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Pregnancies following artificial insemination with spermatozoa from problem stallion ejaculates processed by Single Layer Centrifugation with Androcoll-E.

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Running head: fertility of SLC-selected stallion spermatozoa
Abstract

Some stallions produce ejaculates of low quality and/or low fertility when used for artificial insemination (AI). The purpose of these five case studies was to use Single Layer Centrifugation (SLC) to select the best spermatozoa from “problem” ejaculates for subsequent use in AI. Sperm quality, in terms of motility, morphology and chromatin integrity, was improved in the SLC-selected samples compared to the corresponding uncentrifuged samples, with the exception of one stallion thought to have ampullary stasis. In this stallion, neither the incidence of spermatozoa with detached heads nor the proportion of damaged chromatin was decreased by SLC, in contrast to previous results. Pregnancies were obtained after using SLC-selected spermatozoa from the five stallions for AI, indicating that the spermatozoa were functional after SLC. Overall, the results suggest that SLC may be useful when preparing AI doses from some “problem” ejaculates.

Key words: stallion spermatozoa, SLC, sperm quality, sperm fertility, AI trial.

Introduction

A method for improving quality in sperm doses to be used for artificial insemination (AI) is desirable for dealing with some “problem” stallion ejaculates. These can be considered problematic because of their semen characteristics, for example, a low sperm concentration (less than 100 x10^6/mL) in a large volume (sometimes over 100 mL of gel-free ejaculate), or specific problems with sperm morphology or motility. Alternatively, spermatozoa may not survive cooling, hence restricting their use to on-site AI. Previously some problem ejaculates have been processed by “sperm washing”, centrifuging the extended semen to pellet the spermatozoa and removing some of the
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45 seminal plasma, thereby effectively raising the sperm concentration. However, although this procedure results in a transient increase in progressive motility (Parlevliet & Colenbrander, 1999; Brinsko et al, 2000; Aurich, 2005; Love et al, 2005), it may cause chromatin damage (Morrell et al, 2010a). Density gradient centrifugation (DGC) has been used to select good quality spermatozoa from the rest of the ejaculate and pregnancies have been generated following low-dose AI (deep intrauterine horn; Varner et al, 2008). However, this method of AI is not in widespread use and the sperm yield from DGC is too low (Edmond et al, 2008) for conventional AI. Therefore, new handling strategies are required to enable “problem” ejaculates to be processed for conventional AI.

50 A new method for selecting the best spermatozoa has been developed at the Veterinary Faculty of the Swedish University for Agricultural Sciences (SLU), using Single Layer centrifugation (SLC) through species-specific colloids (Androcoll™). SLC-selected sperm samples have good motility, normal morphology, intact plasma membranes and good chromatin integrity (reviewed by Morrell & Rodriguez-Martinez, 2010). The objective of the current case study was to determine if SLC could be used to select sufficient good quality stallion spermatozoa from “problem” ejaculates to be used in conventional AI.

Materials and Methods

Animals and semen collection

65 Warmblood (2) and trotter stallions (3) were housed under standard husbandry conditions at commercial studs, (the Swedish National Stud, Flyinge, Sweden; Alebäck Stud Farm, Lidköping, Sweden) or at the University of Bologna, Italy. The stallions had
been judged to produce “problem” ejaculates based on semen characteristics and/or low pregnancy rates (0-20%) following AI with cooled shipped semen doses in the field. All of the stallions were in use as breeding sires for commercial AI. Semen was collected by allowing the stallions to mount a phantom and ejaculate into a warmed artificial vagina (Colorado or Missouri type, depending on the individual). The semen was collected into an insulated glass bottle fitted with a filter to capture gel. Immediately after collection, the ejaculate was extended 1:1 with warm extender, which was either Kenney’s extender (Kenney et al, 1975) or INRA96 (IMV, l’Aigle, France) or skimmed milk medium (modified from Heitland et al, 1996, ie without egg yolk and glycerol) at 37°C.

Semen processing by SLC (all stallions)

The SLC procedure was carried out using Androcoll-E (SLU, Uppsala, Sweden), using one of two protocols, according to the volume of extended ejaculate available. Briefly, where possible, the semen was extended to give an approximate sperm concentration of 100x10⁶/mL, which had previously been determined to be optimal for SLC (Morrell et al, 2010c). The following volumes were used: 4.5 mL extended semen on top of 4 ml Androcoll-E (SLC-Small) or up to 18 ml extended semen on top of 15 ml Androcoll-E (SLC-Large). After centrifugation at 300g for 20 min in a bench centrifuge with a swing-out rotor, the resulting sperm pellets were removed and resuspended in fresh extender for cooling and transporting as semen doses for AI. Where possible, an aliquot of the sperm suspension was removed for quality control. The insemination was carried out by the veterinarian at the receiving stud according to standard practice.

Case Studies
Stallion 1 was a warmblood stallion, 8-9 years old, which had had a low pregnancy rate in several breeding seasons. Aliquots from four ejaculates were made available for SLC-Small for laboratory assessment only (subjective motility; Morrell et al, 2009a; morphology and chromatin integrity Morrell et al, 2009b), and a further three ejaculates were processed by SLC-Large for AI doses.

Stallion 2 was a 10-year old warmblood stallion, producing ejaculates with low or no progressively motile spermatozoa and a high proportion of detached heads. A diagnosis of probable ampullary stasis was made by the stud veterinarian. However, repeated collections over a two month period to “clean out” the partial blockage and remove the accumulated spermatozoa did not resolve the issue of poor sperm quality. In the previous breeding season, aliquots of three ejaculates had shown improved sperm quality after SLC, (Morrell et al, 2010b). Aliquots from five ejaculates were used for SLC with Androcoll™-E-Large for AI in five mares. Two mares became pregnant, two mares were not pregnant to this AI and the last mare was barren throughout the season.

Stallion 3, was an 11-year old trotter stallion, which had been producing large volume ejaculates with a low sperm concentration, thus causing a problem in obtaining sufficient spermatozoa for a standard sperm dose (1 billion motile spermatozoa in Sweden) in a manageable volume (ie ≤20 mL). However, on the day of the SLC, the ejaculate characteristics were normal (volume 50 mL, and sperm concentration 235 x10⁶/mL). An aliquot of extended semen (15 mL) was prepared by SLC with Androcoll-E-Large.

Sperm motility was assessed subjectively by observer 3 (GK), and further aliquots were evaluated for viability, morphology and acrosome staining (Kútölgyi et al., 2006). The spermatozoa were classified according to morphology (Table 1), viability and acrosome
status (Figure 1). The SLC-selected sperm sample was used for AI in one mare on the
day of SLC, which became pregnant.

Stallion 4 was a 13-year old trotter, referred to GM by the stud veterinarian because of
low sperm concentration and poor sperm survival in cooled semen doses. No pregnancies
had been obtained from semen doses prepared by conventional semen processing in the
previous breeding season. Two ejaculates were collected with the following semen
characteristics: (i) volume 60 mL, sperm concentration 45 million/mL, motility 70%; (ii)
volume 65 mL, sperm concentration 52 million/mL, motility 65%. On both occasions, the
ejaculate was extended 1:1 with skimmed milk semen extender (after Heitland et al,
1996) and then subjected to cushion centrifugation (Cushion Fluid, Minitube, Tiefenbach,
Germany) at 00 g for 13 min to concentrate the spermatozoa. The resulting sperm sample
was split, with one part of the first ejaculate being used for cushion centrifugation to
remove some of the seminal plasma followed by DGC (their usual method of
preparation), and the other being used for centrifugation using Androcoll –E Small.
Sperm motility was evaluated by CASA (Mari et al, 2010). The motility results showed
increased total and progressive motility in the centrifuged samples than in the
uncentrifuged samples (Table 1), although there was no difference between the two
centrifugation treatments (DGC and SLC). For the first ejaculate, the DGC sperm
preparation was used for AI in one mare but did not result in a pregnancy. For the second
ejaculate, the SLC-sperm preparation was used for AI in one mare, which became
pregnant.

Stallion 5 was a 24-year old trotter, referred to GM by the stud veterinarian because of
large semen volume, relatively poor quality (approximate values: volume 100 mL, sperm
concentration < 200 million/mL, motility 50%), and poor sperm survival in cooled semen. The pregnancy rate after AI with conventionally prepared semen doses in the previous breeding season had been approximately 20%. Eight ejaculates were collected and were immediately extended in skimmed milk extender (as for Stallion 4). One aliquot was used for cushion centrifugation at 400 g for 13 min with 3 ml cushion fluid (T1), and the other (15 mL) was subjected to SLC with Androcoll-E Large at 300 g for 20 min (T2) (Morrell et al, 2009c). Sperm pellets were resuspended in fresh skimmed milk medium, and all sperm samples including the untreated aliquots (N) were used for CASA (Mari et al, 2010), immediately (0h) and after 24h. In some cases, namely the last three ejaculates for Stallion 5, sperm motility was also analysed after 48h. The remainder of the sperm samples from T1 and T2 were used for AI after 24h. Mean values for motility were compared using student`s t test, with significant differences defined as P<0.05. The CASA motility results are shown in Table 1. Total motility and progressive motility were significantly better for T1 and T2 than for N (P<0.05). There was no difference between T1 and T2 at 0h or 24h, although T2 (the SLC-samples) had better mean values than T1 (sperm washing) after 48h cold-storage (P<0.05). For AI with T1, 5 out of 11 mares became pregnant and for T2, three out of eight mares became pregnant.

Discussion

Previously, sperm quality had been improved (according to laboratory analyses) by SLC of stallion spermatozoa using Androcoll-E (Morrell & Rodriguez-Martinez, 2010). The results presented here indicated not only that sperm quality could be improved by SLC for four out of the five stallions, but also that sufficient spermatozoa could be
recovered after SLC-processing by stud personnel for conventional AI resulting in pregnancies.

For stallions 4 and 5, a comparison was made between SLC and “sperm washing” i.e. centrifugation without a colloid. Previous studies have shown that SLC is better than “sperm washing” in that selection is made for motile spermatozoa with good chromatin integrity, whereas sperm washing samples show only a transient improvement in sperm motility and no improvement in chromatin integrity (Morrell et al, 2010a). In the present study, washed samples and SLC-selected samples had better motility than the corresponding uncentrifuged samples, although there was no difference in sperm motility between the two methods at t0 or t24. By t48, however, sperm motility in the washed samples was deteriorating whereas the SLC-samples retained their motility. There was no difference in pregnancy rates between the two treatment groups (AI at 24 h after SLC), although the sample size was small.

Unfortunately it was not possible to perform “control” AI, i.e. using non-SLC selected spermatozoa, for any of the ejaculates described here because the low quality of the untreated ejaculates precluded them from being used for commercial AI doses. A stud does not wish to harm its reputation by supplying poor-quality semen doses for AI.

However, there was sufficient documented evidence of the poor fertility of some of these stallions (stallions 1 and 5 each had 20% pregnancy rate previously from AI with unselected spermatozoa; semen from stallions 2 and 4 was deemed to be of too poor quality to be used for AI doses) to be confident that the improvements in sperm quality seen in the laboratory analyses were likely to be associated with the pregnancies following AI with SLC-selected spermatozoa. Of course, it should be born in mind that
the number of mares used for AI was very low, but even so the results obtained are encouraging and would appear to offer a solution for equine breeders faced with these problem stallions. Our aim in publishing these results is to make public the existence of this procedure so that equine breeders can choose if they want to use it for problem stallions or not. In this way it should be possible to recruit more mares and more stallions to the project, thus eventually achieving a larger sample size than has been possible to date. The problem stallions used thus far represent some examples of the many different types found in the field; the study is continuing, including more stallions with different types of problems.

It was interesting to note that the SLC-samples from stallion 2, which was thought to have temporary ampullary stasis, did not show an improvement in normal morphology or chromatin integrity, compared to the non-SLC samples. It appeared that the detached heads pelleted along with the normal spermatozoa during centrifugation, a phenomenon that has been reported previously (Morrell et al, 2010b). Moreover, the SCSA result indicated more chromatin damage in the SLC-sample than in the non-SLC sample from the same ejaculate. These results are contrary to the results reported for more than 100 ejaculates, where both normal morphology (±11%; P<0.001) and chromatin damage (-5.1%DFI; P<0.001) were improved in the SLC samples compared to the corresponding uncentrifuged samples (Morrell et al, 2010c), and are also in contrast to results from sperm samples from the same stallion taken the previously when he was not affected by ampullary stasis. It is interesting to speculate whether the raised %DFI in the SLC samples from this stallion was due to the presence of the detached heads, which had been concentrated by the centrifugation process. Alternatively, ampullary stasis per se could
have a detrimental effect on chromatin integrity in the centrifuged samples, perhaps due to changes in the seminal plasma affecting the sperm membranes and making them more susceptible to damage during centrifugation, thus causing a small rise in %DFI.

Further studies are underway to identify which types of problem ejaculates can be improved by Single Layer Centrifugation as a practical solution to improving stallion sperm quality. In the meantime, it can be concluded from the results presented here that SLC can be used by stud personnel to process “problem” ejaculates, that sperm quality is generally improved in the SLC-selected samples and that the SLC-selected spermatozoa are capable of fertilization after AI.

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**Conflict of Interest:**

None

**Acknowledgement:**

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**References**


Morrell JM, Johannisson A, Dalin A-M, Rodriguez-Martinez H, 2009: Single Layer Centrifugation with Androcoll-ET can be scaled-up to allow large volumes of stallion ejaculate to be processed easily. Theriogenology 72 879-884.


Table 1: Summary of results for “problem” ejaculates from five stallions processed by SLC.

<table>
<thead>
<tr>
<th>Case Study</th>
<th>Sperm quality before and after SLC</th>
<th>AI results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stallion 1</td>
<td>motility 60±10% versus 77.5±2.9%, normal morphology 34±6.3 versus 59.5±24.3, %DFI 18±4.5% versus 7±1.1%</td>
<td>Control 1/5, SLC 2/3</td>
</tr>
<tr>
<td>Stallion 2</td>
<td>Previous year: motility 65% versus 90%, normal morphology 68% versus 89.5%, %DFI 18.6% versus 3.4% Problem year: viability 67% versus 69%; % DFI 19% versus 24% for SLC-selected and control samples, respectively). Normal morphology in control 67% but 11% detached heads</td>
<td>Control not done (quality too poor); SLC 2/4 (plus one barren mare inseminated)</td>
</tr>
<tr>
<td>Stallion 3</td>
<td>Motility 70% versus 90% at 0h, 40% versus 80% after 24h. Viability 66% versus 81% at 0h, 51% versus 71% after 24h; Morphology 71% versus 86% at 0h, 74% versus 84% after 24h.</td>
<td>SLC 1/1</td>
</tr>
<tr>
<td>Stallion 4</td>
<td>CASA motility: Expt.1 Control TM 65% PM 17%; DGC TM 86% PM 36%; SLCTM 88% PM 30% CASA motility: Expt 2 Control TM 71% PM 23%; DGC TM 85% PM 25%; SLCTM 77% PM 36%</td>
<td>DGC 0/1 (expt. 1); SLC 1/1 (expt. 2)</td>
</tr>
<tr>
<td>Stallion 5</td>
<td>CASA 24h: control TM 20% PM 6%; cushion cent TM 30% PM 12%; SLC 29% PM 14%. CASA 48h: control TM 10% PM 1%; cushion cent TM 19% PM 5%; SLCTM 20% PM 11%.</td>
<td>Cushion cent 5/11: SLC 3/8</td>
</tr>
</tbody>
</table>
Figure 1: Stallion spermatozoa stained with Chicago Sky Blue and Giemsa, showing examples of live and dead spermatozoa (a), acrosome status and various morphological abnormalities (a and b).