

## Proteins as the molecular markers of male fertility

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### Abstract

Proteins play a key role in many functions such as metabolic activity, differentiation, as cargos and cell fate regulators. It is necessary to know about the markers involved in male fertility in order to develop remedies for the treatment of male infertility. But, the role of the proteins is not limited to particular function in the biological systems. Some of the proteins act as ion channels such as catsper and proteins like Nanos acts as a translational repressor in germ cells and expressed in prenatal period whose role in male fertility is uncertain. Rbm5 is a pre mRNA splicing factor necessary for sperm differentiation whose loss of function results deficit in sperm production. DEFBI14 is a beta defensin family protein necessary for sperm motility in LPS challenged mice where as TEX 101 is a plasma membrane specific germ cell protein whose function is not clearly known u to now. Gpr56 is another adhesion protein whose null mutation leads to arrest of production of pups in rats. Amyloid precursor protein role in Alzheimer's disease is already known but it plays an important role in male fertility also but its function is uncertain and has to be considered while targeting APP during the treatment of Alzheimer's disease. The study on amyloid precursor protein in male fertility is a novel thing but requires further study in correlation to alzheimer's disease.

**Keywords:** Nanos2, RBM5, Amyloid precursor protein, DYNLT1.

### 1. Introduction

Nanos is a highly conserved gene in *D.melanogaster* and acts as a RNA binding protein (Mosquera, Forristall, Zhou, King, 1993; Kobayashi, Miyazaki, Natori, & Nozawa, 1991; Pilon, & Weisblat, 1997; Subramaniam, & Seydoux, 1999; Mochizuki, Sano, & Kobayashi, 2000; Kopranner, Thisse, Thisse, & Raz, 2001; Tsuda, Sasaoka, Kiso, Abe, Haraguchi, Kobayashi, & Saga, 2003) It plays several roles in *drosophila* but the role of it in fertility in humans has yet to be studied. Three types of Nanos gene such as Nanos 1, Nanos2 and Nanos3 (Haraguchi, Tsuda, Kitajima, Sasaoka, Nomur'za-Kitabayashid, Kurokawa, & Saga, 2003) are majorly studied and in which Nanos 3 involves in migration of primordial germ cells to gonads (Tsuda, 2013) and its presence in spermatogonia is necessary for differentiation during spermatogenesis.

Some of the proteins like ser/thr protein phosphatases and kinases are well known for regulating metabolism by phosphorylation and dephosphorylation events but their role in male fertility was not well known. However, the Protein phosphatase 4 is one of the ser/ thr phosphatases found to be necessary for preventing errors in the genetic exchange. It plays key role in maintenance of synaptonemal complex and generation of programmed ds breaks which is necessary for cross over events (Sato-Carlton, Li, Crawley, Testori, Martinez-Perez, Sugimoto, & Carlton, 2014) Ser/thr protein kinase causes phosphorylation of H1, H2, H2AX and H3 (Spiridonov, Wong, Zervas, Starost, Pack, Paweletz, & Johnson, 2005) and is required for chromatin remodelling in mitosis and meiosis. It also plays a role in DNA compaction also.

Ion channel proteins like catsper an anion transporter known for maintaining the ion fluxes in the cells and also involves in maintenance of membrane potential and ion balance inside the cell and required for maintaining sperm motility through  $ca^{2+}$  ion fluxes. It plays an important role in male fertility through regulating sperm motility. Slc26

is an anion transporter necessary for transport of monovalent and divalent anions like chloride (Cl<sup>-</sup>), sulphate (SO<sub>4</sub><sup>2-</sup>), iodide (I<sup>-</sup>) and bicarbonate (HCO<sub>3</sub><sup>-</sup>) (Mount & Romero, 2004) and also involved in differentiation in mouse.

Proteins that counteract oxidative stress in seminal plasma include SOD, catalase and GPx where as non enzyme antioxidants like α-tocopherol, urate (Kobayashi, Yamada, Asaoka, & Kitamura, 1996; Padron, Lynne, Brackett, Thomas, Sharma, Agarwal, 1997; Ollero, Gil-Guzman, & Lopez, 2001; Fujii, Iuchi, Matsuki, & Ishii, 2003), naphthoquinone and HCO<sub>3</sub><sup>-</sup> were previously known for protection against oxidative stress but now these are well known for their role in sperm functional parameters also.

Another protein GPR 56 was one of the adhesion G-protein coupled receptor that functions in cell adhesion through G-protein coupled signalling (Yona, Lin, Siu, Gordon, & Stacey, 2008). First report of its involvement in male fertility was known through testis cord remodelling by Guangchun Chen et al., .

DYNLT1 is a 14 Kda protein occupying the L1 inner arm of cytoplasmic and flagellar dynein components (Harrison, Olds-Clarke, & King, 1998; King, Dillman, Benashski, Lye, Patel-King, & Pfister, 1996). It is also present in oocytes, sperm tails (Campbell, Cooper, Dessing, Yates, & Buder, 1998; O'Neill, & Artzt, 1995) and Golgi complexes (Tai, Chuang, & Sung, 1998). It also functions in as dynein independent manner and is a cell fate regulator in neural progenitor cells (Dedesma, Chuang, Alfinito, & Sung, 2006).

There are many proteins that plays a role in male fertility but this review is concise up to some of the major key proteins involved in male fertility.

## **2. Proteins involved in male fertility through phosphorylation and dephosphorylation events:**

Serine /threonine protein phosphatase PP4 homolog PPH 4 was normally found in budding yeast and its presence in *C.elegans* and *D.melanogaster* is proved to be necessary for DNA ds break formation initiation and cross over along with synapsis independent pairing and prevention of non homologous pairing in autosomes during the synapsis . But it doesn't prevent the synapsis dependent pairing of homologous chromosomes. The homology of these proteins is about 92% at aminoacid level in humans to that of mouse. The protein is necessary for formation of chiasma, thus recombination without affecting the loading of recombinant proteins Rad 51. Ds break forming capability with respect to phosphatase enzyme is an age dependent factor and found to be involved in the meiosis, and in division that occur in germ cells (Sato-Carlton, Li, Crawley, Testori, Martinez-Perez, Sugimoto, & Carlton, 2014).

PP4 is necessary for the conversion of foci in to cross overs through COSA-1 (Sato-Carlton et al., 2014) and dephosphorylates SUN protein which is required for synapsis independent pairing (Sato-Carlton et al., 2014) in the yeast .

SSTK is a small protein kinase found on chromosome 8 and distributed with high similarity in mammals and is expressed in almost all the tissues. Phylogenetic analysis showed moderate similarity of SSTK to the testis specific ser/thr kinases TSSK1, TSSK2 and TSSK3, MAP kinase/ microtubule affinity- regulating kinases MARK and MARK4, and the ELKL motif kinase EMK1 (Spiridonov et al., 2005).

This protein consists of N and C lobes of a protein kinase domain containing catalytic and ATP binding domains in which catalytic residues include K41, E60, D135, N140, D154, and T170, a glycine rich motif in phosphate binding loop and the conserved sequence DFG in the active site. SSTK consists of tyr phosphorylation domains which are similar to phosphorylation inhibitory domains of cyclin dependent kinases Cdc2 and Cdk2 and TTY sequence similar to T- X-Y phosphorylation motif found in the activation loop of MAP kinase (Spiridonov et al., 2005).

SSTK even though associate with HSP90-1, HSP70, and HSP70-1 doesn't phosphorylate them but necessary for proper maintenance of structure of sperm head, sperm motility and DNA condensation in sperm head as it phosphorylates H1, H2A, H2AX and H3 but not the H2B, H4 and TP1 (Spiridonov et al., 2005).

Another protein TSSK found to have kinase activity is necessary for male fertility as they are localised in the spermatids which is HSP90 dependent. It was found that this kinase associates with HSP90 as it results in reduced expression after incubation with HSP90 inhibitors and kinases. TSSK-1,2,4,6 phosphorylates H2A histone where as TSSK3 doesn't show any kinase activity. TSSK -1,2,6 reduced in expression when treated with HSP90 inhibitors indicating that HSP90 is required for their maintainance of half life and also catalytic activity of TSSK-4,6. TSSK 2 and 6 undergoes ubiquitination directly when inhibited with inhibitors of HSP90 and undergoes proteosomal degradation but without change in their mRNA levels (Jha, Coleman, Wong, Salicioni, Howcroft, & Johnson, 2013).

### 3. Ion channels and transporters as male fertility factors

Another protein catsper localised in sperm tail is necessary for sperm motility, calcium influx and fertilisation in mice . Catsper 1 was found to be one of the four important proteins of calcium channels necessary for male fertility in mice (Avenarius, Hildebrand, Zhang, Meyer, Smith, Kahrizi, Najmabadi, & Smith, 2009) and also in humans. In case of sperm motility  $ca^{2+}$  influx is necessary for hyper activation in males and in female genital tract the  $ca^{2+}$  influx is necessary for capacitating and high motility penetration of sperm in to oocytes.

Slc26A8 is an testis anion transporter expressed in sperm and necessary for sperm motility and required for fertilisation potential in males. It is an anion transporter and doesn't involve in the maturation of gonads of mice. Slc26A8 is found to be localised in the annulus that connects the midpeice to principal piece of the flagella. It was shown that the ability of consumption of ATP was reduced due to defects in mitochondrial sheath even though the motor protein expression in flagella is normal and was found that null mutation of the protein leads to compromise in normal maturation of sperm and capacitation in the mice and humans (Toure, Lhuillier, Gossen, Kuil, Lhote, Je'gou, Escalier, & Gacon, 2007).

### 4. Is oxidative stress preventing enzymes and structural maintenance proteins are necessary for sperm function

Superoxide dismutase, catalase and glutathione peroxidase plays an important role in relation to oxidative stress. SOD level changes was associated with changes in sperm count where as catalase with sperm morphology and GPx expression levels was found to be not associated with any of the sperm parameters (Macanovic, Vucetic, Jankovic, Stancic, Buzadzic, Garalejic, Korac, Korac, & Otasevic, 2015)

There is no contributinal research of nuclear matrix proteins in relation to sperm parameters up to now. So, study of proteins of nuclear matrix sperm was found to be useful. Proteins identified in the sperm head include mostly chaperons, cytoskeletal proteins, peroxiredoxins, isomerases and other enzymes (Kichine, Falco, Barbara, Robaire, & Chan, 2013)

**Table 1:** Proteins role in male fertility, localisation and functions other than male fertility and their chromosomal distribution

Name of the protein	localisation	Role in male fertility	Functions other than male fertility	organism	Chromosomal distribution	Reference
Protein phosphatase 4 (PP4)	-	Meiotic chromosome dynamics	Dephosphorylation events during cell cycle and DNA damage response	Universal regulator	6	Sato-Carlton et al. (2014)
Catsper	Principal peice	Sperm motility	Not known	Mice, humans	11	Rahman et al.(2014)
Superoxide dismutase(SOD), catalase	Seminal plasma	Sperm count (SOD)  Sperm morphology (catalase)	Free radical scavengers	universal	6 (MnSOD)  11 (catalase) in humans	Borgstahl et al. (1996)
Amyloid precursor protein (APP)	Tail and head region	Sperm motility and sperm-oocyte interaction	Receptor like and adhesive properties in nervous system	ubiquitously	21	Silva et al. (2015)
SSTK(ser/thr kinase)	Heads of elongated spermatids	chromatin condensation, reconstruction	Associates with HSP 90 and HSP70	Rat, dog, mouse, human,		Spiridonov et al. (2005)

		of the sperm cytoplasm, acrosome formation, and development of the flagellar apparatus.		cow	19	
GPR56 an adhesion G-protein coupled receptors	Sertoli cells	Development of male gonads	Inhibition of melanoma progression, brain function	mammals	16	Chen et al. (2010)
TEX101	Plasma membrane of germ cells	Spermiogenesis and fertilization	Binds to uPA/uPAR complex and mediates the effects	mice	19	Schiza et al. (2014)
B-defensin 114	Epididymis and saliva, gingival keratinocytes	Protection against LPS loss of fertility	Anti inflammatory	humans	6p 21	Yu et al. (2013)
Testis anion transporter Slc26a8	Spermatocytes and spermatids	Germ cell function and differentiation	Transport sulphate in a chloride dependent manner	bacteria, yeast, plants, nematodes and mammals	9p24.2	Mount & Romero (2004)
RBM5	Spermatogonia, round spermatids and spermatocytes	pre-mRNA splicing regulator in round spermatids	Apoptosis, lung histology	Humans and mouse	9	O'Bryan et al. (2013)
Nanos2	germ cells.	translational repressor necessary for germ-cell development	Meiosis suppression	Widely spread	19	Kusz et al. (2009)
DYNLT1	head, tail and mid piece	Spermatogonial cell division and differentiation	Protein trafficking, membrane vesiculation, cell cycle regulation, and stem cell differentiation.	Mouse	4 and 17	Indu et al. (2015)

## 5. Other proteins involved in male fertility

Amyloid precursor protein is normally known for its activity in Alzheimer's disease but its role in male fertility was not known. This protein was first identified in testis and studied for its interaction with testis. It is now known to interact with RANB9 protein and 37 proteins in which COPS5 has highest correlation coefficient where as CD81 and CD 99 with  $C=0.029$  and  $0.064$  respectively (Silva, Yoon, Domingues, Guimarães, Goltsev, Silva, Mendes, & Fardilha, 2015).

GPR56 is an adhesion protein necessary for male fertility. It was found to be expressed in testis cords, PM cells, localised in Sertoli cells and germ cells but found to be absent in interstitial cells. GPR56 is highly expressed in

Sertoli cells and spermatogonial cells with reduced expression in PM cells. Production of progeny with defective testis is seen with GPr56 null mice as the spermatic cords are disrupted and scattered instead of forming tubular structures. There is also basement membrane disruption in the testis with no alterations in FSH, LH and testosterone probably showing no effect on these hormones (Chen, Yang, Begum, & Xu, 2010).

TEX 101, a testicular germ cell specific protein is found to be located in the plasma membrane of germ cells. It does not affect the mating but secreted as one of the GPI anchored proteins by TACE in to the seminal fluid. However the mechanism of action of protein was not clearly understood (Schiza, Jarvi, Diamandis, & Drabovich, 2014).

DEFB114 is the  $\beta$ - defensin highly expressed in the caput and corpus regions of epididymis. It shares cysteine conserved regions and cysteine pairing regions (Cys10–Cys24, Cys3–Cys31, Cys14–Cys32) with  $\beta$ - defensin members. It is involved in maintenance of male fertility through preserving sperm motility in LPS challenged mice by neutralising it in a dose dependent manner (Dong, Liu, Xin, Shi, Sun, Zhang, Lin, & Diao, 2013).

Rbm5 is the pre – mRNA splicing factor present in the nucleus and cytoplasm of spermatocytes and round spermatids. R263 was found to be the highly conserved region in the Rbm5 and loss of function allele mutation leads to arrest of pups reduction as it is necessary for RNA binding in the RNA recognition motif and change in  $\beta$ -strand structure necessary for the binding of protein. This mutation doesn't effects the fertility in females. The protein is necessary for sperm differentiation and the mutation leads to in azoospermia due to arrest at stage 8 of haploid sperm development which is independent on hormones FSH, LH and testosterone. The loss of function in this protein leads to testis atrophy as it is highly expressed in testis compared to other tissues (O'Bryan, Clark, McLaughlin, D'Sylva, O'Donnell, Jacqueline, Sutherland, O'Connor, Whittle, Goodnow, Ormandy, & Jamsai, 2013).

St5, is involved in MAP Kinase pathway which is necessary for the cell growth, post testicular maturation and fertilisation and necessary to be explained in detail. The splicing by Rnm5 produce 61 transcripts in which 126 Kda fragment was involved in regulating the MAP kinase pathway and found to contain high amounts of p-ERK1/2 than ERK1/2. Regulation in sperm differentiation by protein in wild type than mutant indicates its regulation in tumor growth (Kusz, Tomczyk, Sajek, Spik, Latos-Bielenska, Jedrzejczak, Pawelczyk, & Jaruzelska, 2009).

Nanos 2 is the protein encoded by the gene Nanos that acts as translational repressor in germ cells and expressed in the cytoplasm of germ cells of seminiferous tubules of testis during prenatal period. The molecular weight of protein was found to be more than 15Kda. It is highly expressed in the peri tubular cells and its localisation is different during prenatal and adult stage. H68Q and H109H are the mutations localised in the zinc finger domain commonly found in Nanos 2. It form complexes with the proteins like (DAZ and BOL, PUMILIO2, NANOS1, respectively) but its role in male fertility is not clearly understood (Kusz et al, 2009).

DYNLT1 is a gene that found associated with t-complex of testis. It is localised in sperm head, mid piece and sperm tail and in infertile men. The localisation was restricted to head and mid piece only. It is known to cause male sterility in loss of function and restoration of the fertility with BAC construct of the protein in mice, fly and humans. Its molecular weight was found to be 14Kda. It is a component of microtubular network and over expression of HSP 90 along with DYNLT1 leads to phosphorylation of Thr 94 which plays a role in cell division. DYNLT1 is found to be involved in sperm division and differentiation (Indu, Sekhar, Sengottaiyan, Kumar, Pillai, Laloraya, Kumar, 2015).

## 6. Discussion

Proteins plays a key role in the biological systems as enzymes, transcription factors, antibodies, cytokines etc., so, instead of focussing on the sperm morphology, and sperm parameters concentrating on molecules involved in maintaining genome integrity proves to be helpful in understanding their role in male fertility. Some of the proteins are ion channels maintaining  $Ca^{2+}$  influx during motility and capacitation. Some of them are useful for DNA compaction as they are involved in phosphorylation of histones in sperm. Absence of some proteins leads to loss of fertility as they are responsible for sperm division and differentiation like DYNLT1 and found to be involved in cargo binding, lymphocyte division, vesicular transport and human embryo implantation.

Most of the proteins which are discussed here are responsible for sperm motility and male fertility but not focussed in the female. Amyloid precursor protein in patients with alzheimers disease should be studied for interactions related to fertility. Some of the proteins acts as antioxidants preventing oxidative stress also play a role in sperm count and morphology where as defensin DEFB114 found to preserve fertility in lipopolysaccharide mice which seems its involvement in immune reactions may be as pattern recognition molecule .

Some of the proteins like Rbm5 are known to interact with other proteins and involved in prevention of continous cell division in germ cells and also acts as splicing factor of pre mRNAs of apoptotic proteins such as caspase, FAS receptor and C- FLIP. It indicates its control on cell division preventing the tumorigenic growth in tissues apart from male fertility.

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