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## Standardization of a method to detect bovine sperm-bound anti-sperm antibodies by flow cytometry

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1	Standardization of a method to detect bovine sperm-bound anti-sperm antibodies by flow
2	cytometry
3	
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15	
16	Abstract
17	
18	The objectives of this study were to standardize some methodological and analytical
19	aspects of a direct technique to detect sperm-bound anti-sperm antibodies (ASAs) in bovine
20	semen using flow cytometry. Four ASA-positive bulls with experimentally induced ASAs and 10
21	reproductively normal ASA-negative bulls were included in the study. The effect of pre-fixation
22	of sperm membranes with formalin buffer solution and inclusion of dead cells in the analysis was
23	evaluated. Fixation of sperm membranes had no significant effect on the percentage of IgG- or

24	IgA-bound spermatozoa detected by flow cytometry. Including dead cells in the analysis
25	increased the percentage of IgG-bound spermatozoa in fixed (live and dead $18.6 \pm 9.7$ % and live
26	$1.3 \pm 0.5$ %) and non-fixed samples (live and dead $18.8 \pm 9.2$ %, live $1.5 \pm 0.6$ %) (P = 0.0029),
27	as well as IgA-bound spermatozoa in fixed (live and dead $16.3 \pm 6.4$ %, live $0.3 \pm 0.5$ %) and
28	non-fixed samples (live and dead 21.4 $\pm$ 4.6 %, live 1.0 $\pm$ 0.5 %) (P = 0.0041) (median $\pm$ SE) in
29	semen from ASA-negative bulls. Intra-sample, intra-assay and inter-assay coefficients of
30	variation (CV) for determination of sperm-bound IgG were 0.8 %, 4.6 % and 5.3 %, respectively.
31	For determination of sperm-bound IgA, intra-sample, intra-assay and inter-assay CV were 2.8 %,
32	8.4 % and 40.3 %, respectively. In spite of the high inter-assay CV for IgA determination, all
33	ASA-positive bulls had high percentages of IgA-bound spermatozoa at all times. Flow cytometry
34	correctly identified ASA-positive bulls. Confocal laser microscopy confirmed the binding of
35	ASAs to the sperm head and cytoplasmic droplets, and less frequently to the mid and principal
36	piece. It was concluded that fixation was not necessary. Dead cells should be excluded from the
37	analysis since ejaculates with large numbers of dead cells can yield false-positive results. Flow
38	cytometry was accurate and reliable for detection of sperm-bound IgG and IgA and
39	discrimination between ASA-positive and ASA-negative bulls.
40	
41	Keywords: Flow cytometry; anti-sperm antibodies; sperm-bound antibodies; immunoinfertility;

42 bovine

43

44 1. Introduction

46 During spermatogenesis, developing germ cells express new surface antigens that are not recognized as self. Sperm-specific surface antigens first appear on pachytene primary 47 spermatocytes [1]. The blood-testis barrier (BTB), removal of antigenic apoptotic cells by 48 phagocytosis and immunosuppressive factors released by Sertoli cells confer the testes an 49 immune privileged status. Disruption of the BTB induced by infectious, inflammatory or 50 degenerative conditions exposes sperm antigens to the immune system and results in formation 51 of anti-sperm antibodies (ASAs) [2]. In bulls, genital infection with Chlamydia sp., Brucella 52 abortus and Infectious Bovine Rhinotracheitis Virus was associated with concomitant presence 53 54 of ASAs [3,4]. Antisperm antibodies were also detected in bulls with seminal vesiculitis [5] and orchitis [6]. The ASAs persisted in a bull with orchitis for 18 m after initial presentation [6]. 55 Persistence of ASAs can account, at least in part, for the long-term effects of genital infection on 56 fertility. Exposure to electromagnetic pulses was also shown to alter the BTB and result in 57 formation of ASAs in mice [7]. Exposure to electromagnetic pulses from electric transmission 58 lines, generators and fences could represent an unidentified risk factor for immune-mediated 59 infertility in bulls. 60

Bovine ASAs can reduce penetration and fertilization of oocytes in vivo and in vitro, 61 sperm-zona pellucida secondary binding, the ability of capacitated spermatozoa to complete the 62 acrosome reaction and the motility of capacitated and non-capacitated spermatozoa [8-10]. 63 Antisperm antibodies can impair fertility by contributing one more factor to an already 64 65 compromised semen sample, or by being the primary cause of idiopathic infertility. Their effect on fertility depends on the location of the ASAs, their regional specificity, the antibody class, 66 isotype and load, and the antigen specificity [11-13]. Antibodies directed against sperm antigens 67 68 can be detected free in seminal plasma or serum. However, only those bound to the surface of

spermatozoa are of significance for fertility [13]. Both IgA and IgG, but not IgM, have a proven
negative effect on fertility [11,14]. Therefore, an ideal diagnostic test should be able to identify
sperm-bound ASAs and provide information about the proportion of ASA-bound spermatozoa in
an ejaculate, the antibody class and load, and the regional specificity [15].

To date, most reports in veterinary medicine have involved the use of indirect techniques 73 74 to detect ASAs in serum or seminal plasma. Sperm agglutination [16,17] and immobilization tests [18] have been used to detect ASAs in bulls. However, these tests are insensitive and 75 nonspecific [15]. Immunofluorescence [17], immunocytochemistry [3,19] and enzyme-linked 76 77 immunosorbent assay [8,20] have also been used in bulls. These techniques require fixation of the cell membranes. Fixation can result in non-specific binding of antibodies, exposure of 78 intracellular antigens, denaturation of sperm antigens or membrane damage, resulting in false-79 positive or false-negative results [15,21]. The mixed antiglobulin reaction and immunobead-80 binding tests are most commonly used in human medicine [22]. These tests provide a semi-81 quantitative estimation of the proportion of ASA-positive spermatozoa, and information on the 82 antibody class and its location on the spermatozoa. However, both tests are based on counting 83 motile spermatozoa bound to beads or latex particles. Therefore, the estimation is subjective. The 84 85 tests require good sperm motility in the samples from the infertile patients if a direct test is used, or availability of a semen donor with good sperm motility if the indirect test is used [15]. Instead, 86 flow cytometry allows objective and quantitative estimation of ASAs on the surface of living 87 88 spermatozoa and is a sensitive, specific and repeatable test [15]. Flow cytometry also allows identification of the antibody class, isotype and load [15]. 89

90 The use of flow cytometry to detect ASAs in bulls was only recently reported [6].
91 Moreover, a standardized direct technique to detect sperm-bound ASAs has not been developed

in veterinary medicine. How the samples are processed and analyzed can have a significant 92 impact on the reliability of the results. When analyzing live cells, cross-linking of surface 93 antigens by multivalent antibodies, or of antigen-antibody (ag-ab) complexes by secondary 94 antibodies can cause aggregation of ag-ab complexes into patches and caps [21,23]. Patching 95 and capping is followed by shedding of the ag-ab complexes. Patching and capping can be 96 97 prevented by fixing the cell membranes prior to incubation with antibodies [21,23]. However, as mentioned before, fixation can alter the membranes or antigens also giving misleading 98 results [15,21]. Another source of error is nonspecific uptake of antibody by dead spermatozoa. 99 100 Nonspecific binding can yield false-positive results if the proportion of dead cells in the ejaculate is high [15]. 101 The objectives of this study were to standardize some methodological and analytical 102 aspects of a direct technique to detect sperm-bound ASAs in bovine semen using flow 103 cytometry. The effect of fixation and inclusion of dead cells in the analysis were evaluated, 104 coefficients of variation for the standardized protocol were calculated and binding of ASAs to 105 bovine spermatozoa was confirmed with confocal laser scanning microscopy. 106 107 108 2. Materials and Methods 109

110 2.1. Animals

111

Four 1-year old *Bos Taurus* bulls of Angus breed were purchased from local producers. The bulls were housed individually or in pairs in pens, and fed brome hay and water *ad libitum*, and 2 lb of sweet feed twice daily. Bulls were allowed to acclimate for one week prior to starting the

115	experiments. To provide a known ASA-positive control, the bulls were immunized with
116	autologous spermatozoa as described below. When the percentage of ASA-bound spermatozoa
117	was $\geq$ 20 %, bulls were considered to have a positive response [24] and experiments were
118	initiated. Additionally, ten privately owned Angus bulls (standardization of the technique, $n = 5$ ;
119	calculation of coefficients of variation, $n = 5$ ) classified as satisfactory breeders during routine
120	breeding soundness examination [25] were included as ASA-negative control bulls. Bulls were
121	considered satisfactory breeders if they had no gross abnormalities of their internal and external
122	genitalia, a scrotal circumference above the minimum recommended value for the age, $\geq$ 30 %
123	individual sperm motility and $\geq$ 70 % morphologically normal spermatozoa [25]. The study was
124	performed following Kansas State University's Institutional Animal Care and Use Committee's
125	guidelines. The bulls with experimentally-induced antibodies were euthanized at the end of the
126	study.

128 2.2. Semen collection and evaluation

129

130 Semen was collected using electroejaculation (SireMaster Original, ICE Corporation,

Manhattan, KS, USA). The accessory sex glands were massaged transrectally with a gloved hand
for 30 to 60 sec. A 6.5-cm in diameter lubricated rectal probe was inserted into the rectum with
the electrodes facing ventrally. Electrical stimulation was applied with increasing intensity until
ejaculation [25]. A complete semen evaluation [25] was performed immediately after collection.
Semen was then used for immunization or for the experiments.

136

137 2.3. Immunization of bulls

139	Immunizations were performed as described before with some modifications [18]. Ejaculated
140	spermatozoa were washed three times by centrifugation at 900 x g for 10 min diluted in warm
141	Dulbecco's phosphate buffered saline (DPBS, Invitrogen, Grand Island, NY, USA). Washed
142	spermatozoa, 1 x 10 <sup>9</sup> , were re-suspended to 1 mL in DPBS. One milliliter of Freund's complete
143	adjuvant (Sigma-Aldrich, St. Louis, MO, USA) was then added. Each bull was immunized with
144	2 mL of inoculum containing 1 x $10^9$ autologous spermatozoa. The inoculum was administered
145	intramuscularly in the neck in four different aliquots of 0.5 mL each. Booster immunizations
146	were administered to three bulls 22 d after the primary immunization. Semen was processed in the
147	same way as for primary immunizations but Freund's Incomplete Adjuvant (Sigma-Aldrich) was
148	used instead of Freund's Complete Adjuvant. One bull did not receive a booster immunization
149	since the response to the primary immunization was satisfactory.
150	
151	2.4. Standardization of flow cytometry for detection of ASAs
152	
153	The effect of fixing spermatozoa with formalin buffer solution prior to labeling on the ability
154	to detect sperm-bound ASAs was evaluated. One ejaculate was collected from each bull with
155	experimentally-induced ASAs $(n = 4)$ and each ASA-negative bull $(n = 5)$ . Each ejaculate was
156	initially divided into two aliquots. Semen was diluted to 50 x $10^6$ spermatozoa /mL in DPBS
157	(non-fixed samples) or formalin buffer solution (FBS, Animal Reproduction Systems, Chino,
158	CA, USA) (fixed samples). Formalin buffer solution had been previously diluted 1:10 in DPBS.
159	After 10 min at room temperature, samples were washed three times by centrifugation and
160	labeled with fluorescein isothiocyanate (FITC)-labeled anti-bovine IgG or IgA, or their

161	respective isotype control antibodies as described below. Samples were analyzed by flow
162	cytometry. The percentage of IgG- and IgA-bound spermatozoa was calculated including the
163	entire cell population (live and dead cells) or live cells only. Comparisons were made among
164	treatment groups: non-fixed samples including live cells only in the analysis, non-fixed samples
165	including both live and dead cells in the analysis, fixed samples including live cells only in the
166	analysis, and fixed samples including live and dead cells in the analysis.
167	
168	2.5. Calculation of coefficients of variation
169	
170	It was determined in the previous experiment that fixation was not necessary and that
171	including dead cells in the analysis yielded false-positive results. Therefore, non-fixed samples
172	were used and only live cells were included in the analysis for calculation of coefficients of
173	variation (CV). One ejaculate from each ASA-negative $(n = 5)$ and ASA-positive bull $(n = 4)$
174	was divided into five aliquots and processed in five replicates to calculate intra-assay CV. One of
175	the aliquots was evaluated five times to assess intra-sample CV. Only semen from ASA-positive
176	bulls was available for assessment of the inter-assay CV. Inter-assay CV was calculated
177	retrospectively from two ejaculates collected from each bull 6 to 20 d apart. The CVs were
178	calculated with the following formula: CV (%) = mean of standard deviations / mean x 100.
179	
180	2.6. Antibody labeling
181	
182	Semen was diluted to a concentration of 50 x $10^6$ spermatozoa /mL in warm DPBS, and was
183	washed three times by centrifugation at 900 x g for 10 min in DPBS. Then, $2.5 \times 10^6$ of washed

184	spermatozoa were added to each of four tubes containing 320 $\mu L$ of DPBS. The corresponding
185	antibodies were added to each tube: $IgG = 30 \ \mu L$ of FITC-labeled polyclonal goat anti-bovine
186	IgG F(ab') <sub>2</sub> (12.5 µg/mL; Cat. No. 101-096-003, Jackson Immunoreseach Laboratories Inc.,
187	West Grove, PA, USA); IgG isotype control = 30 $\mu$ L of FITC-labeled polyclonal rabbit anti-goat
188	IgG F(ab') <sub>2</sub> (12.5 µg/mL; Cat. No. 305-096-003; Jackson Immunoreseach Laboratories Inc.);
189	IgA = 20 $\mu$ L of FITC-labeled polyclonal rabbit anti-bovine IgA (12.5 $\mu$ g/mL; Cat. No. A10-
190	108F; Bethyl Laboratories, Montgomery,TX, USA); or IgA isotype control = 20 $\mu$ L of FITC-
191	labeled polyclonal goat anti-mouse IgA (12.5 µg/mL; Cat. No. A90-103F; Bethyl laboratories).
192	A preliminary study was performed to evaluate saturating concentrations and select the
193	appropriate concentration of each antibody (data not shown). The samples were incubated for 30
194	min at room temperature in the dark, followed by three washes by centrifugation at 900 x g for
195	10 min in DPBS. Propidium iodide (PI, viability stain), 5 $\mu$ L, was then added for simultaneous
196	staining of dead cells.

197

## 198 2.7. Flow Cytometry

199

The percentage of IgG- and IgA-bound spermatozoa was assessed by flow cytometry 200 201 (FACSCalibur, Becton Dickinson, San Jose CA, USA). From each sample, 10 000 cells were analyzed at a rate of 1 to  $2 \times 10^3$  cells /sec using DPBS as the sheath fluid. Data from these cells 202 were collected using forward scatter as the size parameter. A gate containing spermatozoa was 203 selected based on dot plot distribution of forward (size) versus side scatter (complexity 204 205 parameter) to eliminate debris and epithelial cells from the analysis (Fig. 1). The FITC and PI signals were detected using a standard argon laser (488 nm) and emission filters ( $535 \pm 30$  nm 206

207 for FITC and  $585 \pm 30$  nm for PI). The instrument was calibrated daily with standard beads so that the CV of the forward scatter and fluorescence channels were < 5 % on a daily basis. 208 Compensation for FITC emission into the PI detector or vice versa was done by establishing 209 quadrants on spermatozoa labeled only with PI or FITC-conjugated antibodies, followed by 210 electronic substraction of the FITC emission into the PI detector and PI emission into the FITC 211 detector. After color compensation, fluorescence emission data were collected with logarithmic 212 amplification for green fluorescence (FITC using FL1 detector) and orange-red fluorescence (PI 213 using FL2 detector). Quadrant settings were adjusted for each sample. The control quadrant 214 215 (lower left, LL) was marked on samples labeled with the isotype control to include < 1 % of cells as positive in the upper left (UL), upper right (UR) and lower right (LR) quadrants (Fig. 1). The 216 ASA-negative dead cells (PI stained) appeared in the UL quadrant, ASA-negative live cells (no 217 stain) in the LL quadrant, ASA-positive dead cells (dual stained) in the UR quadrant, ASA-218 positive live cells (FITC stained) in the LR quadrant (Fig. 1). The percentage of ASA-positive 219 live spermatozoa (LR quadrant) was calculated considering only live cells (PI negative cells in 220 LL and LR quadrants) in the analysis. When including dead cells (PI positive cells), the 221 percentage of ASA-positive spermatozoa (LR and UR quadrants) was calculated considering all 222 223 quadrants.

224

225 2.8. Confocal laser scanning microscopy

226

Labeled spermatozoa from bulls with experimentally-induced antibodies were evaluated under confocal laser scanning microscopy to confirm binding of the antibodies to the sperm surface. No attempts were made to quantitatively evaluate the percentage of ASA-bound

spermatozoa or the relative distribution of the binding sites. Spermatozoa were labeled with
FITC-labeled anti-bovine IgG or IgA as described above. After labeling, 10 µL of FBS was
added to inhibit sperm motility and facilitate visual evaluation. A drop of sperm suspension was
evaluated on a microscope slide under a cover slide. The FITC signal was excited at 488 nm and
was collected with a band pass filter at a wavelength of 505-550 nm. Samples were assessed at
X20 and X40 and optical sections were collected (LSM 710 META, Carl Zeiss MicroImaging,
Thornwood, NY).

237

238 2.9. Statistical analysis

239

Statistical analysis was performed using SAS package (SAS Institute, Cary, NC, USA). 240 Distribution of the data was tested for normality using a Shapiro Wilk test. Data were not 241 normally distributed. To determine response to immunization, percentages of IgG- and IgA-242 bound spermatozoa before and after the last immunization were compared using a Wilxocon 243 signed test. Only non-fixed live spermatozoa were included in this analysis. To assess the effect 244 of fixation and inclusion of dead cells in the analysis, differences in median percentages of IgG-245 246 and IgA-bound spermatozoa among treatment groups were compared using a Friedman test. The Friedman test is a non-parametric test that compares median values across treatments controlling 247 for bull. Since non-parametric tests were used, data were reported as median  $\pm$  SE. Differences 248 249 were considered significant at P < 0.05.

250

251 3. Results

252

Immunization with autologous spermatozoa induced a significant increase in the

254 percentage of both IgG-bound spermatozoa and IgA-bound spermatozoa. The percentage of IgG-

bound spermatozoa was  $2.9 \pm 2.1$  % and  $89.8 \pm 4.6$  % before and after immunization,

respectively (P = 0.0209). The percentage of IgA-bound spermatozoa was  $7.7 \pm 2.2$  % and  $75.7 \pm$ 

18.9 % before and after immunization, respectively (P = 0.0433) (median ± SE).

There was no significant difference in the percentage of IgG- or IgA-bound spermatozoa between samples fixed with FBS and non-fixed samples (Fig. 2). Including dead cells in the analysis increased the percentage of IgG- (P = 0.0029) and IgA-bound spermatozoa (P = 0.0041) detected in semen samples from ASA-negative bulls (Fig. 2). However, median percentages of ASA-bound spermatozoa did not differ among semen samples from ASA-positive bulls when dead cells were included in the analysis (Fig. 2).

Intra-sample CV for determination of sperm-bound IgG was 0.8 %, intra-assay CV was

4.6 % and inter-assay CV was 5.3 %. For determination of sperm-bound IgA, intra-sample CV

was 2.8 %, intra-assay CV was 8.4 % and inter-assay CV was 40.3 %. Both antibody classes

bound to the acrosomal, equatorial and post-acrosomal areas of the sperm head, and to

268 cytoplasmic droplets (Fig.3). Least frequently, ASAs bound to the sperm midpiece and principal269 piece.

270

271 4. Discussion

272

Systemic immunization with autologous spermatozoa induced an immune response in all
bulls characterized by an increase in sperm-bound IgG and IgA. Immunoglobulin G in genital
secretions is mostly derived from systemic circulation [26]. In the presence of an intact blood-

276 testis or blood-epididymis barrier, IgG reaches the genital tract and binds to spermatozoa at the rete testis or at ejaculation when spermatozoa contact the secretions of the accessory sex glands 277 [27-29]. On the other hand, IgA is produced locally [26]. Systemic immunization can result in 278 increased production of IgA within the genital tract, and increases in antigen-specific B cells in 279 the testis [29]. It is possible that systemic immunization of bulls with spermatozoa induced both 280 281 a systemic and mucosal immune response here. It is also possible that migration of activated IgA-committed B cells from lymph nodes draining the injection site to the genital mucosa 282 contributed to the increase in sperm-bound IgA after immunization, as described in humans [30]. 283 284 Recommendations for processing and evaluating bovine semen samples for detection of sperm-bound ASAs by flow cytometry can be made based on the results of this study. Polyclonal 285 antibodies and F(ab')<sub>2</sub> fragments were used here. Since they are expected to react with all 286 subclasses, use of polyclonal antibodies may decrease the likelihood of obtaining false-negative 287 results [15]. Use of F(ab')<sub>2</sub> fragments is also preferred to prevent non-immune binding of the Fc 288 portion of the IgG molecule to the sperm membrane [15], which occurs via disulfide 289 rearrangement at the cell surface in bulls [31]. Fixation of sperm membranes with formalin 290 buffer solution prior to labeling did not affect the ability to detect sperm-bound ASAs. Fixation 291 292 was performed to potentially prevent patching or capping of ag-ab complexes, which would have yielded false-negative results. Mature spermatozoa have both mobile and non-mobile surface 293 antigens [32,33]. Patching and capping involve redistribution of mobile antigens in response to 294 295 multivalent ligands. Patching is a local clustering of molecules, while capping is the aggregation of the clusters to a single area of the membrane. Following capping, molecules are shed from the 296 cell membrane [23]. While these phenomena were demonstrated in early spermatogenic cells [1], 297 298 patching and capping were not observed in late spermatids [1] or mature spermatozoa [34]. It

was speculated that non-mobile antigens are inserted later in germ cell development, and that cross-linking between mobile and non-mobile antigens results in loss of capping in late spermatids [1] and spermatozoa. While fixation may be necessary to prevent lateral mobility of antigens, it can also alter the results by causing non-specific binding of antibodies, exposure of intracellular antigens, denaturation of sperm antigens and membrane damage [15,21]. It was concluded that since fixation of sperm membranes prior to labeling did not affect the results but increased processing times, this procedure could be avoided.

When dead cells were included in the analysis, false-positive results were obtained in 306 307 samples from ASA-negative bulls. It is likely that non-specific binding of antibodies to dead cells or increased autofluorescence displayed by dead cells accounted for the increase in the 308 percentage of fluorescently-labeled spermatozoa among ASA-negative bulls [15,35]. In ASA-309 positive bulls, the percentage of ASA-bound spermatozoa was already high. Even when non-310 specific binding to dead cells may have occurred, the difference may not have been large enough 311 to be significant. It was concluded that dead cells should be excluded from the analysis to 312 prevent false-positive results in ASA-negative bulls. This limits the use of flow cytometry to 313 detect ASAs in bulls with necrozoospermia. 314

Coefficients of variation were all < 10 %, except for inter-assay CV for IgA-bound spermatozoa. It is not known if this high CV resulted from the low number of samples available, or from different frequencies of ejaculation that resulted in varying storage times and contact with ASA-loaded genital secretions. It is also possible that the variation reflected changes in antibody titers at different times post-immunization and was inherent to the model used rather than the test itself. The reason for the high inter-assay CV of the IgA test requires further investigation with more standardized sampling times. Nonetheless, the percentage of IgA-bound

spermatozoa was  $\geq 20$  % in all samples from all ASA-positive bulls. In spite of the high interassay CV, the test was able to correctly identify IgA-positive bulls. With this exception, CVs in this study were similar to those reported in the human literature [15]. It was concluded that flow cytometry was accurate and reliable for detection of sperm-bound ASAs and discrimination between ASA-positive and ASA-negative bulls.

Confocal laser microscopy confirmed binding of ASAs to the sperm surface. The 327 combination of flow cytometry and fluorescence microscopy provided an ideal diagnostic 328 approach. Flow cytometry allowed identification of sperm-bound ASAs and provided objective 329 330 and quantitative information about the antibody class and load. Additional use of fluorescence microscopy provided information about the regional specificity of the ASAs. Due the lack of 331 reports on presence and behavior of naturally-occurring sperm-bound antibodies in bulls, it is 332 difficult to determine how detection of experimentally-induced antibodies compares with 333 detection of sperm-bound ASAs produced during bacterial infection or trauma. Studies are under 334 way to determine the reference ranges and prevalence of naturally-occurring sperm-bound ASAs 335 in satisfactory breeder beef bulls and bulls with reproductive pathology. 336

In conclusion, a direct technique to detect sperm-bound ASAs in bull semen was developed. Flow cytometry was accurate and reliable for detection of sperm-bound ASAs and discrimination between ASA-positive and ASA-negative bulls. When combined with fluorescence microscopy, this method provided an ideal diagnostic approach for objective and quantitative evaluation of sperm-bound ASAs in bulls.

342

343 Disclosure statement

344

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347	
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349	
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359	
360	References
361	
362	[1] Romrell LJ, O'Rand MG. Capping and ultrastructural localization of sperm surface
363	isoantigens during spermatogenesis. Dev Biol 1978;63:76-3.
364	[2] Comhaire FH, Mahmound AMA, Depuydt CE, Zalata AA, Christophe AB. Mechanisms and
365	effects of male genital tract infection on sperm quality and fertilizing potential: the andrologist's
366	point of view. Human Reprod Update 1999;5:939-98.

- 367 [3] Zralý Z, Benová J, Šišák M, Diblíková I, Švecová D, Zajícová A, Vêžník Z. Occurrence of
- antibodies to sperms in blood sera of bulls and boars. Vet Med Czech 1998;43:197-204.
- 369 [4] Hegazi AG, Ezzo O. Serum auto antibodies in buffaloes and cattle naturally infected with
- 370 IBRV and brucella. Buffalo J 1995;3:325-30.
- [5] Perez T, Carrasco LW. Autoimmunizacion espermatica en sementales bovinos como causa de
- subfertilidad. IVth Int Congr Anim Reprod Artif Insem 1964; 5:527.
- [6] Vlok I, Ferrer M, Sardoy M, Anderson D. Serum anti-sperm antibodies associated with
- orchitis in a bull. Clinical Theriogenology 2009;1:251.
- [7] Wang XW, Ding GR, Shi CH, Zeng LH, Liu JY, Li J, Zhao T, Chen YB, Guo GZ.
- 376 Mechanisms involved in the blood-testis barrier increased permeability induced by EMP.
- 377 Toxicology 2010;276:58-63.
- [8] Kim CA, Parrish JJ, Momont HW, Lunn DP. Effects of experimentally generated bull
- antisperm antibodies on in vitro fertilization. Biol Reprod 1999; Jun;60(6):1285-91.
- [9] Coonrod SA, Westhusin ME, Naz RK. Monoclonal antibody to human fertilization antigen-1
- (FA-1) inhibits bovine fertilization in vitro: application in immunocontraception. Biol Reprod
  1994;51:14-23.
- 383 [10] Coonrod SA, Herr JC, Westhusin ME. Inhibition of bovine fertilization in vitro by
- antibodies to SP-10. J Reprod Fertil 1996;107:287-97.
- 385 [11] Yeh WR, Acosta AA, Seltman HJ, Doncel G. Impact of immunoglobulin isotype and sperm
- surface location of antisperm antibodies on fertilization in vitro in the human. Fertil Steril
- 387 1995;63:1287-92.

- 388 [12] Krause WKH. Sperm functions influenced by immune reactions. In: Immune Infertility,
- Krause WKH and Naz RK, Eds., Springer Verlag Berlin Heidelberg, Berlin, Germany. 2009, p.
  49-65.
- [13] Eggert-Kruse W, Christmann M, Gerhard I, Pohi S. Klinga K, Runnenbaum B. Circulating
- antisperm antibodies and fertility prognosis: a prospective study. Hum Reprod 1989;4:513-20.
- [14] Chamley LW, Clarke GN. Antisperm antibodies and conception. Semin Immunopathol
  2007; 29:169–84.
- 395 [15] Räsänen ML, Hovatta OL, Penttilä IM, Agrawal YP. Detection and quantitation of sperm-
- bound antibodies by flow cytometry of human semen. J Androl. 1992;13:55-64.
- 397 [16] Wright PJ. Serum spermagglutinins and semen quality in the bull. Australian Vet J
  398 1980;56:10-13.
- 399 [17] Purswell BJ, Dawe DL, Caudle AB, Williams DJ, Brown J. Spermagglutinins in serum and
- seminal plasma of bulls and their relationship to fertility classification. Theriogenology1983:20:375-81.
- [18] Menge AC, Christian JJ Jr. Effects of auto- and iso-immunization of bulls with semen and
  testis. Int J Fertil 1971; 16:130-8.
- 404 [19] Fayemi O. Sperm antibodies and reproductive efficiency in the zebu cattle in south-western
- 405 Nigeria. Pakistan Vet J 2005; 25:111-4.
- 406 [20] Zralý Z, Benová J, Diblíková I, Švecová D, Kummer V, Mašková J, Vêžník Z. Antisperm
- 407 antibodies in blood sera of bulls and correlations with age, breed and ejaculate quality. Acta Vet
- 408 Brno 2002;71:303-8.

- 409 [21] Nikolaeva MA, Kulakov VI, Korotkova IV, Gulobeva EL, Kuyavskaya DV, Sukhikh GT.
- 410 Antisperm antibodies detection by flow cytometry is affected by aggregation of antigen-antibody
- 411 complexes on the surface of spermatozoa. Human Reprod 2000;15:2545-53.
- 412 [22] Krapez JA, Hayden CJ, Rutherford AJ, Balen H. Survey of the diagnosis and management
- 413 of antisperm antibodies. Hum Reprod 1998;13:3363-7.
- 414 [23] Santoso S, Zimmerman U, Neppert J, Mueller-Eckhardt C. Receptor patching and capping
- of platelet membranes induced by monoclonal antibodies. Blood 1986;67:343-9.
- 416 [24] Nicholson SC, Robinson JN, Sargent IL, Barlow DH. Detection of antisperm antibodies in
- seminal plasma by flow cytometry: comparison with the indirect immunobead binding test. FertilSteril 1997;68:1114-9.
- [25] Chenoweth PJ, Hopkins FM, Spitzer JC, Larsen RE. Guidelines for using the bull breeding
  soundness evaluation form. Clin Theriogenology 2010;2:43-9.
- [26] Mestecky J, Fultz PN. Mucosal immune system of the human genital tract. J Infect Dis
  1999;179(suppl 3):S470-4.
- 423 [27] Knee RA, Hickey DK, Beagley KW, Jones RC. Transport of IgG across the blood-luminal
- 424 barrier of the male reproductive tract of the rat and the effect of estradiol administration on
- reabsorption of fluid and IgG by the epididymal ducts. Biol Reprod 2005;73:688–94.
- 426 [28] Wang Y, Ben K, Cao X, Wang Y. Transport of anti-sperm monoclonal IgA and IgG into
- 427 murine male and female genital tracts from blood. Effect of Sex Hormones. J Immunol
- 428 1996;156:1014-9.
- 429 [29] Cunningham KA, Carey AJ, Finnie JM, Bao S, Coon C, Jones R, Wijburg O, Strugnell RA,
- 430 Timms P, Beagley KW. Poly-immunoglobulin receptor-mediated transport of IgA into the male

- 431 genital tract is important for clearance of Chlamydia muridarum infection. Am J Reprod
- 432 Immunol 2008; 60:405–14.
- 433 [30] Brokstad KA, Cox RJ, Olofsson J, Jonsson R, Haaheim LR. Parenteral influenza
- 434 vaccination induces a rapid systemic and local immune response. J Infect Dis 1995;171:198-203.
- 435 [31] Richards JM, Witkin SS. Non-immune IgG binding to the surface of spermatozoa by
- disulphide arrangement. Clin Exp Immunol 1984;58:493-501.
- 437 [32] O'Rand MG, Metz CB. Immunofluorescence of rabbit spermatozoa treated with anti-sera to
- 438 a single sperm surface membrane glycoprotein and wheat germ agglutinin. J Cell Biol
- 439 1975;67:317a.
- 440 [33] Gaunt SJ, Brown CR, Jones R. Identification of mobile and fixed antigens on the plasma
- 441 membrane of rat spermatozoa using monoclonal antibodies. Exp Cell Res 1983;144:275-284.
- 442 [34] Koehler JK. Studies on the distribution of antigenic sites on the surface of rabbit
- 443 spermatozoa. J Cell Biol 1975;67:647-59.
- 444 [35] Schmid I, Ferbas J, Uittenbogaart CH, Giorgi JV. Flow cytometric analysis of live cell
- proliferation and phenotype in populations with low viability. Cytometry 1999;35:64-74.

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Fig. 1. Example of dot plot distribution of forward (FSC-H) and side scatter (SSC-H) of a washed sperm sample (left panel). The cells within gate 1 (R1) represent the population of spermatozoa. Example of dot plot distribution of two-color analysis of a sperm sample from a bull with experimentally-induced anti-sperm antibodies stained with FITC-labeled anti-mouse IgA (isotype control) (central panel) or FITC-labeled anti-bovine IgA (right panel). Fluorescence data was collected with logarithmic amplification for green (FITC; FL1-H) and red (PI; FL2-H) fluorescence. The anti-sperm antibody (ASA)-negative dead sperm appeared in the upper left (UL) quadrant, ASA-negative live sperm in the lower left (LL) quadrant, ASA-positive dead sperm in the upper right (UR) quadrant, and ASA-positive live sperm in the lower left (LR) quadrant. 



Fig. 2. Percentage of anti-sperm antibody (ASA)-bound spermatozoa in samples fixed with
formalin buffer solution and non-fixed samples, and including live only or live and dead cells in
the analysis. <sup>a,b</sup>Values with different superscript differ significantly among treatments within
ASA-negative bulls (Median ± SE).



