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The presence of a high- $K_{\rm m}$ hexokinase activity in dog, but not in boar, sperm

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13 Abstract The presence of a high- K_m hexokinase activity was 14 tested in both dog and boar spermatozoa. Hexokinase kinetics 15 from dog extracts showed the presence of a specific activity (dog-16 sperm glucokinase-like protein, DSGLP), in the range of glucose 17 concentrations of 4-10 mM, whereas boar sperm did not show any DSGLP activity. Furthermore, dog-sperm cells, but not 18 19 those of boar, showed the presence of a protein which specifically 20 reacted against a rat-liver anti-glucokinase antibody. This 21 protein also had a molecular weight equal to that observed in 22 rat-liver extracts, suggesting a close similarity between both the 23 24 proteins. This glucokinase-like protein was distributed in the peri- and post-acrosomal zones of the head, and the midpiece and 25 principal piece of tail of dog spermatozoa. These results indicate 26 that dog spermatozoa have functional high- $K_{\rm m}$ hexokinase 27 activity, which could contribute to a very fine regulation of their 28 hexose metabolism. This strict regulation could ultimately be 29 very important in optimizing dog-sperm function along its lifetime.

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Keywords: High-K_m hexokinase; Dog-sperm; Glucokinase-regulatory protein

1. Introduction -36

37 Carbohydrates are, probably, the most important substrates 38 for the maintenance of energy levels in mammalian sperm from 39 fresh ejaculates. Thus, sugars such as glucose, fructose and 40 mannose are utilized as energy sources by sperm cells from 41 species like bull, dog and boar [1]. However, recent findings 42 support the hypothesis that sugars can play another role, at 43 least in some mammalian species, and not only as mere energy 44 substrates. This is especially evident in dog, where the incu-45 bation of sperm cells from fresh ejaculates with either glucose 46 or fructose induced hexose-specific changes in functional pa-47 rameters such as motility [2] or tyrosine phosphorylation

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48 patterns [3]. These glucose- or fructose-specific effects were related to specific actions on the majority of the evaluated 49 metabolic parameters, such as intracellular levels of glucose 6-50 phosphate and glycogen or production of L-lactate and CO₂ 51 [3]. Glucose and fructose also showed separate effects on 52 hexokinase activity [3], and they even induced separate effects, 53 not only on glycogen synthase activity [4], but also on the in-54 55 tracellular, specific location of this enzyme in dog sperm [5]. All of these results indicate that dog sperm would have very 56 57 sophisticated mechanisms to specifically identify sugars that 58 they are consuming in order to direct them to induce the sugar-59 linked, functional effects. These mechanisms might be related 60 to the intake of sugars, since the fructose-specific transporter, GLUT5, and the more glucose-specific transporter, GLUT3, 61 are located in separate zones, not only in dog sperm [3], but 62 also in other mammalian species, such as bull, mice and human 63 [6]. However, there are probably other systems that allow 64 sperm to optimize these hexose-differentiating mechanisms, at 65 least in dog. 66

Vertebrate glucokinase (hexokinase type IV) is a member of 67 the hexokinase protein family which shows some remarkable 68 69 characteristics that clearly differentiate it from the other mammalian hexokinases. In fact, glucokinase does not have a 70 71 strict specificity for substrate, since it can phosphorylate not 72 only glucose, but also fructose or mannose [7]. Nevertheless, 73 glucokinase's elevated $K_{\rm m}$ for glucose, together with its specific 74 expression in the liver and pancreas, allows it to be a sensitive and efficient control step for the maintenance of mammalian 75 glucose metabolism [7]. The existence of a similar, high- $K_{\rm m}$ 76 hexokinase activity in mammalian sperm could be an efficient 77 78 system to control that described above, i.e., hexose-specific 79 functional changes observed, at least in dog. Taking this all 80 into consideration, the main aim of this work is to test the presence of a high- $K_{\rm m}$ hexokinase activity in mammalian 81 sperm, which could act similarly to hepatic glucokinase in the 82 control of sperm's hexose metabolism. For this purpose, sperm 83 cells from dog and boar were used, since they are species which 84 85 show very different functional characteristics, from their mo-86 tion parameters (dog cells are fast and linear, whereas boar 87 cells are much slower, see [2,8]) to their life-span after ejaculation (dog spermatozoa last about one week inside the female 88 genital tract, whereas boar cells last only about 48 h, see [9]). In 89 these cells the total hexokinase activity kinetics was deter-90

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Abbreviations: DSGLP, dog-sperm glucokinase-like protein; PBS, phosphate-buffered saline; SDS, sodium dodecyl sulfate; TRITC, tetramethylrhodamine isothiocyanate

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91 mined, and then the presence of proteins with immunological

92 properties like rat-liver glucokinase and glucokinase regula-93 tory protein was tested, in order to find some glucokinase-like,

94 or at least some high- $K_{\rm m}$ hexokinase, activity in the cells. Our

95 results indicate that dog spermatozoa, but not those of boar,

96 showed a clear high- $K_{\rm m}$ hexokinase activity, as well as a pro-

97 tein which specifically reacts to an anti-rat-liver glucokinase

98 antibody. This high- $K_{\rm m}$ hexokinase activity could be related to

99 the dog-sperm's ability to specificially react in front of glucose

100 or fructose.

101 2. Materials and methods

103 2.1. Animals and sample collection

104 Canine semen was obtained from 11 purebred Beagle dogs ranging 105 from 2 to 7 years of age. The care of the dogs followed the guidelines 106indicated in the Catalan Animal Welfare Law (Generalitat de Ca-107 talunya, Spain). Semen was collected once or twice weekly by manual 108 stimulation. Only the sperm-rich fraction of the ejaculates was used. 109 Porcine semen was obtained from 10 healthy boars ranging from 2 to 110 5 years of age that belonged to a commercial herd (Servicios Genéticos 111 Porcinos, S.L.; Roda de Ter, Spain). Ejaculates were manually col-112 lected and only the rich-sperm fractions were used.

113 2.2. Total hexokinase activity kinetics

114 The kinetics of the total hexokinase activity in dog and boar sper-115 matozoa was determined as in [3]. For this, both boar and dog samples 116 were centrifuged at $600 \times g$ for 10 min and were then resuspended in 50 117 mL of a Krebs-Ringer-Henseleit solution without sugars at 15 °C 118 (KRH- medium; pH 7.4). Sperm were again centrifuged at $600 \times g$ for 119 10 min and sedimented cells were then resuspended in an additional 50 120 mL of the KRH- medium. The centrifugation-resuspension step was 121 repeated once more, in order to completely eliminate all substances 122 included in seminal plasma that could affect kinetic and immunological 123 properties of sperm hexokinases. The final, 50-mL-sperm suspension 124 was again centrifuged at $600 \times g$ for 10 min. After this, the supernatant 125 was discarded and the resultant pellets were immediately homogenized 126 by sonication in 250 µL of an ice-cold buffer (pH 7.4) containing 500 127 mM glycilglycine, 2 M KCl, 100 mM dithiothreitol, 300 IU/mL 128 aprotinin and 100 mM phenylmethylsulfonyl fluoride (hexokinase 129 buffer). Homogenized samples were centrifuged at $10\,000 \times g$ for 15 130 min at 4 °C and hexokinase activity was measured both in the resultant 131 supernatants and in the pellets. For this purpose, the pellets were 132 washed once in 500 µL of hexokinase buffer and were further resus-133 pended in 250 µL of hexokinase buffer. Hexokinase activity was 134 measured as in [10] with the addition of increasing concentrations of 135 glucose in the reaction buffer after adaptation of the technique to a 136 Cobas Bio autoanalyzer (Roche Biomedical, Basel, Switzerland).

137 2.3. Immunological techniques

138 For this experiment, semen samples were pooled both from two 139 (boar semen) or four ejaculates (dog semen). Samples were initially 140 treated through the three times centrifugation/KRH- resuspension 141 washing step described above. After this, both dog and boar sperm 142 cells were suspended in a final 5-mL KRH- medium at 37 °C. Aliquots 143 of the suspension were placed in open vials and incubated with con-144 tinuous shaking at 37 °C, with the addition to the medium of either 145 glucose or fructose at a final concentration of 10 mM in both cases. 146 Concentrations of sperm cells in the final suspension were of 3.5-147 4×10^5 spermatozoa/mL in dog samples and $3-6 \times 10^6$ spermatozoa/ 148 mL in those of boar. Finally, aliquots were taken at the indicated times 149 and processed for immunological detection.

Western blot analyses were performed in samples homogenized by sonication in ice-cold 10-mM Tris–HCl buffer (pH 7.4) containing 1% (w/v) sodium dodecyl sulfate (SDS) and 1 mM Na₂VO₄ (homogenization buffer, proportion 1:5, v/v). The samples were briefly boiled and were then centrifuged at $10\,000 \times g$ for 14 min at 4 °C. Mammalian sperm has very low amounts of cytoplasm and a very compartmentalized structure [11]. These particularities led us to consider the presence of the J.M. Fernández-Novell et al. | FEBS Letters xxx (2004) xxx-xxx

tested proteins in either the supernatant or the pellet obtained after homogenization, centrifugation and the boiling of the samples, since proteins could be in a free form or they could be linked to the internal sperm structures. For this purpose, the obtained pellets were resuspended in 20–30 μ L of the homogenization buffer, and Western blot analyses were performed in both supernatants and resuspended pellets obtained in all of this process.

Western blot was based on SDS gel electrophoresis [12], followed by transfer to nitrocellulose [13]. The transferred samples were tested with the antibodies at a dilution of 1:1000 (v/v). Immunoreactive proteins were tested using peroxidase-conjugated goat, anti-rabbit second antibody (dilution 1:200, v/v) and the reaction was developed with an ECL-Plus detection system (Amersham, Buckinghamshire, England).

Immunocytochemistry was carried out with spermatozoa seeded onto glass coverslips, which were washed with phosphate-buffered saline (PBS; pH 7.4) and were fixed for 30 min in PBS containing 4% (w/v) paraformaldehyde. The fixed samples were incubated with 1 mg/mL NaBH₄ to eliminate autofluorescence, and blocked in 3% (w/v) bovine serum albumin in PBS. Spermatozoa were further incubated with the anti-glucokinase antibody (dilution 1:200, v/v) for 2 h at 15–17 °C, washed with PBS and treated with a tetramethylrhodamine isothiocyanate (TRITC)-conjugated swine anti-rabbit immunoglobulin (Dako, Glostrup, Denmark). Finally, fluorescent images were obtained by a Leica TCS 4D confocal scanning laser microscope (Leica Lasertechnik, Heidelberg, Germany), adapted to an inverted Leitz DMIRBE microscope and a $63 \times$ (NA 1.4 oil) Leitz Plan-Apo Lens (Leitz, Stuttgart, Germany). The light source was an argon/krypton laser (75 mW).

2.4. Suppliers

Anti-rat-liver glucokinase and anti-rat-liver glucokinase regulatory protein were produced and tested in the laboratory of Dr. Guinovart (IRBB, Barcelona Science Park, University of Barcelona; see [14]). All of the reactives used were of analytical grade.

3. Results

3.1. Kinetics of dog and boar total hexokinase activity

191 Supernatants obtained from homogenates of dog sperm 192 showed an increase in total hexokinase activity when it was 193 determined in the presence of glucose in a range from 0.05 mM $(1.4 \pm 0.1 \text{ IU/mg protein})$ to 2 mM $(3.7 \pm 0.4 \text{ IU/mg protein})$ 194 see Fig. 1A). A further, and noticeable increase of total 195 hexokinase activity was then determined in a range of glucose 196 concentration from 4 mM (4.2 ± 0.4 IU/mg protein) to 10 mM 197 198 $(8.4 \pm 0.8 \text{ IU/mg protein}, \text{ see Fig. 1A})$. No further increase in 199 total hexokinase activity was detected at glucose concentra-200 tions above 10 mM. On the other hand, total hexokinase activity from pellets obtained after homogenization of dog sperm 201 from fresh ejaculates also showed an increase in total hexo-202kinase activity in the glucose concentration range from 0.05 203 mM (0.60.1 IU/mg protein) to 4 mM (4.0 ± 0.2 IU/mg pro-204 205 tein). Again, a further and noticeable increase of total hexo-206 kinase activity was observed between 4 mM glucose (4.0 ± 0.2) 207IU/mg protein) and 10 mM glucose (6.2 ± 0.3 IU/mg protein, see Fig. 1B). These results were reflected in the Lineweaver-208Burke representation. Thus, as shown in Fig. 1C, hexokinase 209 210 kinetics of supernatants from dog-sperm extracts described a 211 biphasic diagram, with two separate lines, the first in the glucose range from 0.05 to 6 mM, and the second in the glucose 212 range from 6 to 50 mM. A theoretical, approximate calculus of 213 the $K_{\rm m}$ of both lines resulted in values of 8.5 and 0.08 mM, 214 assuming that these values are only approximate. These results 215 were compatible with the presence of a glucokinase-like ac-216 tivity in dog-sperm supernatants. On the other hand, the Li-217 neweaver-Burke representation of pellets from dog-sperm 218 219 homogenates also showed the presence of two separate lines

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Hexokinase activity (IU/mg Protein)

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8

6

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0

0.05

0.5

1

2





Fig. 1. Kinetics of total hexokinase activity in dog- and boar-sperm extracts. (A and B) Relationship between total hexokinase activity and glucose concentration in supernatants (A) and resuspended pellets (B) from dog (\blacklozenge) and boar (\blacksquare) sperm extracts. (C and D) Semi-logarithmic representation between inverse values of hexokinase-specific activity and glucose concentration in supernatants (A) and resuspended pellets (B) from dog (\blacklozenge) and boar (
) sperm extracts. Lines show the apparent hexokinase activity types from dog (continuous lines) and boar samples (broken lines) revealed by this representation. Results are means ± S.E.M. for eight separate semen samples

220 with very different slopes, the first in the glucose range from 221 0.05 to 6 mM, and the second once again from 6 to 50 mM 222 (Fig. 1D). In this case, the approximate, theoretical values of 223 the $K_{\rm m}$ of both lines were of 6.9 and 0.09 mM.

224 Total hexokinase kinetics of boar sperm showed totally 225 different results. Thus, supernatants from homogenates of 226 these cells did not show an appreciable increase, from 0.05 mM 227 glucose (1.9 \pm 0.2 IU/mg protein) to 50 mM glucose (2.0 \pm 0.3 228 IU/mg protein, see Fig. 1A), thus indicating the presence of 229 only one hexokinase activity that was very sensitive to the 230 presence of glucose in the medium. Slightly different results 231 were observed in pellets from these homogenates, since in this 232 case there was a progressive increase in hexokinase activity in 233 the glucose range from 0.05 mM (3.9 ± 0.7 IU/mg protein) to 234 0.5 mM ($4.9 \pm 0.8 \text{ mM}$, see Fig. 1B), and further increases of 235 hexokinase activity were not noticeable (7.2 ± 0.7 IU/mg pro-236 tein in the presence of 50 mM glucose). These results were 237 reflected in the Lineweaver-Burke representation, where both 238 supernatants and pellets from boar sperm showed only one line from 0.05 mM glucose to 50 mM glucose (Figs. 1C and D). 239 240 The calculated, approximate $K_{\rm m}$ value of this line was of 0.03

mM in supernatants and 0.06 mM in resuspended pellets. 241

242 3.2. Presence of an immunoreactive protein against 243 anti-glucokinase antibody

244 The Western blot using an anti-rat-liver glucokinase anti-245 body revealed the presence of a protein which specifically reacted against this antibody in dog sperm (Fig. 2). This protein 246 was clear in the pellets, with a molecular weight of about 50 247 248 kDa (Fig. 2A). On the other hand, the Western blot from supernatants showed two fainter bands of a molecular weight 249 of about 45-50 kDa (Fig. 2B) The intensity and the molecular 250 251 weight of these bands were not modified after incubation in the



Fig. 2. Western blot against sperm glucokinase. The figure shows two representative Western blots obtained from supernatants (A) and resuspended pellets (B) from dog and boar (P) sperm homogenates. Dog sperm was analyzed from fresh ejaculates (D0) or subjected to incubation with 10 mM glucose for 10 min (DG) before being subjected to Western blot analysis. M: Molecular weight markers. L: Rat-liver extracts. The total number of independent replicates for these Western blots was 5.

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3.3. Detection of sperm glucokinase regulatory protein

252 presence of 10 mM glucose after up to 30 min (Fig. 2, and data 253 not shown). It is noteworthy that liver extracts used as positive 254 controls showed a positive band for glucokinase of about 50 255 kDa, which match to those observed in both supernatants and 256 pellets from dog-sperm homogenates (Fig. 2). Finally, boar homogenates did not show any 50-kDa band which could 257 258 correspond to the result observed in dog extracts, although 259 they revealed a positive signal of about 45 kDa, similar to 260 another non-specific reactivity band detected in liver extracts 261 (Fig. 2).

262 Confocal images showed the presence of a specific, reactive 263 protein against the anti-rat-liver glucokinase antibody in both 264 the head and the tail of dog spermatozoa from fresh ejaculates 265 (Fig. 3). Tail marking was located at both the midpiece and the 266 main piece, whereas head location was established in both the 267 peri-acrosomal and the post-acrosomal zones. These locations 268 were not significantly modified after incubation in the presence 269 of 10 mM glucose after up to 30 min (Fig. 3C, and data not 270 shown). No positive reaction in front of the anti-rat-liver 271 glucokinase antibody was observed in boar spermatozoa (data 272 not shown).

274 Western blot from dog-sperm extracts did not demonstrate the presence of a specific reactivity against an anti-rat-liver 275 glucokinase regulatory protein antibody, neither in superna-276tants nor in resuspended pellets obtained after homogenization 277 of samples (Fig. 4, and data not shown). On the other hand, 278 279 supernatants, but not resuspended pellets, obtained after homogenization of boar spermatozoa showed a clear, specific 70-280 kDa protein, which was equal to that obtained in rat-liver 281 extracts (Fig. 4A, and data not shown). Molecular weight and 282 density of this band in boar sperm were not modified after 283 incubation with 10 mM glucose and 10 mM fructose after up 284 to 30 min (data not shown). 285

4. Discussion and conclusions

Our results indicate the presence of a high- K_m hexokinase 287 activity with a similarity to glucokinase in dog sperm, but not 288 in boar cells. This can be sustained by the following facts: 289



Fig. 3. Immunocytochemistry of dog-sperm glucokinase. The figure shows representative images of dog spermatozoa immunolocalized against glucokinase. A: Negative control. B: Cells from fresh, control samples. C: Cells from samples incubated with 10 mM glucose for 5 min. The total number of independent replicates for these experiments was 5. Images have a multiplication factor of 75. Arrows indicate sperm tails, whereas asterisks indicate sperm heads.

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Fig. 4. Western blot against sperm glucokinase regulatory protein. The figure shows a representative Western blot obtained from supernatants obtained after homogenization of either dog or boar (P) sperm homogenates, following the technique described in Section 2. Dog sperm was from fresh ejaculates (D0) or it was subjected to incubation with 10 mM glucose for 10 min (DG) before being subjected to Western blot analysis. M: Molecular weight markers. L: Rat-liver extracts. The total number of independent replicates for these Western blots was 5.

290 (i) The presence, in dog spermatozoa, of a specific hexokinase 291 activity which shows a high $K_{\rm m}$ for glucose, which was in 292 the range described for mammalian glucokinase (5–10 293 mM, see [7,10]).

(ii) The presence of a dog-sperm protein which specifically reacts against an anti-rat-liver glucokinase antibody and,
moreover, shows a molecular weight similar to mammalian glucokinase (about 50 kDa, see [7]).

298 Thus, immunological and kinetic data point towards this 299 interpretation. Of course, we cannot affirm that this protein is 300 truly glucokinase. Moreover, other proteins different from 301 glucokinase have been described with high- $K_{\rm m}$ characteristics, 302 such as N-acetylglucosamine kinase [15] or the high- K_m 303 hexokinase present in the mhAT3F hepatoma cell line 304 (mhAT3F-HK, see [16]). However, kinetic and immunological 305 characteristics indicate that the dog-sperm, high-Km hexoki-306 nase (DSGLP) is more similar to glucokinase than these pro-307 teins. Thus, N-acetylglucosamine kinase shows a $K_{\rm m}$ for 308 glucose (370 mM, see [15]) far higher than that of DSGLP (7-9 309 mM, see Section 3). Furthermore, the comparison of the 310 structural sequences between glucokinase (reference number 311 NP-034422.2, see [17]) and N-acetylglucosamine kinase (ref-312 erence number Q9QZ08, see [17]) revealed a percentage of 313 overall structural affinity of 11.6%. On the other hand, the 314 structural comparison between the epitope utilized to develop 315 the anti-glucokinase antibody [14] and the whole sequence of 316 the N-acetylglucosamine kinase rendered the following result: 317 Epytope of liver glucokinase:

318 414-KLHPSFKERFASVR-428

319 Sequence of *N*-acetylglucosamine kinase which rendered the 320 higher homology:

321 274-KSWELLKEGFLLALT-288

322 As shown in this comparison, in the best case only four am-323 inoacids, which were not linear (highlighted letters), are the same 324 in both sequences. These data seem to indicate that our antibody 325 has a low degree of affinity for N-acetylglucosamine kinase. 326 Furthermore, the theoretical molecular weight of N-acetylglu-327 cosamine kinase, calculated from its aminoacidic composition 328 [17], is about 37-38 kDa. Our Western blot analysis did not 329 detect any band with could be included in the 35-40 kDa mo-330 lecular-weight range. Summing up all of these data, we can 331 conclude that N-acetylglucosamine kinase does not seem to be 332 DSGLP. On the other hand, mhAT3F-HK also shows a K_m for 333 glucose (40 mM, see [16]) higher than DSGLP. Moreover, al-334 though this protein recognized an anti-glucokinase antibody, its 335 molecular weight was about 30 kDa [16]. Our results did not

reveal the presence of any protein with this molecular weight. 336 The sum of all of these results, of course, does not preclude the 337 presence of *N*-acetylglucosamine kinase and mhAT3F-HK in 338 dog sperm. However, our results also indicate that DSGLP with 339 immunological properties similar to rat-liver glucokinase, regardless of the existence of other high- K_m hexokinases. 341

342 Nevertheless, some differences between glucokinase and DSGLP exist. Thus, glucokinase activity is mainly regulated in 343 two ways, first through controlling its expression [7]. However, 344 mammalian sperm does not have the ability to express its 345 DNA to synthesize proteins [11], so, in this manner, dog sperm 346 cannot control DSGLP activity through this way. The second 347 way is through changes in its spatial position inside the cell 348 [14]. This latter control is achieved through a glucose-modu-349 lated linking between glucokinase and the glucokinase-regu-350 latory protein, which displaces glucokinase in the hepatocyte 351 to areas with or without glucose, thus controlling its ability to 352 phosphorylate the monosaccharide [14]. On the other hand, 353 displacements of glucokinase-regulatory protein through the 354 355 hepatocyte to achieve its control of glucokinase activity are 356 possible due to the hepatocyte being a cell with an active and more-or-less relaxed nuclear structure as well as with a great 357 amount of cytoplasm. Spermatozoa have neither requirement, 358 since their amount of cytoplasm is very scarce and their nu-359 clear structure is condensed and very inactive, totally different 360 from that observed in hepatocytes [18]. Moreover, the presence 361 of a clear immunoreactivity of glucokinase-regulatory protein 362 in boar sperm, which did not show DSGLP activity, suggests 363 the lack of a clear regulatory role for the glucokinase-regula-364 tory protein in dog sperm. Thus, DSGLP activity regulation 365 366 would have probably been achieved by other ways, like phospho-dephosphorylation mechanisms that are also opera-367 tive on glucokinase [19]. 368

The functional role of DSGLP could be explained as a regu-369 370 latory mechanism of dog-sperm hexose metabolism. Previous 371 reports have shown that dog-sperm cells have a very complex hexose metabolism, which includes the presence of anabolic 372 paths like glycogen metabolism, and differentiated, functional 373 roles for separate hexoses [2,4,5]. The elevated $K_{\rm m}$ for glucose 374 shown by DSGLP could be an efficient system to control glu-375 376 cose-specific mechanisms of regulation of dog-sperm function-377 ality, in a similar way that glucokinase controls glucose-induced 378 changes in hepatocytes and pancreatic β -cells [7,10].

379 It is noteworthy that DSGLP is present in dog spermatozoa, but not in boar cells. This would mark great differences in 380 hexose metabolism and, hence, in the energy status manage-381 ment of both species. In fact, hexose metabolism of sperma-382 383 tozoa from several mammals like boar or bull are basically 384 glycolytic, with elevated L-lactate formation rates, low glucose 385 6-phosphate levels, low activity in anabolic pathways like 386 glycogen synthesis and small differences, if any, in the selective utilization of hexoses such as glucose or fructose as energy 387 substrates [20,21]. In these spermatozoa, the presence of 388 DSGLP makes no sense, since they utilize different monosac-389 390 charides in the same way for the same ultimate necessity, the 391 attainment of energy. Nevertheless, as commented above, en-392 ergy management of dog spermatozoa is very different, and the 393 selective, functional utilization that these cells have of glucose and fructose can easily explain the necessity of DSGLP. Thus, 394 395 our results indicate that there is not a single, simple mechanism that explains the regulation of mammalian sperm function 396 397 under the energetic point of view. This has to be taken into
398 consideration when trying to apply new strategies in the con399 servation of sperm, since it would vary depending on the
400 specific metabolic and hexose-related functional profiles of

401 each species.

402 5. Conclusions

403 The presence of DSGLP in dog spermatozoa would play an 404 important role in the control of both the energy management 405 pathways and the hexose-related functional mechanisms that 406 dog spermatozoa show from fresh ejaculates. Notwithstand-407 ing, at this moment we have no real knowledge of the exact 408 physiological role and control mechanisms of DSGLP, and 409 experiments regarding substrate specificity, besides purification, sequencing and cloning of DSGLP, will be needed to 410 411 clarify these important points.

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