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Capture Myopathy in Little Bustards after Trapping and Marking

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ABSTRACT: Four little bustards (Tetrax tetrax) (one adult and three juvenile males), captured with leg nooses and fitted with a backpack radiotag, died after capture. The first bird was found after 16 days with its left foot caught in the harness and died after 1 day. The other birds showed symptoms of capture myopathy after release, such as the difficulty or inability to fly and/or walk. They died after 5, 6, and 8 days, respectively. At necropsy, muscles affected in all cases were those from the legs, and these were diffusely pale and dull, with a soft friable texture. Microscopically these muscles had multiple foci of myofiber fragmentation, loss of striation, and necrosis; a mononuclear cell infiltrate was observed in muscle from two birds. These findings suggest the little bustard is susceptible to capture myopathy and that caution should be exercised during its capture and handling.

Key words: Birds, capture myopathy, leg nooses, little bustard, radiotagging, Tetrax tetrax.

Capture myopathy, also known as exertional myopathy or exertional rhabdomyolysis, is a disease seen in animals during trapping, pursuit, capture, restraint, and transport. It is frequently reported in mammals, mainly in ungulates, but it has also been observed in birds (Williams and Thorne, 1996). Long-legged birds, like flamingoes, ostriches, and cranes, have been reported frequently to suffer capture myopathy (Bollinger et al., 1989). Additionally, this condition has been described in different species of waterfowl, gallinaceous birds, and raptors (Spraker et al., 1987; Domingo et al., 1991; Williams and Thorne, 1996; Höfle et al., 2004). Capture myopathy is characterized by damage to muscle tissues brought about by complex physiological changes that depend upon species and the circumstances of capture, physical exertion, and stress. However, hyperthermia and metabolic acidosis due to elevated levels of lactic acid from anaerobic glycolysis, as a response to intense muscular activity, are the central factors (Williams and Thorne, 1996).

The little bustard (Tetrax tetrax) is a near-threatened bird, about the size of a female pheasant, belonging to the Order Gruiformes, which breeds in open steppe and steppe-like landscapes. In the course of a radio-tracking study carried out in the Lleida plains (41°33′34″N, 0°1′57″–01°E) (Catalonia, NE Spain) in 2002 and 2004, four cases of capture myopathy (one adult and three juvenile males) after their capture, radiotagging, and release were reported. The birds were captured by leg nooses and fitted with backpack tags model TW3 (Biotrack, Dorset, UK). They were part of a larger group of 37 little bustards collected under the authority of permits from the Departament de Medi Ambient of the Catalan government. The length of time that the birds were caught in the leg nooses before they were immobilized was less than 2 min. Measuring and marking were performed at the place of capture, and the total time of restraint was between 25 and 55 min. They were also released at the place of capture. Two little bustards were frozen, and two were refrigerated prior to post-mortem and histopathologic examinations. Tissues examined histologically included lung, heart, liver, spleen, kidney, and skeletal muscle from wings, pectoral, and femoral regions. They were fixed in 10% formalin, embedded in paraffin, sectioned at 5 µm, mounted on glass slides, and stained with hematoxylin and eosin.

All of the trapped birds appeared to be
healthy at the time of capture. The first little bustard was found after 16 days with its left foot caught in the harness of the radiotag. It couldn’t stand and was struggling to escape (flapping its wings) when discovered. The animal died 1 day after it arrived at the Wildlife Rehabilitation Centre of Vallcalent (Lleida). The other three little bustards showed difficulty or inability to fly and/or walk. In two birds, parts of the muscles from the right pectoral region, wing and leg, right lung, heart, gut, and spleen were removed by a bird of prey. However, it is not known if the birds were alive or not when they were predated.

At necropsy, the four little bustards had poor body condition. Muscles of the femoral region of both legs were diffusely pale, dull, and friable. In one bird, the iliotibialis cranialis of the left leg was also pale with streaked areas running parallel to the muscle bundle. Histopathologic examination of affected muscles from the legs showed extensive areas of muscular degeneration and necrosis. Myofiber changes included swelling, increased eosinophilia, multiple foci of fragmentation, loss of striation, and pyknosis of myofiber nuclei (Williams and Thorne, 1996). Although muscular hemorrhages and myocardial lesions have been frequently reported (Spraker et al., 1987; Tully et al., 1996), not all animals develop these lesions (Williams and Thorne, 1996). These lesions may be related to the severity of the disease and the time elapsed from muscular damage to death, which is influenced by the method of capture and other factors previously discussed. Increased cellularity was observed in two little bustards, but it was predominantly a mononuclear infiltrate, rather than the neutrophilic/heterophilic infiltrate previously described for capture myopathy in mammals and birds (Williams and Thorne, 1996).

Renal lesions due to ischemia and myoglobin deposits are usually most pronounced after one week postexertion, but are not always present (Carpenter et al., 1991; Williams and Thorne, 1996) and were not observed in the little bustards.

Macroscopic and microscopic lesions were found only in the musculature of the legs of the affected little bustards, like

(Alectoris rufa) (Höfle et al., 2004), but we did not observe hemorrhages in the affected muscles. Extensive areas of hemorrhage were also observed in lesser flamingoes (Phoenicopterus minor) and wild turkeys (Meleagris gallopavo) after capture (Young, 1967; Spraker et al., 1987). Rocket nets used to capture wild turkeys and chasing and capturing flamingoes may have been more stressful for the animals compared to the leg nooses we used. However, other risk factors must be considered, such as species, sex, age, ambient temperature, pursuit time, restraint time, and transport duration (Williams and Thorne, 1996; Nicholson et al., 2000).

The histologic lesions of affected muscles of the birds in our study were mostly those previously reported for acute capture myopathy cases, such as swelling, multiple foci of fragmentation, loss of striation, and pyknosis of myofiber nuclei (Williams and Thorne, 1996). Although muscular hemorrhages and myocardial lesions have been frequently reported (Spraker et al., 1987; Tully et al., 1996), not all animals develop these lesions (Williams and Thorne, 1996). These lesions may be related to the severity of the disease and the time elapsed from muscular damage to death, which is influenced by the method of capture and other factors previously discussed. Increased cellularity was observed in two little bustards, but it was predominantly a mononuclear infiltrate, rather than the neutrophilic/heterophilic infiltrate previously described for capture myopathy in mammals and birds (Williams and Thorne, 1996). Renal lesions due to ischemia and myoglobin deposits are usually most pronounced after one week postexertion, but are not always present (Carpenter et al., 1991; Williams and Thorne, 1996) and were not observed in the little bustards.

Macroscopic and microscopic lesions were found only in the musculature of the legs of the affected little bustards, like
in the sandhill crane and red-legged partridge (Carpenter et al., 1991; Höfle et al., 2004). Since all striated muscles can be affected, the distribution of lesions in the little bustards may reflect the nature of the exertion that induced the changes (Williams and Thorne, 1996). As legs have to be restrained in a flexed position, when the bustards struggle to escape, the leg muscles suffer intense physical exertion. The constant isotonic state may contribute to hindering blood flow and aggravating muscle hypoxia. It has been suggested that long-legged birds may be more susceptible to capture myopathy than are other birds (Carpenter et al., 1991). The little bustard can be considered a long-legged bird, since it belongs to the Order Gruiiformes. To our knowledge, this is the first time that capture myopathy is described in the little bustard. However, it has been previously reported in other species of bustards, such as the houbara bustard (Chlamydotis undulata) and kori bustard (Ardeotis kori) (Bailey et al., 1996). These findings suggest that the little bustard is also susceptible to capture myopathy and that the family of bustards should be considered as a risk group. Furthermore, it can increase the probability of predation (Spraker et al., 1987), and this was probably the case in two of the little bustards studied here. Since treatment is extremely difficult, caution should be exercised during capture and handling, and every effort should be made to prevent the problem.

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**LITERATURE CITED**


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