The seminal coagulum favours passage of fast-moving sperm into the uterus in the black-handed spider monkey

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Abstract

In addition to gametes, mammalian internal fertilisation has required the evolution of assorted anatomical, physiological and biochemical devices to deal with intra- and inter-sexual conflict such as sperm competition and female cryptic choice respectively. The seminal coagulum of primates and other mammals is viewed as one of such devices. Among primates, the seminal coagulum characteristically occurs in multi-male and multi-female species, leading us to suppose that it intervenes in sperm competition. However, it can also provide cues to the female reproductive tract about male desired or undesired traits, and therefore deter or favour sperm survival and migration. The present work investigates whether the seminal coagulum of the black-handed spider monkey enhances sperm fertilisation chances by improving the female reproductive tract conditions, and if the female reproductive tract is ‘blind’ to semen or behaves selectively towards ejaculates of different males. A series of artificial inseminations were done in five females, using the ejaculates of three different males, one at a time, and measuring the presence of distinct types of sperm inside the uteri at 10, 30 and 60 min following the insemination. The presence of coagulum, menstrual phase, and male and female identity only affected fast, straight-moving sperm, with larger amounts of fast sperm appearing inside the uterus when ejaculates had seminal coagulum, as well as when in the periovulatory phase. There was great intra-uterine fast-sperm variation regarding which male’s semen inseminated which female. The results provide evidence to account for sexual conflict in the spider monkey as well as a methodological approach to this kind of study.


Introduction

In animal species with internal fecundation, fertilising involves more than a proper ejaculation in the female reproductive tract (Gomendio et al. 1998, 2007, Satake et al. 2006, Immler 2008). In particular, if contending males are able to deposit semen in the same female, in turn, the female requires control of the fate of the sperm deposited within its reproductive tract in order to enhance its own and offspring’s fitness and override any reproductive costs inflicted by males. Such was the scenario in which post-copulatory male–male competition (Parker 1998, Birkhead & Pizzari 2002) and female cryptic mate choice (Eberhard 1996, 1998) evolved. Therefore, in addition to ornaments and behaviour traits, females and males must evolve further anatomical, physiological and biochemical devices that account for such observed post-copulatory reproductive success (Eberhard 1996, 1998, Gomendio et al. 1998, Parker 1998, Birkhead & Pizzari 2002, Immler 2008). Such events in most vertebrates, and particularly in mammals, take place within the female reproductive tract, requiring a physiological rather than a behavioural approach for their study.

Among male primates, the seminal coagulum produced by the prostate, the seminal vesicle and the bulbourethral glands is likely one of such devices (Dixson 1998). Comparative analyses on the presence or absence of seminal coagulum in the ejaculate of different mammal species suggest that this structure evolved mainly to deal with male–male competition (Dixson & Anderson 2002, 2004). However, there is a possibility that the seminal coagulum also has a role in female cryptic choice. Such a view is supported by the fact that is a major trait showing variability (Sato et al. 2007, Hernández-López et al. 2008), providing females (or their reproductive tracts) a substrate for male quality assessment. Since different glands contribute to its formation, a small variation in the amounts of constituents contributed by each gland yields structurally diverse seminal coagula, where changes in its components, independently from sperm concentration, severely impair fertility (Rossato et al. 2002, Mikhailichenko & Esipov 2005, Khosrowbeygi & Zarghami 2007, de Lamiranda 2007).
The first barrier met by spermatozoa on their way to fertilise oocytes is vaginal acidity, since an acidic vaginal milieu immobilises and damages sperm (Carr et al. 1985, Boskey et al. 1999, 2001). The vaginal tract acidity of female mammals varies throughout the uterine cycle (Bauman et al. 1982, Dixson 1998, Boskey et al. 1999, 2001, Suarez & Pacey 2006), depending on the fluctuations in steroid hormone levels and bacterial growth. As ovulation approaches, the vaginal pH becomes more acidic due to an increase in glycogen accumulation promoted by steroid hormones and by the conversion of glycogen to lactic acid by vaginal bacteria (Bauman et al. 1982, Papka & Williams 1998, Boskey et al. 1999, 2001). By contrast, glycogen becomes less abundant and vaginal pH turns to neutral throughout the luteal phase (primates) or metestrus (other mammals) of the uterine cycle, due to the decrease in the levels of steroid hormones (Bauman et al. 1982, Papka & Williams 1998, Boskey et al. 1999, 2001). Increased vaginal acidity throughout the periovulatory phase has been ascribed the function of protecting females against sexually transmitted disease (Castle et al. 2002, Olmsted et al. 2005). However, in most primates, sexual intercourse occurs throughout the entire menstrual cycle (Dixson 1998), leaving females at the risk of contracting a venereal disease at all times. Therefore, the acid periovulatory milieu could well be controlling the fate of the sperm deposited within the vaginal tract.

The seminal coagulum is slightly alkaline (7.2–8.0 in humans) and acts to neutralise vaginal acidity (Prins 1998), while ascorbic acid (Song et al. 2006) and l-carnitine (De Rosa et al. 2005) protect sperm DNA from denaturation. The seminal coagulum also provides a viscous matrix preventing sperm diffusion out of the ejaculate, and in many primate species contains semenogelins as well as prostate-specific antigen and prostatic acid phosphatase, the enzymes needed to induce ejaculate clotting and its further liquefaction respectively (Lilja 1993, Dorus et al. 2004, Clark & Swanson 2005). In addition, semenogelins induce sperm quiescence to avoid spending energy during migration throughout the male tract (Robert & Gagnon 1996, Cooper 1998, Rossato et al. 2002, de Lamiranda 2007). Furthermore, prostaglandins found in the seminal coagulum induce contractions of the smooth musculature of the female tract to enhance sperm transport (Prins 1998, Suarez & Pacey 2006). All these add up to protect sperm from a hostile female environment, rather than merely obstruct sperm from other males. However, sperm migration up the female tract is not accomplished merely by sperm movement, but aided by vaginal, cervical and uterine contractions, and by cilia movement of the tract wall (Suarez & Pacey 2006).

The black-handed spider monkey (Ateles geoffroyi) is considered a promiscuous species (Dixson 1998, Dixson & Anderson 2002), although little is known of its sexual behaviour in the wild. Notwithstanding, relative testes size (Harcourt et al. 1981), as well as the presence of genes coding for enzymes promoting clotting and liquefaction of the seminal coagulum (Dorus et al. 2004, Clark & Swanson 2005), supports the notion that post-copulatory sperm competition and female cryptic choice occur in this species. Unlike in man and chimpanzees (Lilja 1993, Dixson 1998), spontaneous in vitro liquefaction of the coagulum does not happen in the spider monkey (Hernández-López et al. 2002a, 2002b). Therefore, activation of the enzymes promoting liquefaction might be dependent on vaginal physicochemical cues, such as temperature or pH (Carr et al. 1985, Gundlach & Luttermann-Semmer 1987, Rossato et al. 2002), or even complex biochemical interactions between the vaginal milieu and the ejaculate, such as enzyme–enzyme interactions. In any case, if coagulum liquefaction is required from the vaginal milieu, it is likely that females are able to control the fate of ejaculates within their reproductive tract, and thereby fertilisation by cues informing about male fitness, histocompatibility, etc., provided by this structure. The present work was done to investigate by means of artificial insemination (AI) the in vivo role of the seminal coagulum in the supporting passage of diverse types of sperm from the vagina into the uterus, along with the part played by the vaginal milieu in such a promotion. We reasoned that coagulum dilution should be reflected in larger proportions of linearly motile spermatozoa inside the uterus, concomitant with a reduction in the percentage of immotile sperm. Since an acidic vaginal milieu immobilises and damages a sperm (Bauman et al. 1982), while coagulum alkalinity overturns it (Prins 1998), we compared sperm motility depending on whether or not the male emitted the seminal coagulum. We tested 1) whether ejaculates containing the coagulum show better pro-fertilising activity that when it is missing (Harper 1994) and 2) if liquefaction of the coagulum and types of sperm passing through the cervix into the uterus is dependent upon natural variation of local vaginal temperature or pH. In other words, we expected to find inside the uterus greater proportions of straightforward-swimming sperm in the follicular and peri-ovulatory phases of the menstrual cycle, when optimal fertilisation is more likely to occur. Finally, looking for evidence of individual female participation in the above-described issues, we tested 3) whether the types of sperm found in the uteri were related to the identity of the sperm donor.

Results

Vaginal pH and temperature variations after and following AI

Basal vaginal pH varied significantly throughout the menstrual cycle ($F_{2,38,13} = 4.74$, $N = 169$, $P = 0.014$); during the peri-ovulatory phase it turned to be significantly
(P<0.05) more acidic (mean±S.E.M.: 6.5±0.09) than that in the luteal phase (7.0±0.13), but not more than that in the follicular phase (6.7±0.12). Basal vaginal temperature did not change throughout the menstrual phases (overall mean: 38±0.08 °C). Semen pH did not vary, being always 8, whether or not the seminal coagulum was emitted. Vaginal temperature following AI was slightly but significantly colder when semen lacked seminal coagulum (coagulum absent: 37.2±0.12 °C; coagulum present: 37.8±0.1 °C; F_{1,33.14}=5.21, P=0.03), menstrual phase having no effect (F_{2,26.69}=0.38, N=169, P=0.7). We found a vaginal pH turnover after performing the inseminations, related to both the menstrual cycle phase and coagulum incidence (F_{2,24.67}=6.16, N=169, P=0.007). Irrespective of the presence or absence of seminal coagulum, semen always turned vaginal pH towards alkaline (follicular–periovulatory: 7.6±0.2), except in the luteal phase, where pH turned to completely basic when the ejaculates lacked coagulum (8.7±0.3), while those where the coagulum was present turned neutral (7.2±0.2).

**Sperm migration**

Movement of sperm was classified into four types (Table 1). Excluding fast, linearly-moving sperm (FLM), there were no significant effects involving types of sperm collected within the uterus in relation to menstrual phase, coagulum incidence and time of collection. FLM occurrence through time significantly changed depending on whether the coagulum was present or absent and the menstrual phase (F_{4,59.05}=3.76, N=1014, P=0.009). As is appreciated in Fig. 1A, when the ejaculate contained seminal coagulum, FLM arrived earlier to the uterus, continuing to do so up to 30 min past the insemination in the periovulatory phase. An hour after the insemination, few FLM were collected from the uteri. In contrast, similar inseminations performed in other menstrual phases showed different time patterns. In the follicular phase, FLM arrival increased at 30 min following the insemination, extending up to 1 h after the insemination. In the luteal phase, few FLM were collected at all times. Different patterns of intra-uterine FLM occurred when the semen lacked coagulum (Fig. 1B). Seemingly, the FLM arrival was greater in the luteal phase, particularly at 30 min past the insemination. However, a single insemination was responsible for such an increase at 30 min; removing this value returns the FLM values to averages observed in other menstrual phases (3.6±3.3%). Although at 10 min the mean number of intra-uterine-collected FLM was not much different from that when inseminations included the seminal coagulum, the large variation suggests a random rather than a coordinated release and migration.

Concerning other types of sperm, immotile sperm (IMM) showed a trend to vary in relation to menstrual phase and coagulum incidence (F_{2,40.98}=3.02, N=1014, P=0.06). When the coagulum was present, intra-uterine IMM occurred less in the periovulatory (39.4±8.03%) than that in the follicular (49±10.38%) and luteal (55.53±10.33%) phases. Contrastingly, in the absence of coagulum, more IMM appeared in the periovulatory phase (60.10±8.43%) than in the follicular (57.05±12.97%) phase, while decreasing in the luteal phase (27.45±13.91%). Although both

### Table 1

<table>
<thead>
<tr>
<th>Description of sperm movement</th>
<th>Classification used in the present paper (abbreviation)</th>
<th>WHO classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fast, directed forward movement</td>
<td>Fast, linearly-moving sperm (FLM)</td>
<td>3, 3+, 4</td>
</tr>
<tr>
<td>Slow, directed forward movement</td>
<td>Slow, linearly-moving sperm (SLM)</td>
<td>2+</td>
</tr>
<tr>
<td>Undirected movement, irrespective of velocity</td>
<td>Non-linear movement (NLM)</td>
<td>1, 2, 3−</td>
</tr>
<tr>
<td>Mostly no movement</td>
<td>Immotile sperm (IMM)</td>
<td>0, 1+</td>
</tr>
</tbody>
</table>

**Figure 1** Mean (±S.E.M.) percentage of FLM collected from the uteri at 10, 30 and 60 min following artificial inseminations done in follicular, periovulatory and luteal phases of female black-handed spider monkeys. (A) Inseminations done with semen containing seminal coagulum. (B) Inseminations done with semen lacking seminal coagulum.
slow, linearly-moving sperm (SLM) and non-linearly-moving sperm (NLM) were recovered from the uteri at constant rates (SLM: 16.57 ± 1.74%; NLM: 22.91 ± 1.80%), their occurrence was unrelated to coagulum incidence or menstrual phase.

**Effects of vaginal pH and temperature on sperm migration**

Only the multiple regression between mean intra-uterine FLM and pHs and temperatures in ejaculates having seminal coagulum accounted for a significant linear relationship \((F_{4,90} = 3.3, R^2 = 0.12, P = 0.014)\). Table 2 shows the results for this multiple regression. Post-insemination vaginal pH and post-insemination temperature exerted significant effects, but not basal pH and temperature. Figure 2 shows the scatterplots and the partial residual plots for the intra-uterine FLM against post-insemination vaginal pH and temperature. Notice that despite the significant linear relationships yielded by the multiple regression analysis, data are quite scattered. An outlier point for post-insemination vaginal pH and two for post-insemination temperature were detected (boxed data points in Fig. 2). Yet, dropping these cases out from the analyses marginally reduced the relationship between the FLM and post-insemination vaginal pH \((\beta \pm \text{s.e.} = 6.87 \pm 3.43, t_{92} = 2.0, P = 0.05)\), while increasing that with temperature \((\beta \pm \text{s.e.} = 6.18 \pm 2.3, t_{91} = 2.69, P = 0.008)\). Although the examination of the P–P plot showed a fair adjustment of FLM to a normal distribution, data points in Fig. 2 do not really align in linear trends. In Fig. 2A and B, two populations of FLM are apparent, the first one is a population that does not appear to respond to vaginal pH turnover, either not appearing inside the uterus or doing so at low rates (0–10% in Fig. 2A; residual values 40–60 in 2B). The other population includes a hump-shaped scattering of data points when pH varied from 7.5 to 9.

This latter population seems to correspond to pH-responsive FLM, showing that passage to the uterus is facilitated at a vaginal pH around 7.3, is maximal at 8, and decreases as the vagina turns much more basic. A similar situation is seen concerning vaginal temperature; once more there is a collection of data points accounting for no intra-uterine occurrence of FLM or doing so at low proportions regardless of temperature variation (0–10% in Fig. 2C; residual values 180–190 in Fig. 2D). The remaining FLM data points, predominantly appearing above the regression line, show a somewhat clear tendency to augment inside the uterus as vaginal temperature increases. It is worth remembering that these vaginal pH and temperatures are local modifications produced by coagulum-containing ejaculates. When ejaculates lacked coagulum, the multiple regression also yielded a significant effect for the intra-uterine FLM \((F_{4,70} = 5.73, R^2 = 0.23, P = 0.0005)\), but not for any other type of sperm motility. Basal vaginal pH and post-insemination temperature were the variables contributing significantly to intra-uterine variation of FLM, while post-insemination vaginal pH and basal temperature did not do so (Table 2). Figure 3 shows the scatterplots and partial residual plots for these results. Besides being less abundant than when ejaculates had seminal coagulum, the intra-uterine FLM decreased as basal vaginal pH went from neutral to basic (Fig. 3A and B). On the other hand, the effect of post-insemination temperature was the same as when ejaculates contained seminal coagulum, intra-uterine FLM arrival increasing as the vagina turned warmer (Fig. 3C and D). Removing two outliers (boxed data points in Fig. 3) from the analysis did not greatly affect the results (basal pH: \(\beta \pm \text{s.e.} = -2.89 \pm 0.42, t_{68} = -6.97, P < 0.0001;\) post-insemination temperature: \(\beta \pm \text{s.e.} = 1.45 \pm 0.32, t_{68} = 4.54, P < 0.0001\)). However, the large unexplained variance in both the entire linear model and the partial correlations of pH and temperature (Table 2), as well as the number of times no FLM was found inside the uteri (zero values in Fig. 2), suggests that males are not entirely responsible for FLM passage through the cervix.

**Table 2** Multiple regression coefficients for intra-uterine collected fast, linearly-moving (FLM) in female black-handed spider monkeys in relation to basal and post-insemination pH and temperature in ejaculates having or lacking seminal coagulum.

<table>
<thead>
<tr>
<th>Variable</th>
<th>(\beta)</th>
<th>Standard error</th>
<th>Partial correlation</th>
<th>t</th>
<th>d.f.</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ejaculates having seminal coagulum</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Basal vaginal pH</td>
<td>-4.39</td>
<td>3.64</td>
<td>-0.12</td>
<td>-1.21</td>
<td>90</td>
<td>0.230</td>
</tr>
<tr>
<td>Basal vaginal temperature</td>
<td>1.01</td>
<td>4.51</td>
<td>0.02</td>
<td>0.22</td>
<td>90</td>
<td>0.820</td>
</tr>
<tr>
<td>Post-insemination vaginal pH</td>
<td>9.15</td>
<td>3.54</td>
<td>0.26</td>
<td>2.38</td>
<td>90</td>
<td>0.011</td>
</tr>
<tr>
<td>Post-insemination vaginal temperature</td>
<td>5.33</td>
<td>2.73</td>
<td>0.20</td>
<td>1.96</td>
<td>90</td>
<td>0.053</td>
</tr>
<tr>
<td>Constant</td>
<td>-262.45</td>
<td>159.17</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Ejaculates lacking seminal coagulum</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Basal vaginal pH</td>
<td>-2.89</td>
<td>0.42</td>
<td>-0.63</td>
<td>-6.97</td>
<td>70</td>
<td>0.001</td>
</tr>
<tr>
<td>Basal vaginal temperature</td>
<td>-0.38</td>
<td>0.51</td>
<td>-0.08</td>
<td>-0.75</td>
<td>70</td>
<td>0.457</td>
</tr>
<tr>
<td>Post-insemination vaginal pH</td>
<td>0.23</td>
<td>0.41</td>
<td>0.06</td>
<td>0.56</td>
<td>70</td>
<td>0.575</td>
</tr>
<tr>
<td>Post-insemination vaginal temperature</td>
<td>1.45</td>
<td>0.32</td>
<td>0.46</td>
<td>4.54</td>
<td>70</td>
<td>0.025</td>
</tr>
</tbody>
</table>

- hernandez-lopez@ucm.es, l.hernandez-lopez@ucm.es (L Hernández-López and others)


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Males and females identity effects

Again, apart from FLM and IMM in the periovulatory phase, results from analysing the males, females and males within females as fixed effects showed no significant variations. Significant variation during the periovulatory phase in FLM was associated with which male’s semen was used to inseminate each female, as well as to the presence or absence of seminal coagulum ($F_{1,38}=11.54, N=87, P=0.002$). Table 3 shows the mean distribution of the intra-uterine-collected FLM in the AI performed in the periovulatory phase in relation to male and female identities, and whether ejaculates had or lacked seminal coagulum. Depending on which male’s semen was used to inseminate which female, if coagulum was present, mean FLM varied from 3.2 to 53.5% and when it was absent, from 0 to 45.6%. In accordance with the above-mentioned results, a larger passage of FLM occurred when the coagulum was present. Even though FLM from one male, LK, was apparently more abundant, there were no between-male differences ($F_{2,38}=1.57, P=0.22$), between-female differences ($F_{4,38}=1.06, P=0.4$) and the significant male–female interaction got lost if the effect of coagulum incidence was not included ($F_{3,38}=2.10, P=0.12$).

Besides FLM, IMM in the periovulatory phase also varied significantly in relation to male and female identities ($F_{4,38}=2.88, N=87, P=0.035$), this time irrespective of coagulum incidence, ranging from 12.7 to 67.5% (Table 4). However, there was significant between-male variation ($F_{2,38}=7.93, P=0.0013$), where the IMM values of KI were significantly larger (post hoc contrasts: $P<0.05$) than those of the other two males. Moreover, including all male values, FLM and IMM were significantly and negatively correlated ($r_s=-0.53, N=45, P<0.001$).

Discussion

Although far from being conclusive, our results give evidence of sexual conflict in the black-handed spider monkey. Similar to other mammals studied so far (see Introduction), the vagina of the female spider monkey is hostile to sperm, particularly around the time of ovulation, when it is quite acidic. The sperm of the spider monkey is fairly labile; just a drop of urine in the ejaculate or cold room temperature (below 25°C) is enough to massively kill spermatozoa (L Hernández-López, AL Cerda-Molina & R Mondragón-Ceballos, unpublished results). Our results show that seminal coagulum not only adequately alkalinises vaginal pH in follicular and periovulatory phases, when fertilisation is likely to happen, but also keeps a satisfactory vaginal temperature, perhaps preventing sperm from thermal shock. Changing to a basic milieu favours spermatic motility in the fertile phases of the female (Bauman et al. 1982, Carr...
et al. 1985). However, the opposing effects of ejaculates having and lacking seminal coagulum in the luteal phase suggest additional female participation in the acidity turnover rather than a pure effect of semen. Otherwise, as semen pH was invariant, expectedly the same turnover would occur in each menstrual phase. It is not known whether in natural conditions emission of seminal coagulum occurs at all times in this species, and it certainly can be an electroejaculation artefact. However, for the purposes of the present work, such variability in seminal coagulum emission provided a control to contrast the putative functions of the coagulum.

The spider monkey is absolutely arboreal (Estrada et al. 2004), only rarely venturing to the ground (Campbell et al. 2005). Therefore, seminal coagulum might serve to prevent sperm loss from the vagina or flow back (Baker & Bellis 1993). However, our results show that as in other mammals (reviewed in Suarez & Pacey 2006), seminal coagulum promotes sperm passage through the cervix into the uterus, specifically in case of straightforward-moving sperm during the periovulatory phase. The timed intra-uterine appearance of FLM suggests a paced release from the coagulum as well as transiting through the uterus, rather than getting stored in it. By contrast, when the ejaculates lacked seminal coagulum, FLM intra-uterine arrival was erratic and inappropriate, sometimes appearing in unusually large proportions in the luteal phase (Fig. 1b), which is certainly not the most suitable time for an optimal fertilisation.

The unvarying intra-uterine presence through all menstrual phases, disregarding the presence or absence of seminal coagulum, of SLM, NLM and to some extent IMM, is likely owing to their being massively moved upward by contractions induced by constituents of the ejaculate, whereas FLM released from the seminal coagulum seemingly swam through the cervix in the periovulatory phase, when the cervical canal conditions were fit to allow such a passage (Suarez & Pacey 2006). The finding that the total intra-uterine FLM was positively correlated with the amount of vaginal alkalinisation and warming induced by ejaculates having seminal coagulum, but not, at least concerning post-insemination pH, when the coagulum was absent, gives further support to the idea of FLM being released from the coagulum and swimming through the cervix. However, the tendency of IMM to decrease in the periovulatory phase when the seminal coagulum was present, as well as the negative correlation with FLM, suggests that either additionally or alternatively the FLM were released in utero from the seminal coagulum.

Although the results described above support the fact that seminal coagulum induces vaginal modifications favourable to FLM passage through the cervix, the findings of what appear to be two populations of FLM, one increasing along with post-insemination pH and temperature, and another one insensitive to these
variations, provide evidence that such migration is not utterly provoked by the male (or the seminal coagulum), but that females are also able to exert control on the crossing of FLM through the cervical canal. The intra-uterine FLM patterning in relation to male and female identities and its negative correlation with IMM credit the latter conclusion by showing enhanced (or lessened) FLM activity, in certain male–female combinations.

Aside from its supposed role in sperm competition (Dixson & Anderson 2002, 2004), the present results acknowledge that the seminal coagulum might have evolved to deal with inter-sexual conflict. Therefore, besides maintaining spermatozoa in a quiescent state while passing from the testicles to the vagina (Carr et al. 1985, Khosrowbeygi & Zarghami 2007), the seminal coagulum turns the female vagina into an appropriate environment for sperm survival and migration (Carr et al. 1985). However, this is not entirely accomplished by the male, requiring female participation. Although still far from being compelling evidence of female cryptic choice in the spider monkey, our results acknowledge that sperm selection starts as early as the vagina, where females seemingly rely on cues provided by the seminal coagulum in favouring or opposing FLM migration.

It is noteworthy that we found no differences between males in the proportions of different types of sperm collected within the uteri, except for a significant larger percentage of IMM in one male. This contrasts with whole ejaculate evaluations where we repeatedly and consistently found significant between-male differences in both sperm concentration and proportions of sperm showing different motility (Hernández-López et al. 2002a, 2002b), but is consistent with the view that whole ejaculate assessment does not accurately reflect spermatic in vivo performance or fertility (Satake et al. 2006, Lewis 2007).

Fast, straight-swimming spermatozoa are related to fertility in other mammals (Olds-Clarke 1996, Malo et al. 2005, Satake et al. 2006, Gomendio et al. 2007), although such a relationship may not be so forthright as proposed by some authors viewing males with faster sperm as having greater chances of siring more offspring because these may be able to reach the ovum first (Malo et al. 2005, Gomendio et al. 2007). Sperm migration up the female reproductive tract is not accomplished solely by its motion, it is aided by components found in the seminal plasma and requires female assistance, such as the presence of highly hydrated cervical mucus and contractions of the uterine smooth muscle, which in turn

Table 3 Mean (S.E.M.) percentages of intra-uterine collected fast linear-moving (FLM) following artificial inseminations of female black-handed spider monkeys performed in the periovulatory phase according to which male’s semen was used to inseminate each female.

<table>
<thead>
<tr>
<th>Male</th>
<th>Female</th>
<th>AD</th>
<th>S.E.M.</th>
<th>N</th>
<th>KI</th>
<th>S.E.M.</th>
<th>N</th>
<th>LK</th>
<th>S.E.M.</th>
<th>N</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coagulum present</td>
<td>AR</td>
<td>3.4</td>
<td>2.9</td>
<td>3</td>
<td>9.3</td>
<td>5.0</td>
<td>3</td>
<td>21.0</td>
<td>7.1</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>CI</td>
<td>19.3</td>
<td>5.7</td>
<td>4</td>
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Table 4 Mean (S.E.M.) percentage of intra-uterine immotile sperm (IMM) following inseminations performed during the periovulatory phase of black-handled spider monkeys according to which male’s semen was used to inseminate each female.

<table>
<thead>
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<th>Male</th>
<th>Female</th>
<th>AD</th>
<th>S.E.M.</th>
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<th>KI</th>
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</table>

*Number of inseminations done per female (rows) using the semen of each male (columns).
can be additionally stimulated by constituents of the seminal plasma (Prins 1998, Suarez & Pacey 2006). Recent experiments (Satake et al. 2006) showed that in boars there is a subpopulation of fast forward-moving spermatozoa highly and positively responsive to the female milieu, able to swim mainly on their own past the cervix and uterus. However, in all the mammals so far studied, including humans, their results being therefore very likely to be the same in other primates, most sperm will get trapped and immobilised in and past the uterotubal junction, which is both a physical (Suarez & Pacey 2006) and chemical (Satake et al. 2006) barrier. In rabbits, the majority of these sperm are non-fertile, perhaps getting mortally damaged while being massively transported by uterine contractions (Overstreet & Cooper 1978, Scott & Overstreet 1998). Only a sub-subpopulation of the original ejaculate, spermatozoa having certain surface proteins besides the straight forceful motility, is able to traverse the uterotubal junction (Suarez & Pacey 2006, Suarez 2007), to be stored in the isthmus and to undergo capacitation and hyperactivation to finally reach the ampulla (Suarez & Pacey 2006, Rodriguez-Martinez 2007); all these involve both female and male participation. Likewise, it is not feasible to regard fast sperm in this work as utterly responsible for fertilisation in the spider monkey. Our results so far only acknowledge an initial selection of fast-swimming sperm past the cervix, which will still be subjected to even harsher selection upon reaching the oviduct (e.g. Satake et al. 2006). This subpopulation could well stand for the so-called vanguard sperm required to induce pro-fertilising changes in the upper female tract (Scott & Overstreet 1998, Suarez & Pacey 2006), whereas fertilisation is achieved either by one among those of a sub-subpopulation reaching the ampulla or by late-coming spermatozoa. Nonetheless, our work gives credit to the idea that spider monkeys can adjust locally their reproductive physiology to contend with promiscuity, as occurs in other animal species (Devine 1977, Donald & Dewsbury 1988, Parga 2003).

Ethics of experimentation
The Bioethics Committee of the Instituto Nacional de Psiquiatría Ramón de la Fuente Muñiz approved the experimental protocol and animal handling. We also followed the Mexican Official Norm of Technique Specifications for the Production, Care and Use of Laboratory Animals (NOM-062-ZOO 1999), and the Policy Statement on Use of Primates for Biomedical Purposes, as adopted by the World Health Organization (WHO) and the Ecosystem Conservation Group.

Assessment of menstrual cycles
The phase of the menstrual cycle was determined by cytological examination. To collect vaginal swabs, the females were trained to enter a wire mesh cage (1 × 1 × 1.5 m) attached to the entrance of the outdoor enclosure, and allow the introduction of a cotton swab into the vagina. For their cooperation, they were rewarded with a marshmallow or small cookies. Vaginal smears were fixed in 96% ethanol for 10 min, and stained using Shorr's Trichromic technique (Hernández-López et al. 1998). The menstrual phases (menses, follicular, periovulatory and luteal) were recognised depending on the proportional presence of four types of epithelial cells (scales, superficial, intermediate and parabasal), in addition to the presence of cervical mucus, erythrocytes and lymphocytes (Hernández-López et al. 1998).

Semen collection
Semen samples were collected by electroejaculation, as described previously (Hernández-López et al. 2002b). Briefly, the males were anaesthetised with ketamine chlorohydrate (10 mg/kg im; Anestek; CpMAX S.A. de C.V. de Mexico, D.F., Mexico); a rectal probe (12.5 mm in diameter, 9 cm long), previously lubricated (K-Y Líquido; Johnson & Johnson, Indústria e Comércio LTDA., Sao Paulo, SP, Brazil), was introduced into the rectum (~5 cm deep), and electrical stimuli were delivered. A round of stimuli (starting at 1 V and ~0.1 mA) consisted of ten electric stimulations, each one lasting 10 s, followed by a 5-s resting period. In successive rounds of stimulation, voltage and current were increased (in increments of 1.0 V and 0.1 mA respectively) until ejaculation occurred. Simultaneously, the penis was manually stimulated. The entire procedure took no more than 15 min (in the absence of ejaculation after 15 min, the procedure was stopped).

Experimental procedure
A replicate consisted of electroejaculation, intra-vaginal AI, and the subsequent recovery of sperm (from the uterus) and its evaluation. Early in the morning (0700–0800 h), one male and one female were removed from their pens. Both the monkeys were anaesthetised (ketamine chlorohydrate, 10 mg/kg im), and placed side by side on a surgical bed, with the male lying on his back and the female lying face downward, with her pelvis resting on a pillow (to elevate it). The penis of the male and the perineum of the female were cleaned with cotton (soaked in warm water), and allowed to dry. Thereafter,
Electroejaculation was done; and semen was collected in a 50 ml sterilised polypropylene conical centrifuge tube (largest diameter, 1 cm; length, 6 cm), from which the tip had been removed. The centrifuge tube, tip downwards, was inserted into the vagina, until it reached the cervix. The semen was allowed to drain by gravity. A human AI intra-uterine cannula (Laboratoire C.C.D., Paris, France) was passed through the cervix and suction applied to recover sperm from the uterus at 10, 30 and 60 min after completing AI. The timing of semen collection was based on the rate of progression of sperm (>5 mm/min) and the length of a spider monkey cervix (~30 mm; L Hernández-López, AL Cerda-Molina & R Mondragón-Ceballos, unpublished data). We did not inseminate the same number of spermatozooids per male, as is usual in this kind of experiment (Malo et al. 2005, Satake et al. 2006), because except by using trypsin (Hernández-López et al. 2002a) it is not possible to dilute the seminal coagulum, and it does not dilute spontaneously outside the vagina. Trypsin severely impairs sperm; to dilute and therefore wash the ejaculate was precluded, since once outside the seminal coagulum, the spider monkey’s sperm becomes extremely liable to temperature changes.

While performing the electroejaculation, the ejaculate was classified according to the visual presence or absence of seminal coagulum and a sample microscopically examined to assess the viability of the sample (>50% of live sperm). Sperm assessment was done by one of the experimenters, while the others proceeded with the AI in order to replicate, as closely as possible, intra-vaginal deposition of semen by natural mating. Consequently, some ejaculates classified as lacking seminal coagulum could have contained small amounts of this fraction.

Upon finishing intra-uterine semen sampling, a rapid clinical evaluation of the male was done, checking for rectal haemorrhage and assessing cardiovascular and respiratory functions. A similar evaluation was done on the females (with assessment for vaginal haemorrhage) 30 min after AI. The monkeys were placed in a 1 × 1 × 1 m stainless steel cage to allow them to recover, and afterwards they were fed and returned to their outdoor enclosure. For 7 days after semen collection and AI, the monkeys were checked daily for general health and behaviour.

Aliquots were done in the early follicular (days 3–5 of a 26-day menstrual cycle), periovulatory (days 8–12) and late luteal phases (days 23–25), with the idea of performing at least two inseminations per menstrual phase per female per male using ejaculates having seminal coagulum, and an equal number using semen lacking coagulum. This was not achieved because two females got pregnant almost at the end of the experiments (October and November 2005 respectively) and were not further used. Around three AI were done weekly, using different male–female combinations until accomplishing 169 replicates (October 4, 2004 to December 16, 2005). Each male was electroejaculated at most once every 7 days; similarly, we inseminated each female no more than once every 7 days. Ninety-four ejaculates had seminal coagulum, while 75 did not, distributed as follows: 20 AI in the follicular, 47 in the periovulatory and 27 in the luteal phases using ejaculates having coagulum; 20 AI in the follicular, 40 in the periovulatory and 15 in the luteal phases using semen lacking coagulum.

### Sperm evaluation

From the each semen sample collected from the uterus, we took two 10 µl aliquots to evaluate sperm motility. Following the WHO (1992) guidelines, we assessed the quantity and quality (movement) of sperm as shown in Table 1, since this method also has proven useful in non-human primates (Gago et al. 1999). We categorised sperm movement into four types: FLM, SLM, NLM and IMM. We reduced WHO motility classification to decrease the number of independent statistical analyses and because otherwise FLM was split into three categories, each comprising very small proportions of sperm. We assessed motility by counting 100 sperm twice, once in each 10 µl aliquot. If the average difference between both counts exceeded 5%, we took two new aliquots to assess.

### Statistical analyses

We planned the experiment to collect data from 15 consecutive menstrual cycles per female (the average cycle in the spider monkey is around 26 days: Hernández-López et al. 1998), performing three inseminations per cycle (in the follicular, periovulatory and luteal phases), each using the semen of one of the three males. These would allow performing 45 inseminations per female, where each male’s semen was used three times per menstrual phase. From such a procedure, we would end with a balanced design consisting of 225 data points (three inseminations/menstrual phase/male/female). However, we knew beforehand that ejaculation (or not) of the seminal coagulum is somewhat random and that most likely we would be unable to collect a balanced sample of ejaculates having and lacking coagulum where, besides the incidence of seminal coagula, all males, females and menstrual phases were at least proportionally, if not absolutely, represented. In addition, 48 inseminations, either because they consisted of entirely dead spermatozoa (n = 41) or the male did not electroejaculate in at most 15 min (n = 7), were not used in analyses, and ten more samples were not obtained because two females got pregnant when the experiments were almost finished. Thus, we ended with an unbalanced design matrix difficult to analyse by conventional parametric or non-parametric tests. Thus, we used general linear mixed models, employing restricted maximum-likelihood estimation (McCulloch & Searle 2001), to analyse the data, given the small number of subjects studied, their repeated sampling and the unbalanced design due to discarding some samples. We used the Shapiro–Wilk test to determine the normality of response variables: percentages of FLM, SLM, NLM and IMM collected phases were at least proportionally, if not absolutely, represented. In addition, 48 inseminations, either because they consisted of entirely dead spermatozoa (n = 41) or the male did not electroejaculate in at most 15 min (n = 7), were not used in analyses, and ten more samples were not obtained because two females got pregnant when the experiments were almost finished. Thus, we ended with an unbalanced design matrix difficult to analyse by conventional parametric or non-parametric tests. Thus, we used general linear mixed models, employing restricted maximum-likelihood estimation (McCulloch & Searle 2001), to analyse the data, given the small number of subjects studied, their repeated sampling and the unbalanced design due to discarding some samples. We used the Shapiro–Wilk test to determine the normality of response variables: percentages of FLM, SLM, NLM and IMM collected
same male and female (n = 1–3 per menstrual phase per coagulum incidence due to discarding some samples) were considered pure random effects. We used a Bonferroni procedure in post hoc paired comparisons between menstrual phases when contrasting significant main effects and interactions. We did multiple regressions followed by analysis of residuals to evaluate the relationship between percentages of intra-uterine collected FLM, SLM, NLM and IMM (averaged over times of collection) with vaginal pH and temperature measures, irrespective of menstrual phase. Significance in all the cases was set at P ≤ 0.05. We used SPSS 15 (SPSS Inc., Chicago, IL, USA) to perform all the analyses.

Declaration of interest
We declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

Funding
This work was supported by the Instituto Nacional de Psiquiatría Ramón de la Fuente Muñiz (Research Grant # 3320-A).

Acknowledgements
The authors thank the two anonymous reviewers for their helpful suggestions, and Mrs Sally Packard for correcting and improving the English.

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Received 25 March 2008
First decision 29 April 2008
Revised manuscript received 19 June 2008
Accepted 22 July 2008