

1 Bull BSEs: Evaluation for Success

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5
6 Abstract

7 Bull breeding soundness exams are among the most critical responsibilities we have as
8 bovine veterinarians. These exams are designed to identify subfertile and infertile bulls, which, if
9 used for breeding, can lead to significant financial losses for the producer. Such losses arise from
10 a decrease in the number of calves born, an extended calving season, and ultimately, fewer
11 pounds of calf at weaning.

12
13 Introduction

14 Bull breeding soundness exams are complex because they involve evaluating multiple
15 factors to determine a bull's potential breeding status, including a physical exam, scrotal
16 circumference, sperm motility, and morphology. In this talk, we will explore some of the most
17 challenging aspects of the exam and discuss strategies for achieving success.

18
19 Physical Exam and Scrotal Circumference

20 Physical exams are a crucial aspect of bull breeding soundness exams because they
21 provide a comprehensive assessment of the bull's overall health and structural soundness, which
22 are essential for successful breeding. During the physical exam, veterinarians evaluate the bull's
23 musculoskeletal system, ensuring that the animal can physically mount and breed without

24 limitations such as lameness, joint issues, or muscular deficiencies. Additionally, examination of
25 the eyes, teeth, and body condition score is critical, as visual acuity and the ability to effectively
26 graze are vital for the bull's stamina and longevity in a breeding program. The physical exam
27 also includes an evaluation of the external genitalia, detecting any abnormalities, lesions, or
28 infections that could impair breeding performance or lead to the spread of disease.

29
30 Scrotal circumference in bulls is a critical component of the breeding soundness
31 evaluation, as it directly correlates with a bull's reproductive potential. Scrotal circumference
32 serves as an indicator of testicular mass, which in turn is associated with sperm production
33 capacity. Larger scrotal circumferences generally suggest higher sperm output, better semen
34 quality, and earlier onset of puberty in both the bull and his offspring. Moreover, bulls with
35 adequate scrotal circumference are more likely to produce calves that reach sexual maturity at an
36 earlier age, enhancing the reproductive efficiency of the herd. This measurement also provides
37 insight into the bull's overall endocrine function, as testicular size is linked to the production of
38 testosterone, which influences libido and mating behavior. Therefore, scrotal circumference is a
39 valuable predictor of a bull's fertility and is essential in selecting sires that will contribute to the
40 genetic and productive success of the herd.

41

42 Slide Preparation

43 Slide preparation is essential in bull breeding soundness exams because it significantly
44 impacts the accuracy of evaluating sperm morphology and motility. Proper slide preparation
45 ensures that sperm cells are evenly distributed, well-preserved, and free from artifacts that could
46 distort the assessment. Any errors in slide preparation, such as uneven smearing or inadequate

47 staining, can lead to misinterpretation of sperm quality, potentially resulting in the incorrect
48 classification of a bull's fertility status. Therefore, meticulous slide preparation is vital to provide
49 reliable data for determining a bull's reproductive viability.

50

51 Select Morphological Defects

52 *Proximal cytoplasmic droplet*

53 Proximal droplets are spherical condensations of cytoplasm 2-3 μm in diameter that surround the
54 neck and proximal midpiece of sperm. A high percentage of sperm with proximal cytoplasmic
55 droplets in an ejaculate is associated with abnormal epididymal function and sperm maturation,
56 or abnormal spermiogenesis. Peripubertal bulls often have a high percentage of sperm with
57 proximal droplets in the ejaculate. As bulls mature, the number of proximal droplets in the
58 spermogram should decrease.

59 *Round Cells*

60 Round cells can be noted during evaluation of semen motility slides especially in young
61 bulls. The evaluator must differentiate between immature sperm cells and white blood cells to
62 allow for accurate diagnosis. Immature sperm cells are quite variable in size, depending on
63 whether the cell is a primary or a secondary spermatocyte or a spermatid.^{11,16} Immature sperm
64 cells must be differentiated from white blood cells in semen. This differentiation can be
65 accomplished by staining a dried semen smear in Diff-quick[®], new methylene blue, or Wright's
66 giemsa. Once the stain is dried the round cells can be evaluated and a final diagnosis of immature
67 sperm cell or white blood cell can be made by the evaluator. If a diagnosis of immaturity is made
68 the bull should be reevaluated in 4-6 weeks to allow for maturation.

69 *Detached heads*

70 Detached but otherwise normal heads are likely due to senescence of normal sperm
71 during storage in the epididymis or ductus deferens, whereas detached abnormal heads may be
72 due to abnormal formation of the basal plate and/or implantation fossa. In normal bulls,
73 peristalsis continually moves sperm from the cauda epididymis into the urethra, ensuring a
74 reserve of fresh sperm for ejaculation.⁹² Failure of this transport mechanism is associated with
75 sperm accumulation and eventual senescence of stored sperm. These bulls are referred to as
76 “sperm accumulators” or “rusty load bulls”. Sperm accumulation in bulls is not associated with
77 age and a predilection to the occurrence accumulation appears to be permanent. Re-occurrence
78 within 1 month after sexual rest is common following resolution of sperm accumulation by
79 frequent electroejaculation. Regardless of cause detached heads are not normal and should be
80 counted as abnormal cells.

81 *Iatrogenic Changes*

82 Iatrogenic changes noted in the spermogram are mostly associated with slide
83 preparation. The most common change noted are those due to hypo-osmotic changes whether
84 that comes from stain, fixing solutions, prolonged drying times, cold slides, or cold shock of
85 ejaculate prior to staining. Hypo-osmotic changes are of high suspicion when there is a high
86 percentage of bent midpieces. Characteristically these midpieces have no retained droplet within
87 the bend which aids in differentiating this iatrogenic defect from DMRs (distal midpiece
88 reflexes). Cold shock may also be noted during evaluation of progressive motility as will be
89 depicted by sperm moving slowly, backwards and circling and in severe cases shimmering in
90 place.

91 Evaluation of semen slides

92 Differential counts of sperm morphology must be performed using 1000X magnification
93 (oil immersion) as lower magnifications may result in failure to recognize some defects. Semen
94 should always be evaluated at 100x oil immersion because this magnification level allows for a
95 detailed examination of sperm morphology, providing the resolution necessary to identify subtle
96 abnormalities in sperm structure that may not be visible at lower magnifications. At 100x oil
97 immersion, individual sperm cells can be closely inspected for defects in the head, midpiece, and
98 tail, which are critical indicators of fertility. This high level of detail is essential for accurately
99 assessing the proportion of normal versus abnormal sperm, aiding in the overall evaluation of the
100 bull's reproductive potential. Additionally, the clarity provided by oil immersion ensures that the
101 observations are precise, reducing the likelihood of diagnostic errors that could affect breeding
102 decisions.

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