 and Indonesia [24]. Since indigenous, unimproved sheep, goat and cattle breeds are usually  
82 highly resistant to the clinical effects of the infection, the vast majority of bluetongue episodes  
83 throughout the world are completely silent [25]. However, considering the recent alarming  
84 global spread of BTV serotypes, it is no longer uncommon to see mortality rates approaching 50  
85 - 100% in susceptible flocks. Other losses are due to morbidity and the need to provide care for  
86 the sick animals. Costs associated with morbidity include weight loss, reduced milk yield,  
87 abortion and associated veterinary costs. This covert presence of the virus, alternating with  
88 occasional outbreaks of severe disease has had a considerable adverse effect upon international  
89 trade of bovine and ovine species, and their germ plasms, as countries free from bluetongue  
90 attempt to maintain that status [25].

91 Diagnostic testing for BTV can be very difficult as the virus cross-reacts with many antigenically  
92 related viruses including palyam virus and the viruses that cause epizootic haemorrhagic disease  
93 of deer and African horse sickness [16][10]. However, the enzyme-linked immunosorbent assay  
94 (ELISA) has proven to be the best serologic test for BTV antibody detection, and can be used to  
95 decrease the chances of cross-reactions [26]. Also, a monoclonal antibody-based competitive  
96 ELISA that can distinguish antibodies to virus in the bluetongue serogroup from antibodies to the  
97 other related viruses has been developed [26].

98 Obi *et al.* [27] found that bluetongue virus antibodies were widespread in southern Nigeria. The  
99 serum samples collected between 1979 and 1981 were examined for bluetongue virus  
100 precipitating antibodies in agar gel precipitin test, and 58% of the sheep and 50% of the goat  
101 samples were seropositive. Recently, Oluwayelu *et al.* [28] obtained a prevalence of 89.2%,  
102 88.0% and 84.4% for sheep, goats and cattle respectively in Oyo state of Nigeria using a  
103 commercial bluetongue ELISA kit.

104 Considering the fact that bluetongue is a re-emerging disease [9](Purse *et al.*, 2005), this study  
105 was carried out to investigate the seroprevalence of the disease in cattle and sheep in Ogun and  
106 Osun States, Nigeria, and to study its effect on sex, breed and age of the animals.

107

## 108 **2. METHODOLOGY**

### 109 **2.1 Experimental Design and Sample Size Determination**

110 The design of this study was based on descriptive study. Sample size was determined according  
111 to the method previously described [29]. The experiment was aimed at comparing the result  
112 obtained from Ogun state to that of Osun state, putting into consideration the age, sex and breed  
113 of the animals. The species of animal of interest were cattle and sheep.

### 114 **2.2 Study Areas**

115 The areas considered for this study are Ogun and Osun States. In Ogun state, the samples were  
116 collected from two different locations which are Abeokuta and Ijebu-Ode while in Osun state  
117 samples were collected from Osogbo, Ejigbo and Iwo. For cattle, the sites for sample collection

118 were ranches, cattle markets and abattoirs while that for sheep were sheep markets and backyard  
119 farms in Osun and Ogun states.

120 Ogun state climate follows a tropical pattern with the rainy season starting from March and  
121 ending in November, followed by dry season. The mean annual rainfall varies from 128cm in the  
122 southern parts of the state to 105cm in the northern areas [30]. The average monthly temperature  
123 ranges from 23<sup>0</sup>C in July to 32<sup>0</sup>C in February. The northern part of the state is mainly of derived  
124 Savannah vegetation, while the central part falls in the rain forest belt [30].

125 The climate of Osun state is slightly similar to that of Ogun state, and has a covering of the  
126 tropical rain forest. However, Osun state has an annual rainfall of about 60cm [31]. The state  
127 climate is less humid when compared to Ogun state although the effects of the harmattan winds  
128 are strongly felt in the dry season. The average monthly temperature ranges from 24.5<sup>0</sup>C in July  
129 to 28<sup>0</sup>C in February [31].

130 The two states are located in the southwestern part of Nigeria and are characterized by a long  
131 rainy season, high humidity and temperature favourable for the breeding of the *Culicoides*  
132 vectors of bluetongue virus. Since the ambient temperature in Africa allows the survival of the  
133 *Culicoides* vectors of BT from January to December, sampling was done without a special  
134 interest on a specific season of the year. Several species of *Culicoides* that feed on domestic  
135 ruminants have been identified in these areas [32][33].

### 136 **2.3 Collection of blood samples**

137 Simple random sampling was done. The age, sex and breed of the animals were taken in the  
138 process of sampling. The ages of the animals in the farms were determined through oral  
139 interview with the farmers, while age estimation was done for the animals in markets and  
140 abattoirs. A total number of 340 samples were collected from 205 sheep and 135 cattle. About  
141 5ml of blood was collected aseptically from each animal by venipuncture of the jugular vein  
142 using sterile Monovette EDTA tubes (Sarstedt, Germany). All samples collected were  
143 transported to the laboratory under a cold chain sustained with ice packs. Antibiotics  
144 (Gentamycin and Amphotericin B) were added to the blood samples after which they were  
145 centrifuged at 2000 rpm for 5 minutes. The plasma (supernatant) were collected into sterile  
146 Eppendorf tubes, labeled and stored in a freezer at -20<sup>0</sup>.

147 **2.4 Testing the samples**

148 The 340 samples collected (135 cattle and 205 sheep) were screened for the presence of anti-VP7  
149 BTV antibodies using the POURQUIER<sup>®</sup> BTV cELISA kit. This was done according to the  
150 manufacturer's instruction.

151 **2.5 Statistical analysis**

152 The data obtained in this study was analyzed using Chi-square test and the level of significance  
153 was set at 95% (0.05). The analysis was done using SPSS version 21.

154

155 **3. RESULTS**

156 Out of the 340 plasma samples from cattle and sheep screened for the BTV antibodies in Ogun  
157 and Osun states, 330 samples were positive which gave an overall seroprevalence rate of 97%.  
158 Furthermore, 195 of the 205 plasma samples from sheep screened were positive which gave a  
159 seroprevalence rate of 95%. All the 135 samples from cattle tested were positive, resulting in  
160 100% seroprevalence rate, irrespective of sex, breeds or age groups. There was a significant  
161 difference in BTV antibody prevalence between the cattle and sheep in Ogun and Osun states  
162 (Table 1).

163

164 **Table 1. Overall prevalence of BTV antibodies in cattle**  
165 **and sheep in Ogun and Osun States**

166

167 Species	168 Number of samples tested	169 Positive
170 Cattle	135	135 (100%)
171 Sheep	205	195 (95%)
172 Total	340	330 (97%)

173

174 Table 2 shows the seroprevalence rates of BTV for sheep in Ogun and Osun states to be 95.0%  
 175 and 95.3% respectively. There was no significant difference. The table also shows the prevalence  
 176 according to sex. In Ogun state, out of 99 samples from sheep screened for BTV antibodies, 38  
 177 (90.5%) and 56 (98.0%) tested positive for males and females respectively, and the difference  
 178 was not significant ( $P < 0.05$ ). In Osun state, out of 106 samples from sheep screened, 44  
 179 (95.6%) and 57 (95%) tested positive for males and females respectively, and the difference was  
 180 not significant ( $P < 0.05$ ).

181 **Table 2: BTV antibody prevalence in cattle and sheep based on sex**  
 182 **(Ogun and Osun State)**

	<b>Ogun</b>		<b>Osun</b>	
Sex	Cattle	Sheep	Cattle	Sheep
Male	33/33(100%)	38/42(90.5%)	38/38(100%)	44/46(95.6%)
Female	35/35(100%)	56/57(98.0%)	29/29(100%)	57/60(95.0%)
<b>Total</b>	<b>68/68(100%)</b>	<b>94/99(95.0%)</b>	<b>67/67(100%)</b>	<b>101/106(95.3%)</b>

183  
 184 Table 3 shows results of the BTV antibody prevalence according to the breeds of sheep in Ogun  
 185 and Osun states. There was no significant difference in the seroprevalence rates among the  
 186 breeds in the two states.

187 **Table 3. BTV antibody prevalence in sheep according to breeds in Ogun and Osun States.**

	<b>Ogun</b>	<b>Osun</b>
--	-------------	-------------

<b>Breed</b>	<b>Positive (%)</b>	<b>Positive (%)</b>
Yankasa	33/35(94%)	28/30(93%)
Balami	19/20(95%)	21/22(95.5%)
Ouda	20/21(95%)	25/25(100%)
WAD	22/23(96%)	27/29(93%)
<b>Total</b>	<b>94/99(95%)</b>	<b>101/106(95%)</b>

188

189

190 Comparison of the BTV seroprevalence rates based on age groups in Ogun state showed that  
 191 sheep less than twelve months of age had 92% BTV seroprevalence while those greater than  
 192 twelve months had a seroprevalence of 96.7%. In Osun state, sheep less than twelve months of  
 193 age had a seroprevalence of 96% while those greater than twelve months had 94.7%  
 194 seroprevalence (Table 4). There was no significant difference in BTV antibodies among the age  
 195 groups.

196

**Table 4. BTV antibody prevalence for sheep in Ogun and Osun States  
 According to age groups.**

197

<b>Age group</b>	<b>Ogun</b>	<b>Osun</b>
1 – 12 months	35/38(92%)	47/49 (96%)
> 12 months	59/61(96.7%)	54/57 (94.7%)
<b>Total</b>	<b>94/99 (95%)</b>	<b>101/106 (95.3%)</b>

198

199

#### 4. DISCUSSION

The high prevalence of BTV infection observed in cattle and sheep in this study indicates the possible emergence of bluetongue disease in Nigerian cattle and sheep populations. In the present study, 100% and 95% seroprevalence rates were obtained from cattle and sheep respectively in Ogun and Osun states which are located in southern Nigeria that is characterized by high rainfall, high humidity and temperature favourable for the breeding of *Culicoides* vectors of Bluetongue disease [32]. There is constant feeding of the virus vectors on the vertebrates resulting in continual spread of the disease [32]. The prevalence rates obtained in this study for sheep in the two states were slightly different amongst the different parameters such as sex, breed and age group. Our prevalence is higher than 28.9% in sheep reported by Taylor and McCausland [34] in Northern Nigeria. It is also higher than the 58% prevalence obtained by Obi *et al* [27] in Southern Nigeria. Furthermore, the result is also higher than that of Oluwayelu *et al.* [28] which reported BTV seroprevalence rates of 84.4% and 89.2% for cattle and sheep respectively in Oyo state. Other reports on the presence of BTV were made over four decades ago [35][36][37].

Preventive measures are not practiced in Nigeria to reduce exposure of the animals to *Culicoides* vectors irrespective of the husbandry system of Agriculture. Nomadism, free-range and backyard systems of farming are the common systems of animal husbandry practices in southwestern Nigeria. Animals are often allowed to roam, almost equally exposed to the vectors of the disease before being taken to the market for sale or slaughter. In addition, even when the animals' movements are restricted, midges easily fly into the pens and ranches through any available space to feed on the confined animals. The 100% seroprevalence rate from cattle in this study could be due to the fact that different breeds of cattle from different parts of the country are usually brought together in the markets before they are slaughtered. The breeds of cattle considered in this study are White Fulani, Red bororo and Sokoto gudali. This convergence could allow for high detection rates of BTV antibodies as the vectors feed on infected animals, become infected themselves and then spread the virus to other ruminants. This could lead to the generation of genetic diversity due to reassortment between BTV strains or serotypes introduced by animals from different parts of the country [9][38]. Moreover, there are so many porous borders in Nigeria, with influx of cattle, sheep and goats from neighbouring West African

231 countries. There is no bluetongue disease monitoring and surveillance programme in Nigeria,  
232 neither is there any quarantine measure to ensure that the animals coming into the country are  
233 free of the disease.

234 Cattle are regarded as the maintenance hosts [16]. It has been documented that the disease does  
235 not often manifest in cattle but with a prolonged viraemic period which increases the likelihood  
236 of feeding *Culicoides* vectors getting infected [16]. Also, the preference for cattle to the other  
237 species of ruminants by the *Culicoides* vectors may be a contributory factor [39]. Moreover, it  
238 has been reported that African *Culicoides* lay their eggs in deposited dung near the ruminant  
239 habitat. Cattle dung, unlike that of sheep, can serve as an efficient environment for breeding of  
240 the midges [40]. In this study, the samples from cattle were collected from abattoirs, cattle  
241 markets, ranches or farms operating semi-intensive management systems. In most cases, the  
242 rearing grounds were covered with animal dung and moist soils, which served as a good habitat  
243 for the breeding of *Culicoides* midges.

244  
245 Global warming and prolonged rainy season in Nigeria cannot be left out. It has been proposed  
246 that the recent environmental changes have facilitated expansion of the range of known vectors  
247 such as *Culicoides imicola* [41]. The insect vectors, biting midges, prefer warm, moist conditions  
248 and are in their greatest numbers and most active after rains [8]. Increase in ambient temperature  
249 increases the feeding and breeding activities of *Culicoides* midges which results in increased  
250 BTV transmission rate [8]. Therefore, the high BTV seroprevalence can be attributed to the  
251 sustained activity of *Culicoides* vector in the study areas.

252 Reports have shown that the recent spread of several arboviral diseases appears to have resulted  
253 from anthropogenic and social factors, rural-urban drift and movement (translocation) of virus  
254 infected vectors. *Culicoides* vectors can be transported on wind to long distances of up to several  
255 hundred kilometers [42][43]. When the enzootic foci of bluetongue disease are geographically  
256 close by, the virus can easily be introduced to the country through the wind-borne insects.  
257 Therefore, there is a possibility of occurrence of genetic reassortments between the new strains  
258 of the virus introduced by the insects and the existing strains in the locality, resulting in a variant  
259 that may be more compatible with the *Culicoides* species in that region. This variant may be  
260 more virulent, and are spread by the vectors to the susceptible hosts.

261

262 It is noteworthy that despite the fact that majority of the animals screened in this study were  
263 apparently healthy, high seroprevalence rates of the virus were obtained. This shows that clinical  
264 manifestation of the disease is not often encountered in Nigeria, even among the sheep  
265 population. This is consistent with other reports that bluetongue is endemic in Nigeria and the  
266 indigenous breeds of sheep exhibit sub-clinical manifestation of the disease [37][44][21]. Also,  
267 Mellor [25] reported that since indigenous, unimproved sheep, goat and cattle breeds are usually  
268 resistant to the clinical effects of the disease, the vast majority of bluetongue episodes are  
269 completely silent. This could therefore account for the high BTV seroprevalence obtained this  
270 study in the absence of clinical disease.

271

272 Since vaccination against bluetongue is not practiced in Nigeria, the detection of high BTV  
273 seroprevalence rates indicates natural infection with the virus, as well as sustained activity and  
274 possible increased competence of the *Culicoides* vectors in transmission of the disease.  
275 Moreover, since field strains of BTV are known to exhibit genetic heterogeneity, it is possible  
276 that the genetic profile of circulating Nigerian field BTV strains could have been altered over the  
277 past three decades resulting in the emergence of hitherto absent serotypes that could have  
278 contributed to the present high antibody prevalence. Therefore, there is a possibility of the  
279 indigenous breeds of ruminants becoming susceptible to the emerging and constantly changing  
280 strains in Nigeria overtime. It has been reported that BTV serotype 8, which was previously  
281 silent is currently causing devastating economic losses in ruminant industries worldwide, and  
282 more especially in Northern Europe [18]. BTV-8 is mainly transmitted by *Culicoides imicola*  
283 which is the traditional Asian/African species of *Culicoides* [5]. It is possible that this BTV  
284 serotype and its principal vector (*Culicoides imicola*) is present in Nigeria without any associated  
285 disease outbreak yet.

286

## 287 5. CONCLUSION

288 The high prevalence of BTV antibodies in cattle and sheep as demonstrated in this study  
289 suggests that BTV infection is widespread in Nigeria and stresses the need for continuous  
290 surveillance of the disease in domestic ruminant populations in Nigeria in order to track the  
291 possible evolution of the virus. The fact that animals are subclinically infected does not mean

292 that the disease should be over looked in Nigeria. Also, the fact that most of the cattle screened  
293 in this study were brought from the northern part of the country suggests that bluetongue may be  
294 highly prevalent in the country. Moreover, considering the genetic heterogeneity of field strains  
295 of BTV that occurs as a consequence of genetic drift and shift, it is possible that new strains and  
296 serotypes of the virus could have emerged in the country. In order to avoid the danger of  
297 bluetongue disease outbreak in Nigeria, there is need for continuous national BTV surveillance  
298 and further studies to isolate and characterize BTV from Nigerian ruminant populations and  
299 *Culicoides* with the aim of identifying currently circulating serotypes and strains which can be  
300 used as potential vaccine candidates towards achieving effective control of the disease in  
301 Nigeria. Uncontrolled movement of animals from neighboring West African countries into  
302 Nigeria should be checked, and absolute quarantine measures enforced to ensure that animals  
303 coming into the country are free of the virus.

304

305

## 306 REFERENCES

- 307 1. Center for Food Security and Public Health at Iowa State University (CFSPH).  
308 Bluetongue: Sore Muzzle, Pseudo Foot-and-Mouth Disease, Muzzle Disease, Malarial  
309 Catarrhal Fever, Epizootic Catarrh, Beksiekte. Institute for International cooperation in  
310 Animal Biologics, College of Veterinary Medicine, Iowa State University, 2015.
- 311
- 312 2. Roy, P. "Functional mapping of bluetongue virus proteins and their interactions with host  
313 proteins during virus replication". *Cell Biochemistry and Biophysics* 2008, 50(3): 143-57.
- 314 3. Bjorn-Patrick Mohl and Polly Roy. Bluetongue Virus Capsid Assembly and Maturation.  
315 PMCID 6(8): 3250–3270: PMC4147694. Published online Aug 21 2014.  
316 doi: 10.3390/v6083250 PMID: 25196482.
- 317
- 318 4. Roy P. Bluetongue virus structure and assembly. PMID 2017Jun;24:115-123: 28609677  
319 DOI:10.1016/j.coviro.
- 320
- 321 5. MacLachlan, N.J and Guthrie, A.J. Re-emergence of bluetongue, African horse sickness,  
322 and other Orbivirus diseases. *Veterinary Research* Nov-Dec; 41(6): 35. Published online  
323 2010 Jan 27. doi: 10.1051/vetres/2010007.
- 324

- 325 6. Maan, S., Maan, N.S., Nomikou, K., Veronesi, E., Bachanet – Bankowska, K.,  
326 Belaganahali, M.N., Attoui, H. and Mertens, P.P. Complete genome characterization of a  
327 novel 26<sup>th</sup> bluetongue virus serotype from Kuwait. *PLoS one*. 2011; 6(10): e26147.
- 328 7. Mellor, P.S. and Boorman, J. The transmission and geographical spread of African horse  
329 sickness and bluetongue viruses. *Annual Tropical Medical Parasitology*, 1995; 89: 1-15.
- 330 8. Mellor, P.S., Boorman, J. and Baylis, M. *Culicoides* biting midges: Their role as arbovirus  
331 vectors. *Annual Review of Entomology*, 2000; 45: 307 – 340.
- 332 9. Purse, B.V., Mellor, P.S., Rogers, D.J., Samuel, A.R., Mertens, P.C. Baylis, M. Climate  
333 change and the recent emergence of bluetongue in Europe. *National Review of*  
334 *Microbiology*, 2005; 3: 171 – 181 ([doi:10.1038/nrmicro1090](https://doi.org/10.1038/nrmicro1090)) [[PubMed](#)] [[Google Scholar](#)].
- 335 10. MacLachlan, N.J. Bluetongue and epizootic haemorrhagic disease. In: US Animal Health  
336 Association, Committee on Foreign Animal Disease. Foreign animal diseases: the gray  
337 book. Ed 7. Part III, Chap 7. Richmond, V.A.: *US Animal Health Association*, 2008; 159 –  
338 66.
- 339 11. USAHA (US Animal Health Association). Committee on Bluetongue and Bovine  
340 Retrovirus. *Committee report*. Oct 21, 2002.
- 341 12. McKercher, D.G., McGowan, B. and Howarth, J.A. and Saito, J.K. A preliminary report  
342 on the isolation and identification of the bluetongue virus from sheep in  
343 California. *Journal of American Veterinary Medical Association*, 1953; 122: 300 – 301.
- 344 13. Singer R. S., MacLachlan N.J. and Carpenter T.E. Maximal predicted duration of  
345 viraemia in bluetongue virus infected cattle. *Journal of Veterinary Diagnostic*  
346 *Investigation*, 2001; 13: 43 -9.
- 347 14. Murphy F. A., Gibbs E. P., Horsinek M.C. and Studdert M.C. Reoviridae. In: *Veterinary*  
348 *Virology*. 1999; Ed 3. San Diego, C. A: Academic Press.
- 349 15. Office International Des Epizooties/World Organisation for Animal Health (OIE). Press  
350 release. Bluetongue detected for the first time in Northern Europe.  
351 <http://www.oie.int/eng/press/en-060823.htm>. Accessed 22 Nov. 2006.
- 352 16. Aeillo, S. ed. Bluetongue disease. In: *Merck Veterinary Manual*. Ed 8. Merck & Co., Inc.  
353 1998. Whitehouse Station, N.J., USA.

- 354 17. Verwoerd, D.W. and Erasmus, B.J. Bluetongue, In: Coetzer J.A.W. and Tstin, R.C.  
355 (Eds.). Infectious diseases of livestock, 2<sup>nd</sup> ed., *Oxford University Press Southern Africa*,  
356 *Cape Town*. 2004; pp. 1201 – 1220.
- 357 18. Schwartz – Cornill, I., Mertens, P.P. Contreras, V., Hemati, B., Pascale, F., Breard, E.,  
358 Mellor, P.S., MacLachlan, N.J. and Zientara, S. Bluetongue virus: virology, pathogenesis  
359 and immunity. *Veterinary Research*, 2008; 39: 46.
- 360 19. MacLachlan, N.J., Drew, C.P., Darpel, K.E. and Worwa, G. The pathology and  
361 pathogenesis of bluetongue. *Journal of Companion Pathology*, 2009; 141: 1-16.
- 362 20. McGrath, M. Dancing disease set for long run. 2008  
363 (<http://news.bbc.co.uk/1/hi/uk/7019511.STM>).
- 364 21. Taylor, W.P. The epidemiology of bluetongue. *OIE Science and Technology Reviewed*,  
365 1986; 5: 351 – 356.
- 366 22. Bida, S.A., Njoka, C. O. and Eid, F.I.A. Bluetongue in Wiltshire horn sheep. *Veterinary*  
367 *Record*, 1975; 97: 946.
- 368 23. Ekue, F.N., Nfi, A.N., Tsangue, P. Taylor, W.P. and Gumm, I. D. Bluetongue in exotic  
369 sheep in Cameroon. *Tropical Animal Healthand Production*, 1985; 17: 187 – 188.
- 370 24. Sudana, I.G. and Malole, M. Annual report on animal diseases investigation in Indonesia  
371 during 1976- 1981. Bogor, Java: Balai Penyidikan Penyakit Hewan, 1982.
- 372 25. Mellor, P.S. Bluetongue. *State Veterinary Journal*, 1994a; 4: 7-10.
- 373 26. Afshar, A., Eaton, B.T., Wright, P.F., Pearson, J.E., Anderson, J., Jeggo, M. and Trotter,  
374 H.C. Competitive ELISA for serodiagnosis of bluetongue: evaluation of group-specific  
375 monoclonal antibodies and expressed VP7 antigens. *Journalof Veterinary*  
376 *DiagnosticInvestigation*, 1992; 4:231 – 237.
- 377 27. Obi, T.U., Taylor, W.P. and Ojo, M.O. Prevalence of bluetongue virus precipitating  
378 antibodies in sheep and goats in Southern Nigeria. *Tropical Veterinarian*, 1983; Volume  
379 1 Number 4 page 205 – 208.
- 380 28. Oluwayelu, D.O., Olatoye, O., Akanbi, M. and Hoffmann. Ra-Emergence of Bluetongu  
381 Virus Infection in Oyo state, Nigeria. In: *Proceedings of 5<sup>th</sup> Pan Commonwealth*  
382 *Veterinary Conference, Accra, Ghana*, 2011; pp 234-238.
- 383 29. Thrusfield, M. (1995). *Veterinary Epidemiology*, Second edition. *Blackwell Science*  
384 *Publications, Oxford*, 1995; pp178-198.

- 385 30. Obot N.I., Emberga, T.T. and Ishola, K.S. 22 Years Characterized Trends of rainfall in  
386 Abeokuta, Nigeria. *Research Journal of Applied Sciences*, 2011; Volume 6 issue: 4 page  
387 264-271.
- 388 31. Omotoso, S. and Omotos, O. (1992). Iree Alalubosa. *An Iree Progressive Union*  
389 *Commissioned History*, Oshogbo: Nigeria.1992; Signs and Wonders Publishers Ltd.
- 390 32. Dipeolu, O.O. and Ogunrinade, A.F. Studies on *Culicoides* species of Nigeria. VII. The  
391 biology of some Nigerian *Culicoides* species. *Z. Parasitenkd.*, 1997; 51(3): 289-290.
- 392 33. Akinboade, O.A., Hassan, J.O., Adejinmi. Public health importance of market meat  
393 exposed to refuse flies and air-borne microorganisms. *International Journal of Zoonosis*,  
394 1984; 1: 111 – 114.
- 395 34. Taylor, W.P. and McCausland, A. Studies with bluetongue virus in Nigeria. *Tropical*  
396 *Animal Health Production*, 1976; 8: 167-173.
- 397 35. Moore, D.L. and Kemp, C.E. Bluetongue and related viruses in Ibadan, Nigeria.  
398 Serologic studies of domesticated and wild animals. *American Journal of Veterinary*  
399 *Research*, 1974; 35: 1115-1120.
- 400 36. Milree, J.M., Walton, C. and Jerome, C.P. Prevalence in sheep and goats in northern  
401 Nigeria of antibodies to bluetongue (type 7). *West African Research Team, Royal*  
402 *Veterinary College*. 1977; Final report, page. 39-44.
- 403 37. Durojaiye, O. Agar gel precipitation antibody to bluetongue virus in Nigerian cattle,  
404 sheep and goats. *Nigerian Veterinary Journal*, 1979; 8: 64-67.
- 405 38. Gibbs, E.P.J. and Greiner, E. C. Bluetongue and Epizootic Haemorrhagic Disease. In *The*  
406 *Arboviruses: Epidemiology and Ecology*, 1988; Vol II, TP Monath (edited), CRC Press,  
407 Boca Raton, pp. 39 – 70.
- 408 39. Cynthia, M., K.(2005). *Merck Veterinary Manual 9<sup>th</sup> edition*, Merck & Co., Inc. 2005,  
409 Whitehouse Station, U.S.A.
- 410 40. Sperlova A. and Zendulkova D. Bluetongue: a review. *Veterinarni Medicina*, 56, 2011  
411 (9): 430-452.
- 412 41. Pinto J., Bonacic, C., Hamilton-West, C., Romero, J. and Lubroth, J. Climate change and  
413 animal diseases in South America. *Review of Science and Technology*, 2008; 27: 599 –  
414 613.

- 415 42. Sellers, R.F., and Pedgley, D.E. Possible windborne spread to Western Turkey of  
416 bluetongue virus in 1977 and to Akabane virus in 1979. *Journal of Hygiene of*  
417 *Cambridge*1985; 95: 149 – 158.
- 418 43. Gibbs, E.P. and Greiner, E.C.The epidemiology of bluetongue. *Companion of*  
419 *Immunology of Microbial Infectious Diseases*, 1994; 17: 207 – 220.
- 420 44. Tomori O. Bluetongue and related viruses in Nigeria: Experimental infection of West  
421 African dwarf sheep with Nigeria strains of the viruses of epizootic haemorrhagic disease  
422 of deer and bluetongue. *Veterinary Microbiology*, Volume 5, Issue 3, September 1980,  
423 Pages 177 – 185.
- 424

UNDER PEER REVIEW