

Mild experimental increase in testis and epididymis temperature in men: effects on sperm morphology according to spermatogenesis stages

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Background: A mild increase in testicular and epididymal temperatures in men, bulls and rams (pendulous scrotum) inhibits spermatogenesis and increases the percentage of sperm with an abnormal morphology. However, the stages of spermatogenesis that are most sensitive to a mild increase in testicular temperature in men are unknown. The aim of the present study was to explore the effects of a mild induced increase in testicular and epididymal temperature (i.e., testicular temperature maintained below the core body temperature) on sperm morphology in humans depending on the physiological time of spermatogenesis and epididymal transit.

Methods: Five healthy volunteers were enrolled in an experimental study in which testicular and epididymal temperatures were increased by maintaining the testes in a supra-scrotal position with a specially designed underwear worn 15±1 h a day for 120 consecutive days. Semen collection was scheduled on specific days depending on spermatogenic stages and epididymal transit.

Results: Sperm morphology and the multiple anomalies index (MAI) were analysed before, during and after heating. This mild induced increase in testicular and epididymal temperatures resulted in a significant rise in the percentage of morphologically abnormal spermatozoa on day 34 of heating, which remained elevated throughout the heating period and persisted until 45 days after cessation of heating. The MAI was significantly increased on day 20 throughout the heating period and persisted 45 days after cessation of heating. An increase in the percentage of anomalies in the sperm head, acrosome or tail occurred on days 34 and/or 45 of heating. Abnormal sperm morphology and MAI reverted to control values 73 days after cessation of heating.

Conclusions: A mild sustained increase in testicular and epididymal temperature in man leads morphological abnormalities in spermatozoa mainly due to an impairment of spermiogenesis and meiosis.

Keywords: Men; meiosis; spermiogenesis; sperm morphology; testis temperature

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Introduction

In men and in most mammalian species, normal physiological spermatogenesis requires a testicular temperature that is 2–6 °C below the core body temperature (1,2). In humans, two thermoregulatory systems are responsible for this lower physiological testicular temperature. The first system consists of a countercurrent heat exchange between arterial blood and spermatic vein blood through the pampiniform plexus. The second system is external heat loss outside the body through passive convection and radiation by the scrotum (2,3). Dysfunctions in thermoregulatory systems or conditions that exceed the efficiency of these systems result in an increase in testicular temperature (4). Furthermore, the complete spermatogenic cycle can be divided into three stages, spermatogonia mitosis (about 28 days), spermatocyte meiosis (about 23 days for meiosis I and 1 day for meiosis II), and spermiogenesis (about 22 days) followed by 12 days of epididymal maturation. Consequently, any change in ejaculate sperm must be interpreted according to this physiology chronology (5).

In mammalian studies including men, testicular and epididymal functions were shown to be acutely sensitive to a temperature increase of only a few degrees (6), and considerable effort has been devoted to analyzing this response. The majority of experimental protocols that induce an increase in testicular temperature in humans by different methods (electric warming bag, polyester-lined supports, testis supra-scrotal localization, sauna, water bath) have a negative impact on spermatogenesis (7-12) and the quality of sperm parameter in most cases. However, depending on the body temperature, an experimental increase in temperature intensity could be classified into two categories: a mild increase when the testicular temperature remains below body temperature, and a high increase when it exceeds body temperature.

To date, four studies have examined the effects of a mild experimental increase in scrotal and testicular temperature on human sperm morphology. In the first study, mild testicular heating was performed using the testis supra-scrotal position (TSP) method which positions the testes near the respective inguinal canals (8,13). This method resulted in a reversible decrease in total sperm output and sperm motility and a reversible increase in morphologically abnormal spermatozoa. In the second study, an improvement in the TSP method resulted in a more intense effect on sperm characteristics (14). Abnormal sperm morphology increased to 180% of the baseline values

during the TSP period (15±1 h daily, 6 to 24 months) and reverted to normal values 10–12 months after cessation of the TSP.

This view has been amply confirmed by Shafik (15), who described the reversible effect of an increase in testis temperature by testicular suspension. In 28 male volunteers the testes were suspended in the superficial inguinal pouch close to the scrotal neck for 12 months. During suspension the percentage of normal sperm morphology decreased over the 12-month period. However, 6–12 months after release sperm morphology returned to normal (15).

Wang *et al.* (16) reported no significant increase in the percentage of spermatozoa with abnormal morphology in men wearing polyester-lined supports to increase scrotal temperature by 0.8 to 1 °C. The absence of negative effects in this study was attributed to an insufficient temperature increase.

In animals, experimental studies explored the effects of a mild temperature increase on sperm morphology in bulls and rams, using scrotal insulation in particular. In these models (scrotal insulation), an increase in testicular and epididymal temperatures induced negative effects on sperm parameters and sperm morphology (17-21).

To date, no study has investigated the effects of a mild increase in testicular temperature on sperm morphology in humans, according to the precise chronology of spermatogenesis and epididymal transit in order to identify at least the onset of morphological changes in sperm.

In this context, the aim of the present study was to explore the effects of a mild induced testicular and epididymal temperature increase (i.e., testis temperature maintained below core body temperature) on human sperm morphology depending on the physiological time of spermatogenesis and epididymal transit. This work is the continuation of our previously published experimental study which explored the deleterious effects of testis temperature on sperm production and DNA fragmentation (5).

Methods

Study population

Five healthy fertile volunteers were recruited (men aged 25–35 years who had fathered at least one child by natural conception). The study was approved by the Ethics Committee (Comité de Protection des Personnes Sud-Ouest et Outre Mer I, ID-RCB 2009-A00977-50) and the protocol has been previously described (5).

The volunteers were enrolled through press advertisement and hospital communication. A total of 34 men volunteered for the study. Further to exclusion criteria and financial limitations, 6 volunteers were selected. These men, who had fathered at least one child, had a normal clinical andrological examination, no current pathology and no medical or surgical history. Their alcohol intake was non-existent or moderate/occasional. They were not exposed to toxic agents and presented no specific occupational fertility risk. One volunteer dropped out of the study on day 73 of heating for personal reasons. Therefore, the data for this volunteer were excluded. The remaining five volunteers followed the instructions and the protocol and completed the study. None of the volunteers reported any discomfort during the entire study period.

In the present study, semen samples from 27 healthy fertile men were analysed to serve as controls before applying mild increase in testicular and epididymal temperature protocol to experimental volunteers (n=5).

Study design

Techniques of testicular exposure to heat

Increased testicular and epididymal temperatures were induced by maintaining the testes in a supra-scrotal position. The testes were maintained in the supra-scrotal position with the underwear provided which had an individually designed orifice to allow exteriorization of the scrotum and penis of each volunteer. The method was previously developed by our group and has been tested safe. The volunteers participating in the heating experiment were told how to put on the underwear by lifting the testes up at the root of the penis on waking up before taking part in the experiment. This specially designed underwear was worn for 15±1 h a day (5,8) for 120 consecutive days. Each volunteer recorded in a diary any event or habits which could modify the protocol. The TSP resulted in a 1.5 to 2 °C increase in testicular temperature (15,22).

Sampling periods

This experimental prospective study was divided into three periods: "before heating" (-40, -15 and 0 days), "during heating" (4, 9, 20, 34, 45, 73, 95 and 120 days) and "after heating" (124, 129, 140, 154, 165, 193, 215 and 300 days). The semen collection days were scheduled according to the physiological chronology of spermatogenesis and epididymal transit. According to these physiological times and the exposure of testicular germinal cells to heat, we

divided the "after heating" period into two phases: (I) phase I when some of the germ cells and different stages of the spermatogenesis cycle were heated, corresponding to 124, 129, 140, 154, 165 and 193 days; (II) phase II when no germ cell or stage of spermatogenesis cycle was heated, corresponding to 215 to 300 days) (*Figure 1*).

Semen collection and analyses

Semen samples were collected by masturbation after a median sexual abstinence of 4.1±1 day. After liquefaction (37 °C for 30 min) semen analysis was performed according to the WHO (23) recommendations as in our previous paper (5). Briefly, semen volume was measured with a graduated pipette and sperm count (×10⁶) was assessed using a Malassez cell (Rogo Sanlab). Semen analyses were performed blindly, and all readings were taken in duplicate within 1 h of sample collection. In this paper, we present only the results of ejaculate volume, sperm count and sperm morphology as all other sperm characteristics from this experimental protocol were reported in the previously published paper (5).

Sperm morphology

Sperm morphology was assessed by observing 100 spermatozoa per sample according to the modified David classification (24,25). We used the Shorr staining procedure for sperm morphology analysis. The type of sperm morphology defects were categorized into head, mid-piece and tail, i.e., head defects (tapered head, thin head, small round-head, macrocephalic, multiple heads, abnormal post-acrosomal region and abnormal acrosomal region), midpiece defects (excess residual cytoplasm, thin midpiece, and bent misaligned midpiece) and tail defects (no tail, short tail, irregularly shaped tail, coiled tail and multiple tails).

The multiple anomalies index (MAI) is the mean numbers of anomalies per abnormal spermatozoon (23). All the head, midpiece, and tail anomalies were included in the MAI calculation.

In order to minimise inter-observer variability, sperm morphology was assessed by a single technician in both the experimental and control groups.

Statistical analysis

Data are presented as median and interquartile ranges Q1–Q3 for tables, due to the number of patients, and as mean and standard deviation for graphs. Sperm morphology and MAI data were compared before, during and after

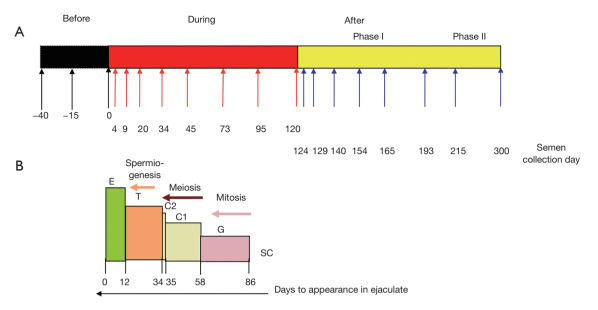


Figure 1 Chronology of semen sampling. (A) Diagram to illustrate the timing of semen sampling during the three study periods before, during and after heating (phase I, phase II); (B) location and evolutionary stages of sperm during the spermatogenic process at heat induction (D0) and their expected appearance in ejaculates. E = epididymal sperm; T = spermatids; C2 = spermatocytes II; C1 = spermatocytes I; G = spermatogonia; SC = stem cells (adapted from: Mild induced testicular and epididymal hyperthermia alters sperm chromatin integrity in men, Fertility and Sterility 2012 with permission from Elsevier).

mild induced testicular and epididymal heating using the Wilcoxon signed rank-sum test, with a P value of 0.50% after a Bonferroni correction. Given the limited number of men included in the experimental protocol, we also conducted a statistical comparison with a control group of 27 fertile men (a single time point evaluation of sperm morphology for each man) using a non-parametric Mann-Whitney analysis.

Considering that there were multiple comparisons, a Bonferroni correction was used and a P value of 0.38% was considered significant. Statistical analyses were performed using SAS software (9.3, SAS Institute). A P value of 5% was considered significant in the absence of Bonferroni correction.

Results

The experimental protocol effectively reduced sperm concentration during heating periods and all men became severely oligozoospermic on D45 with 2 of the 5 volunteers presenting azoospermia on D95 and D120 respectively. From D95 until D154, two men had fewer spermatozoa in their ejaculates while others were either azoospermic or had rare spermatozoa which rendered morphological analysis

invalid at these time points. Therefore, the result of sperm morphology at various time points was presented when the number of volunteers (n=5) was consistent during the study periods (*Figure 2*).

Percentage of normal spermatozoa and MAI

Before heating, the percentage of normal spermatozoa and the MAI did not differ statistically between the control group (n=27) and the experimental group (n=5) (Table 1). Therefore, we compared all experimental values with the values of the control group (see material and methods). The percentage of normal spermatozoa significantly decreased on D34 (during heating), remained significantly lower throughout the heating period until D165 (45 days after cessation of heating) and reverted to control values at D193 (73 days after cessation of heating) (Figure 2). The percentage of morphologically normal sperm was reduced 5-fold during heating. The MAI was significantly increased on D20 (during heating) and returned to the normal mean at D193. Likewise, a drastic decrease in sperm count was observed from D34 to D165 (Figure 2). However, semen volume did not change during the follow-up.

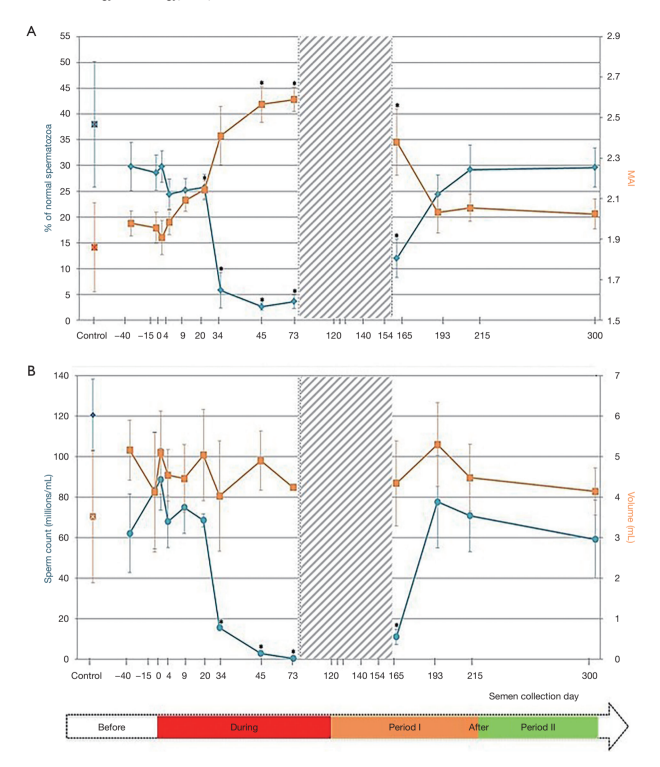


Figure 2 Percentages of spermatozoa with normal morphology, MAI sperm count (millions/mL) and semen volume (mL) before, during, and after (phase I and phase II) mild induced testicular and epididymal heating in men (n=5). (A) Percentages of spermatozoa with normal morphology, MAI; (B) sperm count (millions/mL) and semen volume (mL). Means ± SEM were represented compared to the control group (control; n=27). *, P<0.05. Grey zona: sperm morphology study not possible (see results). MAI, multiple anomalies index; SEM, standard error of the mean.

Table 1 Sperm morphology (%) at follow-up times, before induced testicular and epididymal heating in volunteers, and controls

Variables	Controls (n=27)	Before heating			
variables		-D40 (n=5)	-D15 (n=5)	D0 (n=5)	
Tapered head	1 [0–3]	0 [0–3]	1 [0–1]	0 [0–3]	
Thin head	5 [3–7]	6 [6–7]	9 [2–12]	6 [6–8]	
Small round-head	10 [7–15]	25 [13–26]	20 [19–21]	16 [12–21]	
Macrocephalic	1 [0-2]	0 [0-0]	1 [0-2]	1 [1–1]	
Multiple heads	1 [0-2]	1 [0–1]	0 [0–1]	1 [0–1]	
Abnormal post-acrosomal region	25 [12–31]	20 [14–26]	17 [13–32]	21 [21–22]	
Abnormal acrosomal region	36 [25–47]	45 [38–48]	41 [40–54]	42 [38–47]	
Excess residual cytoplasm	3 [2–6]	3 [2–4]	4 [2–4]	4 [3–7]	
Thin midpiece	2 [0–3]	1 [1–2]	1 [0-4]	0 [0–1]	
Bent or misaligned midpiece	17 [11–21]	19 [17–24]	15 [14–20]	18 [17–22]	
No tail	1 [0-2]	0 [0–2]	1 [0–1]	1 [1–2]	
Short tail	0 [0–1]	1 [0–1]	1 [1–1]	1 [0–2]	
Irregularly shaped tail	0 [0–1]	1 [0–1]	2 [0–2]	1 [1–2]	
Coiled tail	9 [4–13]	13 [9–17]	15 [14–17]	9 [5–11]	
Multiple tails	1 [0-2]	0 [0–1]	1 [1–1]	1 [1–1]	
Normal spermatozoa (%)	38 [26–44]	30 [25–34]	26 [25–30]	28 [25–31]	
MAI	1.86 [1.68–1.94]	2.00 [1.98–2.06]	1.97 [1.84–2.02]	1.86 [1.76–1.98]	

Values are expressed as a median with an interquartile range [q1-q3]. MAI, multiple anomalies index.

Details of sperm morphology changes

Before heating

Before heating, controls (n=27) and volunteers (n=5) had comparable results for sperm morphology.

During heating

Testicular and epididymal heat exposure led to an increase in the percentage of sperm head anomalies compared to the control group (n=27) as early as D34. A notable increase was documented for thin head [27% (10–31%), small roundhead 29% (29–33%)] and abnormal acrosomal region [79% (78–88%)] (*Table 2*). The percentage of spermatozoa with either a thin head or small round-head was 3 times higher than the baseline mean values (D0, D15 and D40) (*Figure 3*).

Similarly, the percentage of spermatozoa with short or irregular tails was significantly higher [2% (1–4%) and 1% (1–5%) respectively] compared to control group values on day 34 whereas the percentages of coiled [29% (21–30%)] or multiple tailed [4% (4–4%)] spermatozoa were

significantly higher on D45 and D73 respectively (*Table 2*). Mean percentages of spermatozoa with short or coiled tails were 2 times higher than the baseline mean values on D45 (*Figure 3*).

After the heating period (phase I)

D165, i.e., 45 days after cessation of heat exposure, was the first time point when ejaculates had an adequate number of sperm to perform a morphology analysis. The percentages of spermatozoa with thin heads [14% (10–18%)] or abnormal acrosomal regions [72% (54–80%)] remained significantly higher (*Table 3*) than in the control group (n=27) until D165.

All morphological characteristics and the overall sperm morphology returned to normal values at D193, i.e., 73 days after cessation of heat exposure (*Table 3*).

After the heating period (phase II)

On D215 (95 days after cessation of heat exposure)

Table 2 Sperm morphology (%) at follow-up times, during induced testicular and epididymal heating in volunteers, and controls

	(20-2) 0000			During	During heating		
valables		D4 (n=5)	D9 (n=5)	D20 (n=5)	D34 (n=5)	D45 (n=5)	D73 (n=5°)
Tapered head	1 [0–3]	0-0] 0	1 [0–1]	0 [0–1]	0-0] 0	0 [0–1]	2 [2–2]
Thin head	5 [3–7]	3 [3–10]	6 [6–12]	8 [7–9]	27 [10–31]*	22 [16–23]*	14 [14–22]*
Small round-head	10 [7–15]	17 [15–24]	18 [17–19]	23 [18–28]	29 [29–33]*	29 [28–44]	41 [34–44]
Macrocephalic	1 [0–2]	0 [0–1]	0 [0–1]	1.5 [0–2]	1 [0–1]	0 [0–2]	0 [0–1]
Multiple heads	1 [0–2]	1 [1–2]	2 [1–4]	1.5 [0–2]	2 [1–2]	4 [4–5]	2 [2–5]
Abnormal post-acrosomal region	25 [12–31]	26 [20–28]	20 [19–32]	18.5 [17–26]	24 [21–26]	32 [17–38]	24 [22–46]
Abnormal acrosomal region	36 [25–47]	50 [48–55]	47 [45–50]	53 [46–54]	79 [78–88]*	*[06-58] 06	88 [76–90]
Excess residual cytoplasm	3 [2–6]	4 [2–7]	4 [2–6]	3 [2-4]	4 [3–6]	3 [3–5]	4 [0–5]
Thin midpiece	2 [0–3]	2 [1–4]	2 [2–2]	2.5 [0-3]	0 [0–1]	1 [0–5]	2 [0-4]
Bent or misaligned midpiece	17 [11–21]	14 [13–22]	22 [12–26]	21.5 [21–28]	29 [25–29]	26 [26–30]	23 [22–26]
No tail	1 [0–2]	2 [1–2]	1 [0–1]	0 [0–2]	3 [2–4]	4 [4–7]	4 [4–8]
Short tail	0 [0–1]	0 [0–1]	1 [0–1]	1.5 [0–2]	2 [1–4]*	3 [1-4]*	3 [2–5]*
Irregularly shaped tail	0 [0–1]	1 [1–1]	1 [0-3]	1 [0–1]	1 [1–5]*	2 [1–4]	2 [0–2]
Coiled tail	9 [4–13]	10 [9–12]	15 [10–17]	16.5 [14–19]	25 [14–26]	29 [21–30]*	21 [18–34]*
Multiple tails	1 [0–2]	1 [0–1]	3 [2-4]	1 [0–1]	2 [1–2]	3 [1–3]	4 [4–4]*
Normal spermatozoa (%)	38 [26–44]	27 [22–29]	26 [21–30]	26 [24–27]	5 [0–19]*	3 [3–3]*	5 [1–6]*
MAI	1.86 [1.68–1.94]	1.97 [1.95–2.07]	2.10 [2.04–2.16]	2.14 [2.09–2.2]*	2.50 [2.42–2.58]	2.46 [2.42–2.75]*	2.52 [2.49–2.68]*

Values are expressed as a median with an interquartile range [q1-q3]. °, at D73, 50 spermatozoa were analysed instead of 100 for 3 patients; *, Values differ significantly from the control group values.

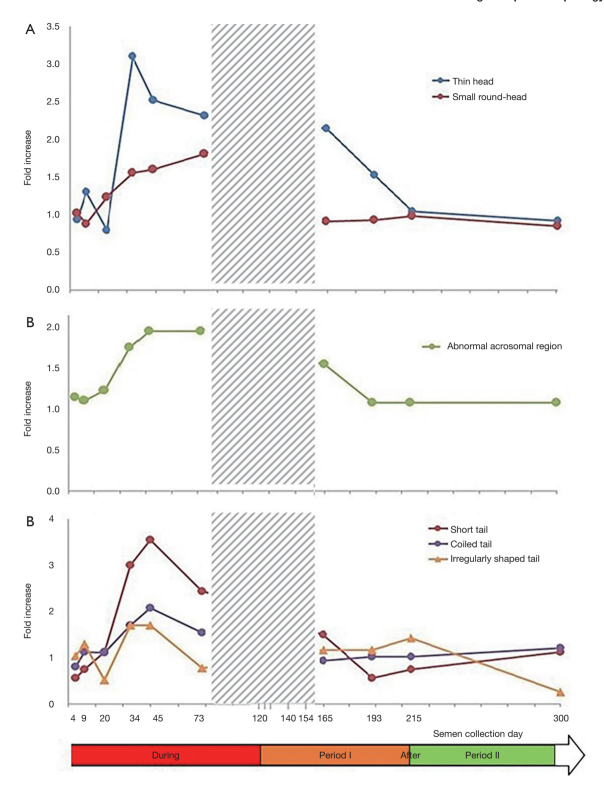


Figure 3 Description of increase in some specific sperm morphology anomalies during and after (phase I and phase II) mild induced testicular and epididymal heating in men: (A) head abnormalities; (B) acrosome-related abnormalities; (C) tail abnormalities. Data concerning fold increase in sperm anomalies during and after heating were compared to baseline mean values at -40, -15 and 0 day before heating. Grey zona: analyses between day 73 and day 154 were not possible (see results).

Table 3 Sperm morphology (%) at follow-up times, after induced testicular and epididymal heating in volunteers and in controls

Variables	Controls (n=27)	After heating			
		D165 (n=5)	D193 (n=5)	D215 (n=5)	D300 (n=5)
Tapered head	1 [0–3]	3 [2–6]	1 [0–2]	1 [0–1]	0 [0–1]
Thin head	5 [3–7]	14 [10–18]*	7 [7–16]	6 [5–9]	6 [5–10]
Small round-head	10 [7–15]	14 [8–26]	18 [18–21]	15 [13–25]	14 [9–21]
Macrocephalic	1 [0–2]	3 [1–4]	1 [0–1]	0 [0–0]	1 [1–1]
Multiple heads	1 [0–2]	2 [2–3]	2 [1–2]	1 [0–3]	2 [0-3]
Abnormal post-acrosomal region	25 [12–31]	36 [34–39]	22 [22–32]	21 [20–21]	21 [21–29]
Abnormal acrosomal region	36 [25–47]	72 [54–80]*	42 [41–54]	41 [39–54]	46 [40–56]
Excess residual cytoplasm	3 [2–6]	8 [8–12]	5 [4–5]	5 [5–6]	2 [2–5]
Thin midpiece	2 [0–3]	2 [1–2]	2 [1–2]	1 [1–2]	3 [1–4]
Bent or misaligned midpiece	17 [11–21]	25 [21–27]	23 [18–28]	19 [16–24]	16 [14–22]
No tail	1 [0–2]	2 [2–3]	1 [0–1]	1 [1–1]	1 [1–5]
Short tail	0 [0–1]	1 [1–3]	0 [0–1]	0 [0–1]	1 [1–1]
Irregularly shaped tail	0 [0–1]	1 [0-4]	0 [0–3]	1 [1–2]	0 [0–1]
Coiled tail	9 [4–13]	13 [11–13]	11 [9–17]	10 [10–18]	17 [9–22]
Multiple tails	1 [0–2]	3 [3–4]	2 [2–3]	1 [1–1]	2 [1–3]
Normal spermatozoa (%)	38 [26–44]	11 [4–20]*	26 [18–29]	33 [21–37]	33 [23–33]
MAI	1.86 [1.68–1.94]	2.42 [2.1–2.73]*	2.10 [1.84–2.16]	2 [1.97–2.08]	1.98 [1.91–2.04]

Values are expressed as a median with an interquartile range [q1-q3]. *, values differ significantly from the control group values.

sperm morphology and percentage of sperm anomalies returned to baseline values (*Table 3*) which indicates that the effects of heat exposure were reversible but at least one spermatogenesis cycle is required before sperm morphological characteristics return to normal.

The incidence of men with abnormal sperm morphology values

In this study, we stated that a man was deemed to have an abnormal morphology parameter when the individual value recorded during the experimental temperature increase was higher than the maximum value observed before initiating experimental heating in all volunteers. This allowed us to calculate the frequencies with which the men presented abnormal values. The results (*Figure 4*) indicate an increase in the number of volunteers with abnormal sperm head morphology during heat exposure which started on D34 for thin head, small round-head, abnormal acrosomal region and coiled tail.

No difference was observed in the number of volunteers with abnormal midpiece morphology at any time point (4 to 300 days). However, an excess of residual cytoplasm was noted in 4/5 men on Day 165. More volunteers showed abnormal sperm tail morphology after 34 days and throughout the heat exposure period, except for irregularly shaped tail defects.

As indicated in *Figure 4*, most of the volunteers presented abnormal sperm morphology and the MAI as early as D34, which persisted throughout heat exposure until D165 (45 days after cessation of the temperature increase).

Discussion

To date, this is the first study to report detailed changes in the morphological characteristics of sperm induced by a mild increase in testicular and epididymal temperature based on the physiological chronology of spermatogenesis and epididymal transit in humans. The key feature of the TSP method is that the increase in testicular and epididymal

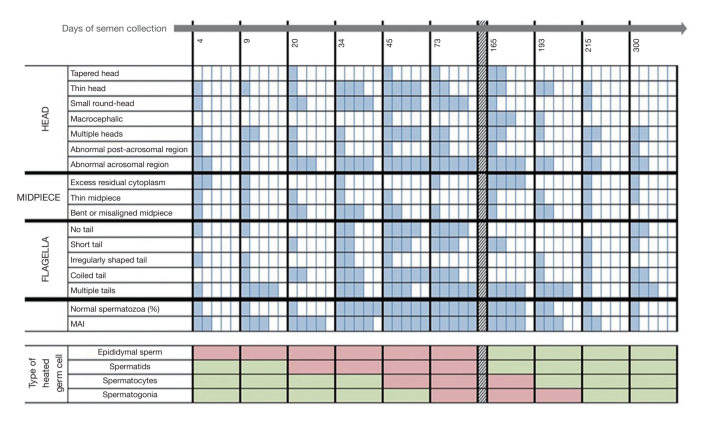


Figure 4 Diagram to illustrate the number of volunteers with a higher percentage of abnormalities during and after mild induced testicular and epididymal heating than at baseline before heating (see results) and which correspond to the spermatogenesis stage and epididymal transit heating chronology. Each cell represents one volunteer. For one anomaly, a blue cell means that a volunteer has an abnormal percentage. A white cell means that a volunteer has a normal percentage. A green cell means that the stages of spermatogenesis or epididymis were not heated. A red cell means that the stages of spermatogenesis or epididymis were heated.

temperature is mild and remains below body temperature. We previously reported a drastic decline in sperm production and an increase in sperm DNA fragmentation using this method (5). In this study, we focus on the effects of a mild induced increase (1.5–2 °C) in testicular and epididymal temperature on sperm morphology.

No effects on epididymal spermatozoa

According to epididymal transit, spermatozoa collected on D4 and D9 of heating were more likely to have been in the epididymis when the temperature was increased. In this study, sperm morphology was not affected by this mild increase in epididymal temperature at these time points (D4, D9). There is a lack of information in the literature and, to date, no study has been conducted in men to show the effect on sperm morphology during the transit of spermatozoa through the epididymis on exposure to a mild temperature

increase.

Nonetheless, in bulls and rams, two species with a pendulous scrotum, a mild increase in scrotal temperature was documented through scrotal insulation (a summary of these studies is available in Figure S1). In bulls, 2 out of 8 studies reported an effect on sperm morphology during spermatozoa epididymal transit on the time of heating introduction (26,27). In rams, studies by Junior et al. (18) and Rocha et al. (28) reported the effect on sperm morphology of an increase in testicular temperature by scrotal insulation, during the transit of sperm through epididymis. Rocha et al. (28) observed a significant decrease in the percentage of morphologically normal sperm from 92%±2.7% on day 0 (pre-insulation) to 39%±4.2% on day 8 of scrotal insulation. These effects were caused by a temperature increase in scrotal skin from 31.2±0.2 °C preinsulation to 35.2±0.3 °C during insulation for 24 h/day over 8 consecutive days. Similarly, Junior et al. (18) reported that

an increase in testicular temperature by scrotal insulation caused an increase in the percentage of sperm with abnormal morphology in the first week post-insulation. The main defects observed were bent tail (35%), coiled tail (16.25%), distal cytoplasmic droplet (14%) and normal detached head (20%).

The discrepancy between our study and animal studies reporting morphological changes in epididymal spermatozoa might be attributed to the variability in the physiology of species and/or in the methods and duration of heating. The epididymis of rams appears to be more sensitive to temperature increases than the human epididymis. This might be due to two physiological factors: (I) in humans the core testicular temperature gradient is approximately 2.8 °C (29) in contrast to 6-8 °C in rams with a body temperature that was modified by scrotal insulation (30); (II) the ram epididymis has a significantly longer tail (29,31). However, while no modifications in sperm morphology have been observed in spermatozoa localised in the epididymis during heating, it should be remembered that epididymal sperm maturation is a heat-sensitive process and alterations in sperm functions such as embryo development, without modifications in classical sperm parameters, have been reported as early as day 4 of mild scrotal heating in rams (30).

Effects on testicular germ cells

According to spermatogenesis and epididymal physiology (5), spermatozoa collected on D20 were at the elongated spermatid stage in the testes when heating was induced. Those collected on D34 were at the late spermatocyte stage (end of meiosis) or the spermiogenesis stage. Those collected on D45 were in the meiosis stage and finally those collected from D56 to D86 were at the mitosis stage of spermatogenesis.

The significant increase in the MAI, which is an integrated index of sperm morphological anomalies and a good predictive value in couples' fertility assessment (32), on D20 suggests that late spermiogenesis was affected by heat induction. However, the drastic fall (5-fold) in the percentage of normal sperm on D34 might be due to effects on early spermiogenesis and the late stage of meiosis. These results are consistent with the fact that pachytene spermatocytes and round spermatids are the testicular cells most sensitive to heat stress (33).

To date, no human study has reported such an early effect (before 34 days during heat induction) of a mild

testicular temperature increase on sperm morphology. Using a similar method of heat induction (TSP) in 10 men, a significant increase in the percentage of sperm with abnormal morphology was observed in semen collected between weeks 5 and 8 after the start of heating, i.e., between day 29 and day 56 of heating (8). In another study including 28 male volunteers the testes were suspended in the superficial inguinal pouch close to the scrotal neck 24 h/day for 12 months (15). The percentage of sperm with a normal morphology dropped from 60% before heating to 27%, with 12% at 3 and 12 months respectively during heating. However, 6–12 months after release the percentage of sperm morphology returned to normal (15). Three other studies have reported the effects of scrotal insulation on semen parameters in men (11,16,34). Two reported a significant decrease in sperm count (11,34) but sperm morphology was not analysed. The other study did not report any change in sperm count or morphology. However, the temperature increase in this last study was very low (increase in scrotal temperature of 0.8–1 °C) (16).

A recent study examined the sperm morphology in fertile men before, during and after scrotal heating with an electric warming bag to increase testicular temperature to 43 °C for 30–40 min 2 days per week for 3 months (12,35), which is not a mild testicular increase but a high one. The percentage of sperm with normal morphology was significantly decreased at 1, 2, and 3 months of scrotal heating compared to the percentage before heating. Overall, the results of Zhang *et al.* (12,35) and our previous studies (8,13,14) indicate that both a mild sustained induced increase in testicular temperature and a high short-term induced increase in testicular temperature result in a decline in normal sperm morphology which is consistent with the results of the present study.

One important aspect of our study is the description of abnormalities induced by the mild testicular-epididymal temperature increase. It is interesting to note that, in the present study, thin heads, small round-heads, and sperm with abnormal acrosomal regions were the head anomalies induced by the mild temperature increase. Tails anomalies such as short tail—an irregular shaped tail which could indicate anomalies of per-axonemal structures (36,37) were significantly increased from D34. These changes on D34 suggest an alteration in the spermiogenesis processes. A significant increase in the percentage of sperm with coiled tails was noted on D45 and multiple tails on D73 suggesting that germ cells in earlier stages of spermatogenesis were affected to greater extent.

Abnormal sperm morphology has been connected to sperm DNA damage (38-40), and we (5) previously reported a significant increase in sperm DNA fragmentation index (DFI) and mean sperm high DNA stainability (HDS) after 20 to 45 days of heat exposure. Moreover, in bulls, scrotal exposure to heat (scrotal insulation method) is related to defective nuclear chromatin condensation during the acrosome and Golgi phases of spermiogenesis (17,19). The significant increase in protamine deficient sperm cells during this period (spermiogenesis and spermatocytes) indicates that scrotal insulation is, at least partly responsible for defective chromatin protamination (41). The same outcome in terms of protamine modifications was also observed following an episode of fever in men (42).

The percentage of normal sperm remained significantly depleted and the MAI increased until D165 which corresponds to 45 days after the cessation of heat (post heating phase I). The recovery of sperm morphological values comparable to the control group was observed on D193. These results indicate that normal sperm morphology values did not reappear until one spermatogenesis cycle had been completed after the cessation of heating. In contrast, Mieusset et al. (14) reported the recovery of a percentage of normal sperm only at 10–12 months, after the completion of TSP. However, in their study the duration of experimental heating was prolonged (24 months). Using another scrotal heating method with an electric warming bag (43.0±0.5 °C, 30–40 min 2 days per week for 3 months), Zhang et al. (12,35) noted that the percentage of sperm with a normal morphology gradually returned to normal levels two months after scrotal heating.

In bulls, significantly higher percentages of sperm abnormalities such as head abnormalities, acrosome defects, pyriform-shaped heads, micro- and macro-cephalic heads and tail defects were observed from 14 to 42 days after heating in the majority of studies (17,19-21,43) and from 10 to 11 days in two studies (27,44). These results were consistent with the effect of heating during the spermiogenesis and meiosis stages. In rams, the effects lasted longer, up to 77 days (more than one spermatogenesis cycle) after heat induction while the duration of heat exposure was brief (duration of exposure: 3 to 8 days) (18,28,45). The mechanism of a long-term effect must be clarified.

In humans, such a mild temperature increase could be caused by several conditions. Fever or occupational heat exposure has been connected to an increase in abnormal sperm morphology. Andrade-Rocha (46) reported a decrease

in sperm concentration and an increase in the percentage of abnormal forms, particularly small-head sperm following high fever (38–39 °C). Working conditions such as welding also affect the percentage of normal forms. Bonde (47) observed a decrease in normal sperm morphology in men 4 to 6 weeks after heat exposure (on average, the skin temperature in the groin increased 1.4 °C) which is consistent with the notion that an induced rise in testicular temperature affects the spermiogenesis—meiosis stages. Similarly, Figa-Talamanca *et al.* (48) reported an increase in sperm morphology abnormalities in workers exposed to high temperatures in the ceramics industry.

In general, the human sperm morphology assessment is regarded as an important semen parameter that is correlated with fertility potential, the probability of conception and pregnancy outcome and pregnancy loss in *in-vivo* and *in-vitro* studies (49-52). Therefore, abnormal sperm morphology can lead to delayed and/or failed conception.

The first striking feature of our study is the experimentally controlled mild increase in testicular and epididymal temperature, which remains below core body temperature, to which men are exposed during routine daily activities. The second strength is the precise chronological analysis of sperm morphology throughout the spermatogenic cycle and epididymal transit. However, one major limitation is the small number of volunteers enrolled in the study.

It is interesting to note that, in our current study and studies reported by others, the changes observed in human and animal sperm morphology following a mild temperature increase were always reversible.

Conclusions

This experimental study shows that even a mild induced increase in testicular and epididymal temperature can severely compromise sperm morphology which is an integral sperm parameter in fertility assessments and in the predicted outcome of assisted reproductive technologies. Continuous and long-term occupational/environmental exposure to slightly higher temperatures could impair sperm morphology and, ultimately, the likelihood of natural conception. Therefore, the history of occupational and environmental exposure to increased temperature should be considered in fertility assessments. Physicians must counsel patients to avoid any risk of exposure to testicular temperature increase, however mild.

On the other hand, to given the reversibility and safety of

this experimental method, a mild testicular and epididymal temperature increase may be considered an alternative approach to male contraception.

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Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was approved by the Ethics Committee (Comité de Protection des Personnes Sud-Ouest et Outre Mer I, ID-RCB 2009-A00977-50) and the protocol has been previously described.

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Supplementary

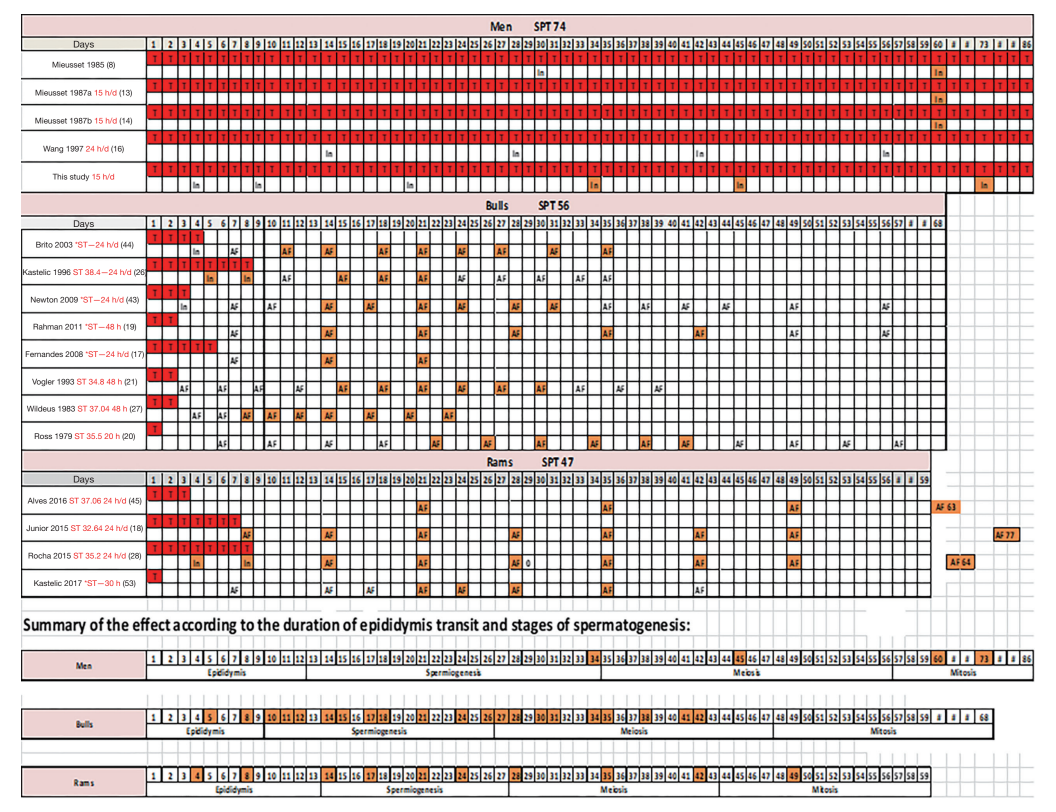


Figure S1 Presentation of the effects of a mild testicular temperature increase on sperm morphology in men, bulls and rams according to the duration of epididymis transit and stages of spermatogenesis (8,13,14,16-21,26-28,43-45,53). Animal studies used scrotal insulation. Red squares represent heating periods. Orange squares represent a significant effect on sperm morphology. White squares represent an absence of significant effect. In: during heating; AF: after heating; STP: duration of spermatogenesis; ST: scrotal temperature. *, ST: no scrotal or testicular temperature measurement.

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