Ultra-differentiation of sperm head in Egyptian lesser jerboa, Jaculus jaculus (Family: Dipodidae).

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Abstract:

In the present study, events of sperm head differentiation in Lesser Egyptian Jerboa, *Jaculus jaculus* were studied for the first time. Adult males of *J. jaculus* were collected during their period of sexual activity from sandy regions of Marsa-Matrouh at north-west of Egypt. Tissues of their testes were prepared for ultrathin sections, which examined under a Joel "JEM- 1200 EXII" operating at 60-70kv. Early and late spermatids were photographed to describe successive stages of sperm head differentiation.

Early spermatids have rounded or oval nuclei with fine chromatin granules and their cytoplasm showed numerous mitochondria, and one or more chromatoid bodies and segments of rough endoplasmic reticulum. The first stage of spermatid development usually starts when Golgi body produces secretory vesicles. These vesicles usually differentiate into an oval dense acrosomal granule, the rest forming a thin layer of acrosomal cap which extends to cover the anterior half of the nucleus and stop on at the nuclear shelf in the equatorial nuclear region. This cap is separated from the nuclear envelope by a narrowed subacrosomal space. Novel and complex structures are observed in the developing acrosome, which are, the crown, anterior, and posterior acrosomal segments, anterior and posterior acrosomal caps, as well as a long dorsal and a short ventral acrosomal caps; posterior subacrosomal spaces and subacrosomal cone at the tip of the elongated nucleus.

Cytoskeletal elements are responsible for re-shaping of the nucleus. A light comprehensive strength of cytoskeletal elements usually induces nuclear prolongation and formation of implantation fossa that appears in the ventro-dorsal region at the posterior side of the nucleus. Manchette microtubules, solitary microtubules and microfilaments may generate gentle compressive strength, to accelerate nuclear prolongation.

Manchette microtubules which disposed parallel to one another and to the long axis of the nucleus could exert the force, required to produce the spermatid nucleus elongation forward and perhaps backward and to protect DNA during nuclear condensation. A translucent space appears to surround the posterior half of the nucleus in order to mitigate the pressure on the nucleus and regulate the elongation with the protection of genetic material during nuclear condensation. Worth mentioning, that the translucent perinuclear space is a unique structure was not described or discussed before.

Keywords: Ultrastructure, sperm head, Lesser Egyptian Jerboa, Jaculus jaculus, Rodents,

1. Introduction

In the mammalian seminiferous tubules, two stages in which spermatogenic cells produced male gametes by way of mitosis and meiosis. First, known as spermatocytogenesis, spermatogonia developed into spermatids; the second is called spermiogenesis, spermatids reformed into spermatozoa [1].

After second meiotic division, Spermatids derives from the division of secondary spermatocytes, and undergoes a series of morphological and cytological changes leads to formation of fully developed spermatozoa. Under the subcellular level, stages of sperm head and/or sperm tail differentiation were taken into consideration by the scientists [2-7]. In mammalian spermiogenesis, the nucleus of spermatid becomes elongated; its chromatin condensed; the anterior nuclear protrusion is capping by acrosome [6-7]; the centrioles migrate to the prospective posterior end of the nucleus then situated into a depression, nuclear implantation fossa, where the connecting piece, mitochondrial and fibrous sheath are formed followed by extension of an axial filament surrounded by special fibrous sheath [8].

Publications interests of spermiogenesis are focused on several themes include differentiation of normal and abnormal sperm-head [2]⁻ and/or sperm-tail [12, 15] as well as other detailed investigations directed to study the role of some organelles such as Golgi apparatus [3-7], microtubules [8-12] and chromatoid bodies [13] involved in the formation of acrosome [14], manchette formation of spermatid cytoskeleton

according to sperm needs [10-13], and configure the fibrous sheath [15-20] along the axial filament [21] in the human [22, 24] and mammalian spermatozoa [25-32].

In order Rodentia [33-36], few investigations are deled with the ultrastructure of sperm head or sperm tail differentiation, especially in Egyptian fauna [37-39]. Moreover, mechanisms of the interesting formative events in sperm head of Egyptian mammals are still elude the present researcher. This work aimed to investigate specific characterization of sperm head differentiation in the Lesser Egyptian Jerboa, *Jaculus jaculus* as a one of the threatened species [40].

2. Material and Methods

2.1. Animals

Adult males of *J. jaculus* were collected during their period of sexual activity from sandy regions of Marsa-Matrouh at north-west of Egypt.

2.2. Tissue preparation

After light ether anesthesia, testes were immediately extracted, washed in cacodylate buffer solution adjusted at pH7.2, cut into two halves and fixed in 2.5% glutaraldehyde in the refrigerator. 2hrs later, thin slices were taken and cut into smaller specimens (about 0.5-1.0 mm thick), rewashed in fresh cold buffer to remove depressed tissues and transferred again into fresh cold 2.5% glutaraldehyde for 4-6 hours. The fixative removed, and specimens washed 3 times in buffer and post-fixed 3 hrs. in 1% osmium tetroxide/buffer, then washed in buffer followed by dehydration, clearing using propylene oxide and embedding in Epon-Araldite mixture.

Semithin sections about 1µm thick were taken, stained using toluidine blue, examined under light microscope to select best locations of spermiogenesis activities, especially sites of sperm head differentiation. Ultrathin sections were stained using aqueous uranyl acetate (1.5%) and lead citrate then examined under a Joel "JEM-1200 EXII" operating at 60-70kv. Early and late spermatids were observed and magnified and photographed to describe successive stages of sperm head differentiation.

3. Results

In the seminiferous tubules, the epithelium consists of sustentacular cells known as Sertoli cells (Ser), which are tall, columnar cells that line the tubules and spermatogenic cells, which differentiate through meiosis to spermatids and subsequently undergoes morphological stages to form sperm cells (fig. 1). Early spermatids have rounded or oval nuclei with fine chromatin granules and their cytoplasm showed numerous mitochondria (M), one or more chromatoid bodies (CB) and segments of rough endoplasmic reticulum (RER) (fig. 2). The first stage of spermatid development usually starts when Golgi body (G) produces secretory vesicles (SV) (figs. 3, 4). These vesicles usually differentiate into an oval dense acrosomal granule (AG), the rest forming a thin layer of acrosomal cap (AC) which extends to cover the anterior half of the nuclear membrane and separates from it by a narrowed subacrosomal space (SAS). This cap surrounds the anterior half of nuclear envelope and stop on at the nuclear shelf (NS) (fig. 5). The next stages show the nuclear re-shaping, chromatin condensation, reformation of cytoskeleton, completion of acrosome and other regions of the developing sperms (figs 6-20).

3.1. Role of manchette in the shaping of the sperm nucleus

Mainly, the cytoskeleton in active spermatid is formed of microtubules and microfilaments (figs. 3, 5, 7). They pass in circular pathways, thus, they are distributed under the cell membrane, in the cytoplasm and around the nuclear envelope (figs. 3-5). Some elements of cytoskeleton re-orient the posterior end of nuclear envelope to form a posterior concavity, implantation fossa, to lodge the proximal centriole (fig. 5). Figure 6 proves that this concavity appears in the posterior ventromedial side of the nucleus. Another large unit of microtubules (MtU) move to surround the equatorial of the nucleus at the nuclear shelf (NS) where the acrosomal cap stops on (fig. 7). At this point, front half of circular microtubules that surround the nucleus are reduced pushing the cell organelles backwardly, in the same time; some of the posterior half microtubules attach themselves to the MtU, re-oriented parallel to one another and to the long axis of the nuclear envelope then end at the rear when docking with the basal plate (BP). In addition, some solitary smaller microfilaments are arranged in a transverse direction to surround the posterior perinuclear space (PP)

that firstly appears to surround the posterior ventral side of the nucleus (figs. 8-13). Both MMs, solitary Mts and microfilaments (Mfs) may be generate gentle compressive strength for the nucleus elongation forward and perhaps backward and to protect DNA during nuclear condensation. The translucent perinuclear space appears to surround the posterior half of the nucleus in order to mitigate the pressure on the nucleus and regulate the elongation with the protection of genetic material during nuclear condensation (fig. 8). In the present stage, the nuclear condensation starts in parallel with the nuclear elongation. Next stage showed that MMs extend to the posterior side of the nucleus and unite with the basal plate (BP) at the posterior nuclear shelf (PNS) that shown in figures 8-13. Also, numerous Mfs appear on a single or double row surrounding the posterior half of the nucleus as they run in circular path (fig. 9).

In late spermatid, as shown in figures 16 and 17, it is worth mentioning that the cytoskeleton is formed of solitary microtubules (Mts) and microfilaments (Mfs). The former are distributed along the anterior (ASS) and posterior subacrosomal spaces (PSS) and some Mts are observed in the outer side of the acrosomal cap and near the posterior nuclear shelf (PNS), in addition, some microfilaments appear in the subacrosomal cone. Moreover, the cytoskeleton in the posterior region of the nucleus showed longitudinal MMs, or solitary microtubules (Mts), in addition numerous microfilaments that run in circular pathways to support the connections between the sperm head (SH) and caudal structures via the neck piece (NP) (fig. 19, 20).

It is not easy to conceive of any mechanism whereby the manchette microtubules disposed parallel to one another and to the long axis of the nucleus could exert force perpendicular to their own axis. Yet, force in this direction would be required to produce the flattening of the sperm nucleus that is common to nearly all mammals.

3.2. Nuclear Shaping

Early spermatids have round nucleus (N) filled with aggregations of fine chromatin granules (Chs), the surrounding cytoplasm illustrated numerous mitochondria (M), and few segments of RER, in addition, the presence of Golgi apparatus (G) starts to produce secretory vesicles (SVs) that will aggregate to form acrosome (figs. 3-6). Figure 7 illustrates the distribution of cytoskeletal-elements responsible for re-shaping of the nucleus. Later, a light comprehensive strength of cytoskeletal elements induced nuclear prolongation and formation of implantation fossa that appears in the ventro-dorsal region at the posterior side of the nucleus (figs. 5, 6). As the nuclear elongation proceeds, the fine chromatin granules aggregate and chromatin condensation begins (figs. 8, 9).

In the next stage, when the posterior peri-nuclear space appears, the MMs and other cytoskeletal organelles, that surround it, generate compression strength to accelerate the nuclear prolongation towards the posterior direction (fig. 10) and consequently to the anterior end (figs. 11-13). The chromatin condensation proceeds then slows in parallel with the reduction of the perinuclear space. Thus, these thinsections explain the relationship between the nuclear elongation, the formation of perinuclear space and MMs strength. In the same time, the caudal-ventral side of the nucleus is thickened to form a basal plate at which the posterior nuclear shelf appears (fig. 12), while in figure 13, the posterior ventro-medial portion, the implantation fossa (IF) is formed to lodge the proximal centriole (PC) in the sperm neck (SN). However, there is a unique smaller space that appears on one side at the pre-equatorial plane (PeS). This space appears in conjunction with the elongation of the nucleus (fig. 10) and disappears after the completion of chromatin condensation (figs. 14). It is not observed before in mammalian spermatids.

In late spermatids, the nuclear elongation and chromatin condensation is configured. At the present stage, the nucleus acquires a long conical shape with acute tip that end at the subacrosomal cone, filled with dense chromatin, and the presence of numerous nuclear canals (NCa). This new shape is supported by a well-organized cytoskeletal structures that are distributed in the anterior (ASS) and posterior subacrosomal space (PSS), in addition, the microfilament in the subacrosomal cone (figs. 16, 17).

Also worth mentioning, in the nucleus of the present spermatid, there are an anterior and posterior nuclear shelves (Fig. 18) and unique implantation fossa (IF) at the posterior ventro-dorsal side of the nucleus (figs. 19, 20).

3.3. Differentiation of acrosome

Fortunately, novel and complex structures are observed during the formation of the acrosome in the present spermatid as will be explained. The first sign of acrosomal formation is the activity of Golgi apparatus (G), which secretes elements of the acrosome at the prospective anterior half of the nucleus in the form of acrosomal vesicle (AV) and acrosomal granule (AG) (figs. 3, 4). Then, during nuclear elongation, both the acrosomal vesicle (AV) and granule (AG) spreads to occupy the upper half of the nuclear envelope and stop on at the nuclear shelf (NS) (figs. 5-7).

In the next stage, after the beginning of nuclear elongation and chromatin condensation, the acrosomal granule appears as a dense large triangular acrosomal mass that will be differentiate into new described four regions. The former is a translucent terminal segment (TS) (fig. 9), the remaining three portions appeared as a novel triangular acrosomal mass, which will be differentiates as shown in figures 10-13 into the crown (SC), anterior (AAS) and posterior acrosomal segments (PAS) at which a long acrosomal structure capping the anterior two-third of the nuclear envelop "anterior acrosomal cap "(AAC)" and separating from it by a narrow subacrosomal space (SAS) and a smaller subacrosomal cone (SC) (fig. 13). Note that the acrosomal cap, observed in figure 10, extends towards the posterior direction of the nucleus as anterior acrosomal cap (figs. 11-13), which terminates at the equatorial region of the nucleus, which is called anterior nuclear shelf (ANS). Moreover, there is a novel triple branched structure, which is formed of a terminal dense rod (TDR) and two lateral dense arms (LDA) (figs. 12, 13).

In late spermatid, as shown in the sagittal ultrathin section (figure 14), proved that the acrosomal cap differentiates into, a novel, long dorsal (DAC) and short ventral acrosomal cap (VAC), and consequently, both of them ends at the dorsal nuclear shelf (DNS) and a ventral one (VNS), respectively. Meanwhile, in the anterior longitudinal section, as shown in figure 15, the anterior and posterior acrosomal cap terminate at the anterior nuclear shelf (ANS) that appears at the lateral equatorial portions of the nucleus in parallel to the dorsal acrosomal cap (DAC) in the previous figure. From the above observations, the acrosomal cap covers about the anterior two-thirds of the nucleus (figs. 14, 15) at the dorsal and both lateral sides, while less than half nuclear length of the nucleus at its ventral side (fig. 14). At this point, the anterior acrosomal cap (PAC) that destines at posterior nuclear shelf (PNS) forming small swallow fusiform caudal fusiform end (CFE) of the posterior acrosomal cap (PAC) (figs. 15-18).

At this point, the anterior acrosomal cap swollen at its terminal ends as a posterior dense fusiform structure (PFE), and then, it extends towards the caudal

nuclear region forming posterior acrosomal cap (PAC) that destines at posterior nuclear shelf (PNS) forming small swallow fusiform caudal fusiform end (CFE) of the posterior acrosomal cap (PAC) (figs. 16-18).

Moreover, the crown segment (CS) appears as club-shaped, which is surrounded by crown dense diadems (CDD) that imprecate along the anterior acrosomal cap as a chain of dense diadems (ChD) (fig. 18).

4. Discussion

Present results describe stages that will help us to conceive the mechanism in which MMs control the nuclear elongation, formation of the implantation fossa at which the connecting piece of the tail formed. Also, the MMs and microfilaments reorient their direction in the anterior and posterior halves of the nucleus, in parallel to its long axis and/or to horizontally surround the posterior nuclear region.

Similar longitudinal MMs reported, just behind the distal extremity of the acrosomal cap in bat spermatids [41]. In mice spermatids, a cone-shaped bundle of MMs encases the nuclear posterior pole [10]. However, longitudinal and circular bundles of MMs are described in marsupial spermatids [42].

Also, the Manchette was described in mammalian spermatids as a transient sleeve-like organelle that appears as longitudinal microtubular elements, displaced laterally parallel to the long axis of the spermatid during early spermiogenesis, encircling the caudal pole of the nucleus, and then extending back to connect the sperm head with neck region [8, 9, 10, 27]. However, Fouquet *et al.* [11] follow the localization of dynactin complex, associated with dynein of microtubules. They concluded that in round spermatid it appears near the centrosome and at the Golgi apparatus, and in elongated spermatids, it was arranged along microtubules of the manchette and at their attachment sites to the nuclear envelope, while disappeared in the testicular spermatozoa. They added that the various localizations of the dynactin might contribute to the activities of the centrosome and of the Golgi apparatus, as well as the shaping of the nucleus by MMs.

The present author believes that the solitary microtubules observed in round spermatid de-polymerized then re-polymerized in longitudinal and circular bundles tangential to the caudal half of the nucleus. These bundles may generate intrinsic compression strength perpendicular to their long axis induced nuclear prolongation. It is worth mentioning that, the generated strength is gently forces the peri-nuclear space that surround the posterior half of the nucleus to accelerate the anterior and posterior nuclear elongation without disturbing the chromatin condensation.

The Manchette acts as a track for the transport of cellular components between the nucleus and distal cytoplasmic regions of the elongating spermatid [12, 38, 43].

Present results showed the formation of oval or rounded acrosomal granule and a thin layer of acrosomal vesicle that capping at the anterior proximity of the nucleus. The former develops into a novel acrosomal structures include crown, anterior and posterior segments that extend posteriorly to capping two-thirds of the nuclear envelope. Similar, acrosomal vesicle was described in the early spermatid of the musk shrew [45, 46], marmoset monkey [6, 7, 29], also, a membrane-bounded acrosomal granule in New Zealand white rabbits [44]. However, several proacrosomal vesicles were observed in marsupial and bat early spermatids [41, 42]. These acrosomal vesicles became flattened and extend to form an acrosomal cap [6, 7, 29. 41, 42, 44, 46] that oriented in a longitudinal direction and capping the elongated nuclei. In the marsupials, the acrosomal cap is re-oriented to the horizontal plane and protrudes as a projecting U-shape [42], while in the musk shrew, a flat fan-like acrosome appears in the late spermatid [45, 46].

On the other hand, a new triple acrosomal structure is observed in the present spermatid that formed of anterior acrosomal rod and two short lateral acrosomal arms, such structures are not described in mammalian spermatids before. In addition, the club-shaped crown segment is surrounded by imprecated chain of dense diadems that extends along the anterior acrosomal cap. Such structures could be specific for the present species, hence it is not observed, yet, in other rodents.

As regards to the posterior nuclear shelf appeared after the formation of the anterior one. Most publications described only one acrosomal shelf as a ring structure at the posterior end of the nucleus of the developing spermatids of rabbits, mice, monkeys, bats and in fat-tailed jerbil, *P. duprasi* [11, 38, 41, 44].

Conclusion

From the present study, it is strongly concluded that the differentiation of sperm head of Egyptian lesser jerboa, *Jaculus jaculus* is controlled by cytoskeletal elements, especially manchette microtubules which master the shaping of the spermatid nucleus during its successive development. Present results showed the

formation of oval or rounded acrosomal granule and a thin layer of acrosomal vesicle that capping at the anterior proximity of the nucleus. The former develops into a novel acrosomal structures include crown, anterior and posterior segments that extend posteriorly to capping the nuclear envelope. A new triple acrosomal structure is observed in the present spermatid that formed of anterior acrosomal rod and two short lateral acrosomal arms, such structures are not described in mammalian spermatids before. In addition, the club-shaped crown segment is surrounded by imprecated chain of dense diadems that extends along the anterior acrosomal cap. Such structures could be specific for the present species, hence it is not observed, yet, in other rodents.

It is worth mentioning that a new translucent perinuclear space appears at the posterior end of the nucleus. It receives a gently generated compression induced by the cytoskeletal elements to produce the anterior and posterior nuclear elongation and to protect DNA during nuclear condensation.



Figure 1: Electronmicrograph showed epithelium of a seminiferous tubule. Note that germinal epithelium (GE) and Sertoli cells (Ser) are situated on a special basement membrane (BM), while other spermatogenic cells can be seen. Primary spermatocytes (PSp) contain large nucleus (N) with accumulating chromatin (Ch), early spermatids (ES) has small nucleus with fine chromatin granules and their anterior side capping with an acrosomal granule (AG) and acrosomal vesicle (AV). Bar: 2µm, X 3000. **Figure2:** Electronmicrograph showed early spermatid contains round nucleus (N) filled with aggregations of fine chromatin granules (Ch), the cytoplasm illustrated numerous mitochondria (M), and few segments of RER. Bar: 1µm, X 2000. **Figure 3:** Electronmicrograph showed active spermatid in which Golgi apparatus (G) starts to produce secretory vesicles (SV) that will be aggregates to form acrosome. Bar: 500nm, X 10000. **Figure4:** Magnified portion from figure 3 showed rough endoplasmic reticulum (RER), units of numerous microtubules (Mt) and microfilaments (Mf) that run in circular patterns. Bar: 400nm, X 2000.



Figure 5: Electronmicrograph showed two stages of active spermatids (ES). Right: shallow and early implantation fossa (EIP) formed in the posterior side of the nucleus (PoS) in which proximal centriole will be enclosed. The anterior side of the nucleus (AS) showed early acrosomal granule and acrosomal cap surround its anterior half and stop on a region at the equatorial level, nuclear shelf (NS) and separated from the nuclear envelope by a narrow subacrosomal space (SAS). Left: Active spermatids represent a later stage next to that observed in the right. Note deeper implantation fossa (IP) at the posterior side of the nucleus (PoS), larger acrosomal granule (AG), acrosomal vesicle (AV), long segments of RER and numerous microtubules run in circular patterns (Mt). Bar 0.5 µm. X 7500. **Figure 6:** Electronmicrograph showed that the proximal centriole (PC) is situated inside the implantation fossa (IF) that appears at the posterior ventro-dorsal side of the nucleus. Bar 1µm. X 4000.



Figure 7: Electronmicrograph of early spermatid just before nuclear elongation showed the formation of large unit of microtubule (MtU) that appears at the nuclear shelf (NS), in addition, numerous solitary microtubules run in a circular direction to surround the upper nuclear portion (Mt), its equatorial and sub-equatorial half of the nucleus that appear to surround the nucleus and run in circular pathways (MMs). These MMs will be re-orienting into longitudinal direction as shown in figures 8-13 in order to nuclear re-shaping. Bar: 500nm, X 10000. **Figure 8:** Electronmicrograph showed the formation of acrosomal cap (AC), subacrosomal space (SAS), anterior (ANP) and posterior nuclear protrusion (PNP), chromatin condensation, the presence of anterior (ANS) and posterior nuclear shelf ((PNS) at juxta-equatorial region, and implantation fossa (IF) at the posterior side of the nucleus, where the MMs are run parallel to its long axis and attach to both the MtU at the anterior (ANS) and posterior nuclear shelves (PNS), in addition, the presence of the posterior perinuclear space (PP) along one side of the posterior edge of the nucleus. Note the migration of cellular organelles at the posterior direction of the nucleus. M: Mitochondria. Bar 0.5 μm.



Figure 9: Electronmicrograph showed that the acrosome start to differentiate into four regions that are considered as new subdivisions in the acrosome. The former is a translucent terminal segment (TS), the remaining three portions appeared as a novel triangular acrosomal mass, which will be differentiates into the crown (SC), anterior (AAS) and posterior acrosomal segments (PAS) at which a long acrosomal structure capping the anterior two-third of the nuclear envelop "anterior acrosomal cap "(AAC)" and separating from it by a narrow subacrosomal space (SAS) as observed in figure 12. Moreover, there is a novel triple branched structure, which is formed of a terminal dense rod (TDR) and two lateral dense arms (LDA). Also, this figure clarifies the re-orientation of MMs and Mf into longitudinal or circular directions to surround the posterior half of the nucleus and the posterior perinuclear space (PP) that appears in the ventral side of the posterior nuclear protrusion (PNP), in addition, the presence of anterior (ANS) and posterior nuclear shelves (PNS), basal plate (BP). Note that the chromatin granules (Ch) are condensed at the peripheries of the nuclear envelop, the distal centriole (DC) is situated near the posterior ventro-lateral side of the nucleus, and the presence of few mitochondria (M). Bar: 500nm, X 12000.



Figure 10: Electronmicrograph showed successive nuclear elongation towards the anterior (ANP) and posterior sides of the nucleus (PNP), chromatin (Ch) condensation is markedly increase, the perinuclear space expands to surround the posterior two-third of the elongated nucleus. This space is bounded between the anterior (ANS) and posterior nuclear shelves (PNS). Also, there is a new smaller space appears on one side at juxta-equatorial nuclear region, pre-equatorial space (PeS). Moreover, the cytoskeleton is formed of MtU, MMs, in addition, solitary Mt and Mf appear to organize the nuclear elongation, the formation of the implantation fossa (IF). The MMs attach to the MtU at the equatorial region posterior to the anterior (ANS) and posterior nuclear shelves (PNS). The MMs still generate a compressive strength on the posterior perinuclear space (PP) causing more nuclear prolongation. Some microfilaments (Mf) appear at posterior end of the nucleus that may be involved in the formation of the implantation fossa (IF). Other cytoskeletal elements are distributed in the posterior regions of the developing spermatid. Note that the proximal centriole (PC) in enclosed in the implantation fossa (IF). As regards to the acrosomal regions, the present thin-section showed the lateral dense arm (LDA), posterior acrosomal segment (PAS), and the acrosomal cap (AC). Bar: 500nm, X 15000.



Figure 11: Electronmicrograph showed that the compressive strength on the posterior perinuclear space (PP) causing nuclear constriction (NC) and increase the anterior (ANP) and posterior nuclear prolongation (PNP), the ANP is covered by an anterior acrosomal cap (AAC) and separated from it by a narrow subacrosomal space (SAS). The triple branched structure of the acrosome appears including the terminal dense rod (TDR), and two lateral dense arms (LDA). In the present spermatid, the posterior side of the nucleus showed posterior space (PS) that fused with the posterior perinuclear space (PP), both of them is supported with longitudinal MMs that attach to the large units of microtubule (MtU) and run posteriorly to the connecting piece in the sperm neck (SN). Note that the presence of pre-equatorial space as it seen in the previous stage. Bar: 500nm, X 12000.



Figure 12: Electronmicrograph showed advanced stage of spermatid with three different regions of acrosomal segments, anterior (ANP) and posterior nuclear elongation (PNP), chromatin condensation and nuclear constriction (NC). The precruser portions of the acrosome explained in figure 9 are differentiated in the present stage, into crown, anterior, posterior acrosomal segments (PAS) that attach with the anterior acrosomal cap (AAC), which extends to the anterior nuclear shelf (ANS). The later is separated from the tip of the conical nuclear prolongation by a narrow subacrosomal space (SAS). In the terminal segment, there is triple structure that formed of terminal dense rod (TDR), and two lateral dense arms (LDA). Moreover, more solitary microtubules appear at the posterior end of the nucleus. They arranged in a circular pathway at the posterior end of the nucleus to support the attachment between sperm head (SH) and neck (SN). Note that the presence of pre-equatorial space (Pes) and posterior peri-nuclear space (PP). Concerning with the cytoskeleton, the described elements (Mt and Mf), of the previous stage, control the migration of subcellular organelles towards the posterior end of the present spermatid at the sperm neck (SN). Bar: 500nm, X 12000.



Figure 13: Electronmicrograph showed continuous chromatin condensation and nuclear prolongation to the anterior (ANP) and posterior (PNP) directions. The space of this cone is in a direct attachment with the subacrosomal space (SAS) on both sides. Also, the PAS is attached with a long anterior acrosomal cap (AAC) that capping the anterior conical protrusion of the nucleus and this cap extends along the two third of the nucleus then terminates at the anterior nuclear shelf (ANS) at the equatorial level of the sperm head. In this point, there is a small pre-equatorial space (PeS) that appears on one side at juxtanuclear region. In addition to the MMs, the cytoskeleton in the present stage showed other solitary microtubules that are situated in the posterior side of the nucleus, where they are arranged in longitudinal and transverse directions for tight attachment between head (SH) and neck (SN) of the developing sperm. However, this stage proved that the posterior perinuclear space starts to be decreased and consequently, the nuclear prolongation is continued in a slow rate, meanwhile, the chromatin condensation will be continued in the next stages. The posterior end of the nucleus illustrates the presence of implantation fossa (IF) that lodge the proximal centrille (PC) and the connecting piece in the sperm neck (SN) showed some organelles that include the distal centriole (DC) situated in the implantation fossa (IF), chromatoid body (CB), MMs, solitary microtubules (Mt) that run in longitudinal and transverse directions, mitochondrion (M). Bar: 0.5 µm, X 7500.



Figure14: Electronmicrograph showed sagittal section that clarifies the developing sperm head (SH), neck (SN) and middle piece (MP). This stage is the most important because it is clarified that chromatin condensation ends, the ventral acrosomal cap (VAC) is shorter than the dorsal one (DAC) and both of them stopped at the ventral (VNS) and dorsal nuclear shelves (DNS), respectively. Note the presence of nuclear constriction (NC) and nuclear canal (NCa), posterior ventro-dorsal protrusion of the nucleus (VDP) that previously described in figure 6, the other portions of the acrosome can be observed, which include terminal (TS), crown (CS), anterior (AAS) and posterior acrosomal segments (PAS), subacrosomal cone (SC), subacrosomal space (SAS). The crown segment gives an acrosomal crown (ACr) that formed of a thin chain of dense diadems (DD) which surround the crown segment and extend along the dorsal acrosomal cap (DAC). Bar: 0.5 µm, X 6000.



Figure 15: Electronmicrograph showed anterior longitudinal section of late spermatid. This ultrathin section gives us a complementary imagination about new structures that appeared on both right and left sides of this spermatid and numerous nuclear canals (NCa) that appear in the middle region of the nucleus after completion of chromatin condensation. The present stage proved that the crown segment gives acrosomal crown (ACr) that covered with a thin chain (ChD) formed of numerous small dense diadems (DD) which surround the crown segment and extend posteriorly along the anterior acrosomal cap (AAC) that ends at the anterior nuclear shelf (ANS). Moreover, additional posterior acrosomal cap originates at the anterior nuclear shelf and extends posteriorly as a thin layer to the posterior nuclear shelf (PNS). Hence, the acrosomal cap differentiates into a long anterior (AAC) and short posterior acrosomal cap (PAC) that separated from the posterior region of the nuclear envelope by a narrow space called posterior subacrosomal spaces (PSS). Bar: 0.5 µm, X 7500.

Figures 16 & 17: Two reconstructed electronmicrograph showed anterior longitudinal section in very late spermatid. This stage illustrated that both lateral dense arms (LDA) are developed, capping both the crown (CS) and anterior segments of the acrosome (AAS), while the posterior acrosomal segment is separated from the anterior by an one electron translucent region (ETR), in addition, the posterior acrosomal cap (PAC) complete its trip to most posterior end of the nucleus at the posterior nuclear shelf (PNS) and separating from the nucleus by additional called space posterior narrow subacrosomal space (PSS). Worth mentioning, the cytoskeleton in the present stage is formed of solitary microtubules (Mts) and microfilaments (Mf). The former are distributed along the anterior and posterior subacrosomal spaces and some Mts are observed in the outer side of the acrosomal cap and near the posterior nuclear shelf. Bar: 250nm, X 15000.





Figure 18: Electronmicrograph showed three advanced stages of late spermatids. This stage gives us the final configuration of the different acrosomal regions. The crown segment (CS) takes club-shape, which is surrounded by Crown dense diadems (CDD) that imprecate along the anterior acrosomal cap as a chain of dense diadems (ChD). The anterior acrosomal segment (AAS) have trapezoid shape and fused with the posterior acrosomal segment (PAS) as a dense region, the caudal end of the anterior acrosomal cap enlarges as fusiform structure (PFE) at the anterior nuclear shelf (ANS), also the posterior acrosomal cap (PAC) terminates as smaller fusiform structure (CFE) at the posterior nuclear shelf (PNS). The subacrosomal spaces include subacrosomal cone at the tip of the nuclear prolongation, Anterior (ASS) and posterior subacrosomal spaces (PSS). Bar: 0.5 µm, X 7500.



Figure 19: Electronmicrograph showed longitudinal section in late spermatid through the sperm head, neck, and middle piece. This stage illustrated the structures that appeared at the posterior end of the nucleus. Note that the implantation fossa lodges the proximal centriole and the distal one is situated perpendicular to it. Elements of cytoskeleton consists of longitudinal MMs, or solitary microtubules, in addition numerous microfilaments that run in circular pathways to support the connections between the sperm head and caudal structures. Bar: 500nm, X 10000. **Figures 20:** Low magnification electronmicrograph showed developed spermatid. The sperm head () provided with acrosome, elongated nucleus that ends at the nuclear fossa (IF), neck region include the both the proximal (PC) and distal centriole (DC), the middle piece include the axial filament (AX) surrounded by mitochondrial sheath, the principal piece appears posterior to the end of the middle piece (MP) at the annulus (An). Bar: 0.5μ m, X 3000.

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Abbreviations

| AAC | Anterior acrosomal cap | MP | Middle piece |
|-------|--|-------|--|
| AAS | Anterior acrosomal segment | MS | Mitochondrial sheath |
| Ac | Acrosomal Cap | Mt | Microtubule(s) |
| ACr | Acrosomal crown | MtU | Microtubule unit |
| AG | Acrosomal granule | Ν | Nucleus |
| An | Annulus | NC | Nuclear constriction |
| AS | Anterior side of the nucleus | NCa | Nuclear canal(s) |
| ANP | Anterior nuclear protrusion | NP | Nuclear protrusion |
| ANS | Anterior nuclear shelf | NS | Nuclear shelf |
| ASS | Anterior subacrosomal space | PAC | Posterior limb of acrosomal cap |
| AV | Acrosomal vesicle | PAS | Posterior acrosomal segment |
| AX | Axoneme (Axial filament) | PC | Proximal centriole |
| BM | Basement membrane | PeS | Pre-equatorial space |
| ВР | Basal plate | PFE | Posterior fusiform end of the anterior acrosomal cap |
| СВ | Chromatoid body | PNP | Posterior nuclear protrusion |
| CDD | Crown dense diadems of the acrosome crown segments | PNS | Posterior nuclear shelf |
| CFE | Caudal fusiform End of posterior acrosomal cap | PP | Posterior perinuclear space |
| Ch(s) | Chromatin | PoS | Posterior side of the nucleus |
| ChD | Chain of lateral dense diadems covering the acrosomal segments | PS | Enlarged posterior nuclear space |
| СР | Connecting piece | PSp | Primary spermatocyte |
| CS | Crown segment of the acrosome | PSS | Posterior subacrosomal space |
| DAC | Dorsal acrosomal cap | RER | Rough endoplasmic reticulum |
| DC | Distal centriole | SAS | Subacrosomal space |
| DD | Dense diadems of the acrosome crown segments | SC | Subacrosomal cone |
| DNS | Dorsal nuclear shelf | Ser | Sertoli cell |
| DS | Dorsal side of the nucleus | SH | Sperm head |
| EIP | Early implantation fossa | SN | Sperm neck |
| ES | Early spermatid | SV(s) | Secretory vesicle |
| ETR | electron translucent region | тс | Terminal crest of acrosomal crown |
| G | Golgi body | TDR | Terminal dense rod |
| GE | Germinal epithelial cell | TS | Terminal segment of acrosome |
| IF | Implantation fossa | VAC | Ventral acrosomal cap |
| LDA | Lateral dense arm | VDP | Ventrodorsal protrusion of the nucleus |
| м | Mitochondria | VNS | Ventral nuclear shelf |
| Mf | microfilaments | VS | Visicular end of lateral acrosomal arm |
| MMs | Manchette microtubules | | |