Evidence for Compromised Metabolic Function and Limited Glucose Uptake in Spermatozoa from the Teratospermic Domestic Cat (*Felis catus*) and Cheetah (*Acinonyx jubatus*)

Kimberly A. Terrell, David E. Wildt, Nicola M. Anthony, Barry D. Bavister, Stanley P. Leibo, Linda M. Penfold, Laurie L. Marker, and Adrienne E. Crosier

Center for Species Survival, Smithsonian Conservation Biology Institute, Front Royal, Virginia
Department of Biological Sciences, University of New Orleans, New Orleans, Louisiana
University of Puerto Rico, Medical Sciences Campus, San Juan, Puerto Rico
White Oak Conservation Center, Yulee, Florida
Cheetah Conservation Fund, Otjiwarongo, Namibia

ABSTRACT

Cheetahs and certain other felids consistently ejaculate high proportions (>60%) of malformed spermatozoa, a condition known as teratospermia, which is prevalent in humans. Even seemingly normal spermatozoa from domestic cat teratospermic ejaculates have reduced fertilizing capacity. To understand the role of sperm metabolism in this phenomenon, we conducted a comparative study in the normospermic domestic cat versus the teratospermic cat and cheetah with the general hypothesis that sperm metabolic function is impaired in males producing predominantly pleiomorphic spermatozoa. Washed ejaculates were incubated in chemically defined medium containing glucose and pyruvate. Uptake of glucose and pyruvate and production of lactate were assessed using enzyme-linked fluorescence assays. Spermatozoa from domestic cats and cheetahs exhibited similar metabolic profiles, with minimal glucose metabolism and approximately equimolar rates of pyruvate uptake and lactate production. Compared to normospermic counterparts, pyruvate and lactate metabolism were reduced in teratospermic cat and cheetah ejaculates, even when controlling for sperm motility. Rates of pyruvate and lactate (but not glucose) metabolism were correlated positively with sperm motility, acrosomal integrity, and normal morphology. Collectively, our findings reveal that pyruvate uptake and lactate production are reliable, quantitative indicators of sperm quality in these two felid species and that metabolic function is impaired in teratospermic ejaculates. Furthermore, patterns of substrate utilization are conserved between these species, including the unexpected lack of exogenous glucose metabolism. Because glycolysis is required to support sperm motility and capacitation in certain other mammals (including dogs), the activity of this pathway in felid spermatozoa is a target for future investigation.

INTRODUCTION

An interesting trait of certain felid species and genotypes is the production of unusually high proportions of sperm malformedations. Species, populations, or individuals that express this condition are called teratospermic [1]. This phenomenon is especially common in species or subspecies that have low levels of gene diversity (cheetah [2–4], Florida panther [5, 6], and Asian lion [7]) and in domestic cats that have been purposefully inbred [8]. Teratospermia (defined here as the production of >60% structurally abnormal spermatozoa) also is common among men. A recent meta-analysis of semen characteristics by the World Health Organization revealed that >95% of men can be classified as teratospermic under this definition [9]. Our laboratory has used certain felid species and genetic lineages to better understand the impact and etiology of teratospermia. Various studies have revealed that spermatozoa from teratospermic ejaculates demonstrate delayed capacitation [10], compromised acrosomal function [10], disrupted protein tyrosine phosphorylation [11, 12], increased osmotic sensitivity [13, 14], reduced zona penetration ability [15], and increased sensitivity to cooling [16] and cryopreservation [17]. These mechanisms no doubt contribute to the reduced fertilizing ability of teratospermic ejaculates in vitro, even after processing to isolate structurally normal spermatozoa for insemination [15].

Some of these physiological impairments (e.g., tyrosine phosphorylation) could be related to a diminished capacity for energy production in malformed spermatozoa, but there is currently no knowledge of gamete metabolism in felids. Studies of mammalian sperm energy production, although conducted since the 1940s, have generally been confined to humans and fewer than 10 domesticated species [18]. Yet as Storey detailed in a recent review [18], there are considerable differences in metabolic function of male gametes, even within this small group of species. It is known that spermatozoa are capable of generating energy in the form of ATP through glycolysis and/or oxidative phosphorylation. However, the relative importance of each pathway to sperm functions, such as motility and capacitation, varies among species [18–25]. Oxidative phosphorylation is 18 times more efficient than anaerobic glycolysis and provides a significant proportion of the ATP supply in spermatozoa of most species [21].

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2Correspondence: Adrienne E. Crosier, Smithsonian Conservation Biology Institute, Center for Species Survival, 1500 Remount Rd., Front Royal, VA 22630. FAX: 540 635 6506; e-mail: crosiera@si.edu
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notable exception is the human, whose sperm appear to rely entirely on glycolysis for motility and hyperactivation [26]. Despite the efficiency of oxidative metabolism, its ability to fulfill energy demands in the distal flagellum is questionable [23, 24, 27], as mitochondria are confined to the sperm midpiece. Therefore, glycolysis may be an important supplemental source of ATP to fuel sperm motility, and glycolytic enzymes have been localized along the fibrous sheath of the flagellum in the boar, bull, rat, stallion, human, and mouse [28, 29]. Sperm production of lactate (presumably by glycolysis) is correlated positively with motility, normal morphology, acrosomal integrity, and osmotic resistance in the boar and donkey [30, 31]. One of the latter studies has suggested that these relationships are more than casual in that litter size in pigs is enhanced after artificial insemination using sperm producing high lactate concentrations [30].

In contrast to livestock species, there is a lack of information on sperm metabolism in carnivores. Gamete metabolism has been fairly well studied in the domestic dog [19, 32–39], but, to our knowledge, these pathways have not been investigated in any other carnivore species. The dog apparently is uniquely capable of sperm gluconeogenesis [19], which is surprising given that glucose synthesis requires three times more ATP than is produced by glycolysis [40]. Utilization of this pathway may explain how dog spermatozoa are able to maintain motility and achieve capacitation in a medium without glucose [19, 38]. This is especially interesting as glucose is required for capacitation in the mouse [41] and human [26], but inhibits sperm motility in the sheep [42].

Felids are attractive models for studying gamete metabolism. The availability of multiple species within the family Felidae provides opportunities for comparative studies to understand the conservation (or diversification) of physiological processes. Furthermore, the existence of teratospermia in certain species or genetic lineages provides the opportunity to explore linkage between a complex biological phenomenon and potential causative factors. Our aim in this study was to determine the relationship between rates of glycolytic and oxidative sperm metabolism and conventional indices of cellular function (i.e., structural morphology, motility, and acrosomal integrity). The approach was unique because we took advantage of two domestic cat populations that consistently produce differing proportions of pleiomorphic spermatozoa. To increase the robustness of our findings, we conducted a cross-species comparison using the cheetah, a species that is well known to be teratospermic regardless of season or living conditions (free-range versus captive) [2–4]. Our hypotheses were that 1) metabolic rates are useful indicators of sperm quality in felids and 2) metabolic function is compromised in spermatozoa from teratospermic ejaculates compared to normospermic counterparts. We expected that elucidating the pathways of felid sperm energy production not only would provide insight into the physiological basis of teratospermia, but also might yield a reliable, quantitative indicator of ejaculate quality. The latter information has potential applied benefits. For example, identifying metabolic substrate requirements would be highly informative for enhancing the use of certain types of assisted reproductive technologies for genetically managing wild felid species [44] or domestic cat lineages studied as models of human disorders [45].

MATERIALS AND METHODS

Animals

Electroejaculates were collected from adult domestic cats (ages 1.5–8 yr) that were known to consistently produce either normospermic (>60% normal sperm per ejaculate, n = 3 males) or teratospermic (<40% normal sperm per ejaculate, n = 3 males) ejaculates. In all, 15 ejaculates were collected from normospermic males (three to eight per individual) and 19 ejaculates from teratospermic males (three to 14 per individual). Males were housed individually in 2.7-m² indoor cages at the Smithsonian Conservation Biology Institute (Front Royal, VA), maintained on a 14L:10D cycle, and provided dry, commercial cat food (Purina Cat Chow; Ralston Purina Co., St. Louis, MO) and water ad libitum.

Electroejaculates (one per male, 22 males) were collected from adult cheetahs (ages 2.5–10 yr) housed at the Cheetah Conservation Fund (CCF; Ongava, Namibia: n = 18), White Oak Conservation Center (WOCC; Yulee, FL; n = 3), or the Smithsonian’s National Zoological Park (NZP; Washington, DC; n = 1). Males at CCF were wild-born and housed as described previously [46]. Males at WOCC were wild-born, housed together (in a group of three) in a 2000-m² outdoor enclosure, and fed a commercially produced Nebraska Carnivore diet (Central Nebraska Packaging Inc., North Platte, NE). The single male at NZP was captive-born, housed on exhibit with two other males in a 1400-m² outdoor enclosure, and fed a commercially produced carnivore diet (Carnivore Diet 10; Natural Balance Pet Foods Inc., Pacoima, CA).

Semen Collection and Evaluation

A surgical plane of anesthesia was induced in domestic cats and cheetahs according to protocols determined by institutional veterinarians and similar to those previously used for semen collection in these two species [10, 15, 46]. All animal procedures were approved by NZP’s Animal Care and Use Committee (ACUC) and the WOCC ACUC. Methods for semen collection and evaluation were similar to those described in previous studies [10, 15, 46]. A rectal probe (1 cm (domestic cat) or 1.9 cm (cheetah) in diameter with three longitudinal electrodes and an electrostimulator [Electrostim; OR]) were used to deliver 80 stimuli (at a low voltage of 2–5 V) over a 30-min interval [47]. Ejaculates (n = 56 total) were collected in sterile, prewarmed vials as previously described [4, 47].

The volume of each ejaculate was measured using a pipette, and 3 µl of ejaculate were immediately assessed visually for sperm percentage motility and forward progressive status (FPS or speed of forward progression; scale 0–5, with a rating of 5 equivalent to most rapid, straightforward progress [47]). A sperm motility index (SMI) was calculated using the formula (percent motility + (FPS × 20) ÷ 2 [48]). A 5-µl sample of raw semen was fixed in 0.3% glutaraldehyde in PBS (pH 7.4, 340 mOsm) for subsequent assessment of sperm morphology [13]. For each sample, 100 spermatozoa were assessed (1000× magnification) and classified as normal or having (in order of precedence) a head, acrosomal, midpiece, flagellar, or other abnormality, as previously described [47]. For all ejaculates, a second 5-µl aliquot of raw semen was fixed in 4% paraformaldehyde to evaluate acrosomal integrity (% IA). Fixed samples were processed using a modified Coomassie Blue G-250 (Fisher Biotech, Springfield, NJ) staining technique, as described earlier [49, 50]. Spermatozoa were evaluated (1000× magnification) and classified as having an intact or nonintact acrosome. Spermatozoa with an intact acrosome exhibited a uniform blue staining pattern overlying the acrosomal region, whereas nonintact cells had clear or patchy staining over this region [50].

Sperm Processing and Metabolic Assessments

Each ejaculate was diluted immediately with an equal volume of a chemically defined, protein-free, modified mouse tubal fluid medium (cMTF) [51] supplemented with 2% polyvinyl alcohol (PVA) [52]. The final cMTF medium (pH 7.45) contained 98.4 Mm NaCl, 4.78 mM KCl, 1.19 mM MgSO₄, 1.19 mM KH₂PO₄, 25 mM NaHCO₃, 1.71 mM CaCl₂, 1 mM glucose, 1 mM Na-pyruvate, 25 mM NaCl (N-morpholino) propanesulfonic acid (MOPS) buffer, and 0.02 µg/ml phenol red. All reagents were purchased from Sigma Aldrich (St. Louis, MO). The cMTF medium was prepared fresh daily from five concentrated stock solutions containing 1) NaHCO₃ and phenol red, 2) CaCl₂ 3) glucose and pyruvate, 4) MOPS and phenol red, and 5) all remaining reagents. All stock solutions were kept at 4°C and discarded after 2 wk (stocks 1–3) or 3 mo (stocks 4 and 5). PVA was added, and the medium was sterilized through a 0.22-µm syringe filter immediately prior to use. Osmolality of the final working medium (295–341 mOsm) was determined using a vapor pressure osmometer (Wescor, Inc., Logan, UT) and was within 10% of the physiological value of domestic cat semen (323 mOsm [53]).

Diluted ejaculates (maintained at ambient temperature, 19°–24°C) were washed by centrifugation (8 min; 300 × g for domestic cat, 200 × g for cheetah) and resuspended in fresh cMTF at a concentration of 3 × 10⁶ motile sperm per milliliter. Sperm concentration was determined using a Nucleocounter SP-100 (Chemometec, Allerød, Denmark) [54]. Sperm samples were incubated (37°C) in microcentrifuge tubes under oil (200 µl) to prevent evaporation. Based on
rates of sperm oxygen consumption in the dog and fox [55], we estimated that dissolved oxygen in cat/cheetah sperm samples would decrease by <1% after 24 h. Because sperm respiration is not limited until 90% of oxygen is depleted from the medium (starting at the atmospheric value) [56], hypoxia due to culture under oil was not of concern. A sample (130 μl) of sperm suspension was taken at 0, 1, 3, 7, and 24 h of incubation, and cells were removed by centrifugation (3 min; 1000 × g) through a Costar Spin-X 0.22-μm nylon filter tube (Coming Incorporated, Corning, NY). The filter was removed from the tube, and the sperm-free medium was stored at −80°C until analysis. Acrosomal membrane integrity and sperm motility were assessed at 0, 1, 3, 7, and 24 h as described above and are reported as average values over each time interval to facilitate comparison with metabolic rates.

Samples of sperm-free medium were analyzed for glucose, pyruvate, and lactate concentrations using enzyme-linked fluorescence assays [51, 57]. Each assay was linked to the oxidative status of the coenzyme NADP (glucose) or NAD (pyruvate and lactate). The reduced forms of these coenzymes (NADPH and NADH) fluoresce at 445 nm when excited at 340 nm, whereas the oxidized forms do not. For the glucose assay, sperm-free medium (10 μl) was incubated (5 min, 37°C) with an enzyme cocktail (200 μl) containing 0.42 mM dithiothreitol, 3.1 mM MgSO4, 0.42 mM ATP, 1.25 mM NADP, and 0.1 U/ml hexokinase/glucose-6-phosphate dehydrogenase (HK/G6PDH) in 50 mM EPPS buffer (4-[2-hydroxyethyl]-1-piperazine propane-sulfonic acid; pH 8.0). The cocktail was stored in the dark at −80°C for up to 3 mo prior to use. The conversion of glucose to 6-phosphogluconate was carried out as shown in Equation 1. Glucose concentration determined by this assay was directly proportional to NADPH fluorescence.

\[
\text{HK} \\
\text{glucose} + \text{ATP} \rightarrow \text{glucose-6-phosphate} + \text{ADP} \\
\text{G6PDH} \\
\text{glucose-6-phosphate} + \text{NADP}^+ \rightarrow \text{6-phosphogluconate} + \text{NADPH} + H^+ 
\]

For the pyruvate assay, sperm-free medium was incubated with an enzyme cocktail (as above) containing 0.14 mM NADH and 0.12 U/ml lactate dehydrogenase (LDH) in 50 mM EPPS buffer (pH 8.0). The cocktail was stored in the dark at −80°C for up to 3 mo prior to use. The conversion of pyruvate to lactate was carried out as shown in Equation 2. Pyruvate concentration measured by this assay was inversely proportional to NADH fluorescence.

\[
\text{LDH} \\
\text{pyruvate} + \text{NADH} + H^+ \rightarrow \text{lactate} + \text{NAD}^+ 
\]

For the lactate assay, sperm-free medium (25 μl) was incubated (5 min, 37°C) with an enzyme cocktail (250 μl) containing 1.92 U/ml LDH, 0.2 U/ml glutamate-pyruvate transaminase (GPT), 0.42 mM NAD+, and 100 μM glutamate in 1 M glycine buffer containing 5.6 mM ethylenediaminetetraacetic acid. The buffer was stored at 4°C for up to 1 mo prior to use, and the cocktail was prepared fresh daily using NAD+ and glutamate stock solutions stored at −80°C for up to 3 mo prior to use. This assay is a nontoxic alternative to the LDH/hydrazine assay. The conversion of lactate to alanine was carried out as shown in Equation 3. Lactate concentration in this reaction was directly proportional to NADH fluorescence.

\[
\text{LDH} \\
\text{lactate} + \text{NAD}^+ \rightarrow \text{pyruvate} + \text{NADH} + H^+ \\
\text{GPT} \\
\text{pyruvate} + \text{glutamate} \rightarrow \text{alanine} + \text{NADH} + \text{H}^+ 
\]

Enzymes (LDH, HK/G6PDH, and GPT) were purchased from Roche Applied Science (Indianapolis, IN). Fluorescence was analyzed using a Spectra Max Gemini XS fluorescence plate reader (Molecular Devices, Sunnyvale, CA) and SoftMax Pro 5 software (Molecular Devices, Sunnyvale, CA). Metabolic rates were calculated as the change in substrate concentration over time divided by sperm concentration and are reported in nmol/10^6 sperm/h.

### Statistical Analyses

Data were analyzed with Statistical Analysis Software (SAS) version 9.1 (SAS Institute, Cary, NC), and percentage data were arc sine-transformed before evaluation. Differences in ejaculate characteristics and sperm morphology among animal groups (normospermic domestic cat, teratospermic domestic cat, and cheetah) were assessed using SAS General Linear Model Procedures (GLM) [58]. To evaluate changes in SMI, % IA, and metabolic rate over time, data were analyzed using a separate GLM for each animal group [58]. Within each domestic cat group (normospermic and teratospermic), there was no interaction (P > 0.05) between individual and time, as well as no main effect of individual (P > 0.05) on SMI, % IA, or metabolism; thus, these variables were omitted from the final model. Differences in SMI, % IA, and metabolic rate among animal groups were assessed using a separate GLM for each time interval [58]. To determine if variation in sperm motility was responsible for differences in metabolic rates among animal groups, data from all individuals and time points were combined and analyzed using a GLM, with percent motile spermatozoa included as a covariate. When a significant (P < 0.05) F-statistic was measured in any GLM, differences among means were assessed using the Duncan multiple-range test. The Pearson correlation was used to evaluate the relationships between metabolic rate and sperm morphology, SMI, and % IA within and across animal groups. Results were considered significant at P < 0.05 and are reported as least-squares (LS) means ± SEM unless otherwise stated.

### RESULTS

**Ejaculate and Sperm Characteristics**

Semen volume and sperm concentration were similar (P > 0.05) in normospermic and teratospermic domestic cats (Table 1). Cheetah ejaculates were less concentrated (P < 0.05) than those from domestic cats, but due to larger (P < 0.05) seminal volumes, the total number of spermatozoa per ejaculate did not (P > 0.05) differ among the three animal groups (Table 1). The
average SMI and percentage of structurally normal spermatozoa were similar \((P > 0.05)\) between teratospermic domestic cats and cheetahs, both of which were less \((P < 0.05)\) than in normospermic cats (Table 1). A bent midpiece encompassing a cytoplasmic droplet was the most prevalent deformity observed in each animal group and constituted \(\approx 45\%\) of all abnormalities (Table 1). This was followed by acrosomal abnormalities and proximal droplets, which were more \((P < 0.05)\) common in the cheetah (19\% and 14\% of all deformities, respectively) compared to the domestic cat (8\% and 5\%, respectively). A bent flagellum encircling a cytoplasmic droplet was a less frequent \((P < 0.05)\) deformity in the cheetah (3\%) than in the domestic cat (12\%). In both species, spermatids and midpiece bends (without a droplet) constituted <10\% of all abnormalities. More than a dozen other deformities were observed rarely (<5\%) in each group but collectively comprised a significant proportion (~15\%) of total anomalies. These malformations were classified as “other” and included macro-/microcephaly, bi-/tricephaly, a misshapen head, residual cytoplasm attached to the head, a bent neck, partial or complete midpiece aplasia, a distal midpiece droplet, a misshapen midpiece, a coiled flagellum with or without a droplet, and a bi-/triflagellate spermatozoon. Images of the more prevalent deformities (i.e., constituting \(\geq 10\%\) of abnormalities in any group) are provided in Supplemental Figure S1 (available online at www.biolreprod.org). These depictions, along with most of the uncommon malformations listed above, are also available in earlier publications [2, 4].

All three animal groups ejaculated high percentages of acrosome-intact spermatozoa (>80\% overall; Table 1), although teratospermic cats produced a smaller proportion \((P < 0.05)\) of cells with intact membranes compared to normospermic counterparts or cheetahs.

**FIG. 1.** A) SMI and (B) % IA in normospermic domestic cats (solid bars), teratospermic domestic cats (lined bars), and cheetahs (open bars). Among animal groups and within each time point, bars with different lowercase letters differed \((P < 0.05)\). Error bars represent means ± SEM.

**FIG. 2.** Sperm pyruvate uptake (A), glucose uptake (B), and lactate production (C) in normospermic domestic cats (solid bars), teratospermic domestic cats (lined bars), and cheetahs (open bars). Among animal groups and within each time point, bars with different lowercase letters differed \((P < 0.01)\). Error bars represent means ± SEM.

### Sperm Motility, Acrosomal Integrity, and Metabolism over 24 h

Within each time interval, SMI and % IA were similar \((P > 0.05)\) in teratospermic cats and cheetahs, but reduced \((P < 0.05)\) compared to normospermic cats (Fig. 1). Sperm pyruvate uptake from 0–1 h was higher \((P < 0.05)\) in teratospermic cats compared to cheetahs, and rates in normospermic cats were similar \((P > 0.05)\) to both of these groups (Fig. 2A). However, after 1 h, rates were ~70\% lower \((P < 0.05)\) in teratospermic cats and cheetahs compared to normospermic cats. Due to this inconsistency and large standard error values for the 0– to 1-h interval, rates of pyruvate uptake for this time interval were omitted from the Pearson correlations.

Exogenous glucose was minimally utilized by domestic cat spermatozoa, and rates of uptake were similar \((P > 0.05)\)
Relationship Between Metabolic Rates and Sperm Quality

When data from all ejaculates (n = 56) were combined, rates of pyruvate uptake were positively correlated (P < 0.001) to SMI (r = 0.44) and % IA (r = 0.43; Table 3). Because rates of lactate production were correlated positively (P < 0.001) to normal sperm morphology across all ejaculates, as well as with SMI and % IA (r = 0.42–0.50; Table 3), this metric was found to be a more accurate indicator of overall cellular quality than pyruvate uptake. Furthermore, rates of lactate production were correlated positively (P < 0.05) to SMI and % IA within each group (r = 0.34–0.67). Rates of glucose uptake were not (P > 0.05) correlated with SMI or % IA in any group.

To determine if decreased metabolic rates in spermatozoa of teratospermic cats and cheetahs were an artifact of reduced motility in these cells, data were reanalyzed with sperm motility (% motile and SMI) included as covariates in the GLM. Results were consistent with the previous analysis: sperm pyruvate uptake and subsequent lactate production were decreased (P < 0.05) in teratospermic cats and cheetahs compared to normospermic cats, thereby indicating a direct relationship to the teratospermic condition.

Given that rates of lactate production were correlated positively to normal sperm morphology across all feline ejaculates, we were curious about the metabolic function of cheetah ejaculates containing relatively high proportions (~40%) of structurally normal spermatozoa. Specifically, do these high-quality cheetah ejaculates demonstrate normal metabolic function when compared to domestic cat counterparts? Conversely, is metabolic function more severely compromised in cheetah ejaculates with very low (~5%) proportions of structurally normal spermatozoa? To address these questions, cheetah ejaculates from the existing dataset were ranked in order of increasing percentages of structurally normal sperm (% N), and those with the lowest and highest percentage values were selected for comparison of lactate production (n = 5 per group; Fig. 3). The mean % N in each ejaculate group was 7% ± 3% (range 5%–10%) and 42% ± 3% (range 35%–58%), respectively. Domestic cat ejaculates (n = 9 total, 4 males) having % N values within the 35%–58% range (mean 44% ± 2%) were selected from the existing dataset as a control for normal rates of lactate production. Despite the large difference in % N between the two cheetah groups, rates of sperm lactate production were similar (P > 0.05) and were reduced (P < 0.05) compared to the domestic cat control group (Fig. 3). Pyruvate and glucose metabolism was not reassessed, because rates of uptake were not correlated with any sperm quality index in the cheetah (Table 3).

Discussion

This was the first study of sperm metabolism in any feline species, and we made four significant discoveries. First, we determined that felid spermatozoa (at least from the two species studied here) did not rely on exogenous glucose as a source of energy. Rather, based on the observed ratios of substrate uptake/production, it appeared likely that these cells generated ATP from the catabolism of one or more unidentified endogenous sources. Secondly, certain cellular mechanisms related to sperm energy production were conserved between the domestic cat and cheetah, indicated by similar patterns of substrate metabolism between the species. Third, metabolic

TABLE 2. Substrate uptake/lactate production ratios in domestic cat and cheetah spermatozoa.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Normospermic cat</th>
<th>Teratospermic cat</th>
<th>Cheetah</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of males</td>
<td>3</td>
<td>3</td>
<td>22</td>
</tr>
<tr>
<td>No. of ejaculates</td>
<td>15</td>
<td>19</td>
<td>22</td>
</tr>
<tr>
<td>Glucose</td>
<td>0.2 ± 0.4</td>
<td>0.1 ± 0.4</td>
<td>ND</td>
</tr>
<tr>
<td>Pyruvate</td>
<td>0.6 ± 0.2</td>
<td>0.7 ± 0.2</td>
<td>0.7 ± 0.1</td>
</tr>
</tbody>
</table>

* ND, not detected.

Overall between normospermic and teratospermic males (Fig. 2B). Extracellular glucose concentration of cheetah sperm medium samples did not change (P > 0.05) from 0–24 h, indicating no metabolism of this substrate by this species. Sperm lactate production occurred at comparatively high rates in all three groups, given the expected ratio (2:1) of lactate production to glucose uptake (Fig. 2C). Compared to normospermic cats, rates of sperm lactate production were ~60% less (P < 0.01) in teratospermic cats and ~80% less (P < 0.001) in cheetahs. Extreme variation in stoichiometric ratios (glucose or pyruvate uptake as a proportion of lactate production) was observed within each time interval for all three groups, with most coefficient of variation (CV) values being ≥50% (data not shown). This variation likely was related to minimal (or sometimes zero) changes in metabolic substrate concentration between consecutive time points (even in samples with high percentages of motile spermatozoa). In support of this supposition, normospermic cats demonstrated higher sperm metabolic rates and, overall, less variation in stoichiometric ratios (CV = 12%–392%) compared to the other two groups. To eliminate as much assay “noise” as possible, stoichiometric ratios were recalculated based on the total change in substrate concentration from 0 to 24 h of incubation. Using this method, most (60%–70%) lactate production was attributed to the uptake and reduction of pyruvate, whereas a lesser portion (0%–40%) was credited to the metabolism of exogenous glucose metabolism (Table 2).

DISCUSSION

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TABLE 3. Correlation coefficient (r) values for metabolic rate versus sperm quality in domestic cats and cheetahs.*

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Overall</th>
<th>Normospermic cat</th>
<th>Teratospermic cat</th>
<th>Cheetah</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of males</td>
<td>28</td>
<td>3</td>
<td>3</td>
<td>22</td>
</tr>
<tr>
<td>No. of ejaculates</td>
<td>56</td>
<td>15</td>
<td>19</td>
<td>22</td>
</tr>
<tr>
<td>Pyruvate uptake (nmol/10^6 sperm/h)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sperm motility index</td>
<td>0.44b</td>
<td>0.44a</td>
<td>0.50b</td>
<td>NS</td>
</tr>
<tr>
<td>Intact acrosomes (%)</td>
<td>0.43c</td>
<td>0.41a</td>
<td>0.52b</td>
<td>NS</td>
</tr>
<tr>
<td>Structurally-normal spermatozoa (%)</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>Lactate production (nmol/10^6 sperm/h)</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Sperm motility index</td>
<td>0.45b</td>
<td>0.37a</td>
<td>0.50b</td>
<td>0.62b</td>
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<tr>
<td>Intact acrosomes (%)</td>
<td>0.42b</td>
<td>0.34a</td>
<td>0.51b</td>
<td>0.62b</td>
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<tr>
<td>Structurally-normal spermatozoa (%)</td>
<td>0.50b</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
</tbody>
</table>

* NS, not significant (P > 0.05).
ab P < 0.05; b P < 0.001.
function was impaired in spermatozoa from teratospermic ejaculates, as revealed by relatively low rates of pyruvate uptake and lactate production in males producing high proportions of pleiomorphisms. This observation was consistent with previous reports that linked teratospermia to disruptions in multiple components of sperm function, including several energy-dependent processes [10, 11, 15, 59]. Finally, rates of lactate production were correlated positively to multiple measures of sperm function in both the domestic cat and cheetah. Therefore, this substrate may prove to be a valuable indicator of ejaculate quality.

The lack of glucose uptake by domestic cat and cheetah spermatozoa is unexpected, given that glycolysis is required to support sperm motility [21, 26, 34] and capacitation [19, 26, 60, 61] in the human and domestic dog. This finding perhaps could result from low hexokinase activity in felids compared to other species or, in the case of the cheetah, the complete absence of this enzyme. Reduced hexokinase activity would limit NADPH production, a key component of the glutathione-mediated defense system that protects cellular membranes against lipid peroxidation damage [62]. This is an intriguing possibility, as spermatozoa from teratospermic felids (including the domestic cat and cheetah) are unusually susceptible to membrane damage [13, 16, 46]. However, it also remains possible that felid spermatozoa possess fully functional hexokinase but metabolize glucose at modest rates relative to oxidative substrates. We currently are using the glycolytic inhibitor α-chlorohydrin to more thoroughly understand the role of glycolysis in felid sperm cellular function.

Given the observed lack of glucose uptake by cat and cheetah spermatozoa, we were surprised to discover that these cells consistently produced lactate, which is an end-product of glycolysis [63]. Under our experimental conditions, we consider there to be three possible sources of lactate: (1) endogenous glycogen, (2) endogenous phospholipids, and/or (3) imported pyruvate. These potential mechanisms of lactate production are not mutually exclusive and may also contribute, to varying degrees, in the generation of NADH and ATP (Fig. 4). In each case, lactate formation could occur in cytosol or mitochondria, as sperm-specific lactate dehydrogenase has been found in these compartments [64–67]. Lactate and NADH production also could occur in separate cellular locations, since reducing equivalents can be transferred between the cytosol and mitochondria by the malate-aspartate shuttle (present in spermatozoa of several species [68–70]).

Glycogen is known to be metabolized by spermatozoa of the domestic dog, another carnivore [36]. Intracellular glycogen breakdown would yield ATP, and cytosolic NADH would be

![FIG. 3. Sperm lactate production in cheetah ejaculates selected for either normal (lined bars) or abnormal (open bars) morphology, compared to a domestic cat control group (closed bars). Among animal groups and within each time point, bars with different lowercase letters differed (P < 0.05). Error bars represent means ± SEM.](image-url)

![FIG. 4. Theoretical model showing three possible mechanisms of lactate production by domestic cat and cheetah spermatozoa, with NADH, lactate, and ATP generated from the metabolism of endogenous glycogen (blue), phospholipid (green), and/or extracellular pyruvate (purple).](image-url)
regenerated by lactate production (Fig. 4). However, phospholipid is considered the primary endogenous substrate for most mammalian spermatozoa [21] and could provide greater amounts of cellular ATP. Phospholipid hydrolysis would yield glycerol and fatty acids (Fig. 4). Glycerol would enter the glycolytic pathway via conversion to dihydroxyacetone phosphate and would be metabolized to produce lactate, NADH, and ATP [71]. Mitochondrial oxidation of fatty acids would provide substantial amounts of NADH, which could contribute to ATP and/or lactate production (Fig. 4).

Because we used a protein-free, chemically defined medium, the only possible exogenous source of lactate in our study was the uptake and reduction of extracellular pyruvate. Although pyruvate reduction would require an NADH source and would not generate ATP [63], these molecules could be provided by endogenous substrate metabolism (as described above). Pyruvate uptake likely would occur via the facilitative transporter SLC16A7 (monocarboxylic acid transporter 2, previously known as MCT2), the primary monocarboxylate transporter in mature spermatozoa of species studied to date [72–74]. Given the starting composition of our c-MTF medium (1 mM pyruvate, 0 mM lactate) and the kinetic properties of SLC16A7 and LDH [75–77], rapid pyruvate uptake and reduction theoretically should occur independently of sperm energy production. Examination of stoichiometric ratios revealed that 60%–70% of produced lactate could have been explained on the basis of pyruvate uptake and reduction (in contrast to only 0%–40% generated from exogenous glucose catabolism). These are important observations given that lactate production (measured by enzyme-linked fluorescence) has been used as an indicator of glycolytic metabolism for other species [25, 26, 31]. Ongoing research in our laboratory supports this mechanism of lactate production in felid spermatozoa. Initial findings determined that sperm lactate production was approximately four times greater in the presence of exogenous pyruvate compared to equimolar amounts of exogenous glucose. Furthermore, in the absence of both substrates in the culture medium, no lactate was produced. Collectively, these observations suggested that mammalian spermatozoa may produce lactate independently of glycolysis, implying that the functional importance of this pathway could have been misinterpreted in earlier studies. More detailed studies are in progress in our laboratory using chemical inhibitors of oxidative and glycolytic metabolism to identify the primary energy substrates for felid spermatozoa.

Although the source of lactate was unclear, rates of production provided a consistent indicator of sperm motility and acrosomal function in both felid species. Intriguingly, lactate production also was correlated to proportions of structurally normal spermatozoa among domestic cat (normospermic and teratospermic combined), but not cheetah, ejaculates. This finding may be related to previous reports that even structurally intact cells from teratospermic ejaculates can be functionally compromised. Specifically, these spermatozoa may demonstrate increased osmotic sensitivity, delayed acrosome reaction, or reduced zona penetration ability [10, 11, 15]. Therefore, rates of lactate production may provide an accurate indicator of ejaculate quality in both species, and may reveal disrupted cellular physiology in apparently normal cheetah spermatozoa. However, lactate production should be validated against a more direct measure of sperm fertilizing ability such as the zona pellucida assay, particularly given its unclear relationship to ATP generation. Because spermatozoa must be capable of multiple energy-dependent processes to achieve successful fertilization, a zona penetration assay could provide a more robust test of the functional relevance of lactate production than any single measure of sperm function (e.g., motility). Understanding the source and biological significance of lactate produced by felid spermatozoa could yield a quantitative, field-friendly indicator of ejaculate quality, which would significantly facilitate developing and refining reproductive technologies for improving felid management and conservation.

Because so little is known about gamete metabolism in felids, this taxon is an excellent target for more detailed investigation in this area of study. The existence of >30 species in the family Felidae (including several with the teratospermic phenotype [78]) also provides important opportunities to determine the etiology and evolution of certain physiological attributes influencing reproductive success. Indeed, direct comparisons across species are fundamental to identifying differences in the mechanisms involving energy production in mammalian gametes, particularly given the ambiguities of published findings relating to sperm metabolism. For example, Carey et al. [79] have reported that mouse spermatozoa metabolize endogenous oxidative substrates to remain motile for >4 h in vitro in the absence of supplemental energy sources. Yet Mukai and Okuno [80] have found that mouse spermatozoa become nonmotile within 30 min in a substrate-free medium and have presented evidence for an obligate role of glycolysis in supporting cellular motility. Contradictory findings also have been reported regarding the need for exogenous glucose to achieve capacitation in dog spermatozoa [61, 81]. Such equivocations can be avoided and more confidence generated by comparatively evaluating two taxonomically related species using a standardized experimental protocol. In the present study, we found similar patterns of substrate uptake/production in the domestic cat and cheetah, suggesting that spermatozoa of both species relied on the same energy source(s). Relatively low rates of substrate uptake/production in cheetah ejaculates revealed that sperm metabolic efficiency was compromised in this species. More importantly, comparison with the teratospermic cat model then allowed us to link this finding to ejaculate phenotype. Still, we uncovered a subtle but key physiological difference between these species. Specifically, in contrast to the domestic cat, even apparently “high-quality” cheetah ejaculates (i.e., containing higher proportions of structurally normal spermatozoa) demonstrated compromised metabolic function. Thus, while the collective results confirmed that the teratospermic cat was an excellent model for expanding knowledge about gamete physiology in wild felids, findings revealed important functional differences between these two related species.

In conclusion, study results underscored the importance of species diversity in fundamental reproductive phenomena, as has been emphasized recently in the contexts of biological conservation and human health [82]. We predict that understanding the unique mechanisms of energy production in felid spermatozoa will facilitate increased efficiency of assisted reproductive technologies that have been used for producing offspring in wild felids [83, 84]. These technologies also are important for propagating at least eight domestic cat models of human genetic disease (W.F. Swanson, personal communication) [85]. For example, such information would be important for optimizing medium composition to maximize sperm survival for artificial insemination, in vitro culture (including for in vitro fertilization), and cryopreservation. Furthermore, we observed that metabolic efficiency reflected other sperm quality metrics across both individuals and species, thus providing the first objective, field-friendly indicator of gamete function in felids. Such findings also offer new insights for improving reproduction in small populations of endangered
species or rare biomedical genotypes. Indeed, a basic understanding of the reproductive uniqueness of a previously unstudied species has been critical to recovery or development of self-sustaining populations [82]. The opportunity to conduct these fundamental studies is one of the invaluable contributions of ex situ (hedge) felid populations to the conservation of Earth’s biodiversity [86].

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