LRGUK-1 Is Required for Basal Body and Manchette Function during Spermatogenesis and Male Fertility

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Abstract

Male infertility affects at least 5% of reproductive age males. The most common pathology is a complex presentation of decreased sperm output and abnormal sperm shape and motility referred to as oligoasthenoteratospermia (OAT). For the majority of OAT men a precise diagnosis cannot be provided. Here we demonstrate that leucine-rich repeats and guanylate kinase-domain containing isoform 1 (LRGUK-1) is required for multiple aspects of sperm assembly, including acrosome attachment, sperm head shaping and the initiation of the axoneme growth to form the core of the sperm tail. Specifically, LRGUK-1 is required for basal body attachment to the plasma membrane, the appropriate formation of the sub-distal appendages, the extension of axoneme microtubules and for microtubule movement and organisation within the manchette. Manchette dysfunction leads to abnormal sperm head shaping. Several of these functions may be achieved in association with the LRGUK-1 binding partner HOOK2. Collectively, these data establish LRGUK-1 as a major determinant of microtubule structure within the male germ line.

Author Summary

Male infertility affects one in six couples in western societies and approximately half of these are the result of male factor disorders. The most common clinical presentation for male infertility is a complex mixture of abnormal sperm output, shape and motility referred to as oligoasthenoteratozoospermia (OAT). In an effort to define an origin of OAT we have analysed a mouse model of leucine-rich repeats and guanylate kinase-domain containing isoform 1 (LRGUK-1) dysfunction. Herein we show that LRGUK dynamically redistributes during the process of haploid germ cell maturation (spermiogenesis) and that
LRGUK-1 function is required for multiple aspects of sperm centriole and tail development and sperm head shaping. Further, we have identified HOOK2 as a novel LRGUK-1 binding partner, thus raising the possibility that several aspects of LRGUK-1 function are achieved in partnership with HOOK2.

Introduction

Male infertility affects at least 5% men of reproductive age in the western societies [1]. Normal male fertility requires sufficient numbers of morphologically normal and motile sperm [2]. Oligoasthenoteratozoospermia (OAT), is the term used to describe semen containing low numbers of sperm with poor motility and abnormal shape, and is the most common clinical phenotype in human male infertility [3]. The underlying aetiology in the majority of men presenting with OAT is unknown, and as such, there remains an absence of specific treatments, an absence of knowledge of associated somatic pathologies, and potential consequences for offspring conceived with OAT sperm via assisted reproductive technologies (ART).

The morphological and motility aspects of OAT likely have their origins in spermiogenesis, the process wherein round haploid germ cells are transformed into highly polarized sperm with the potential for motility and fertility. This process takes approximately two weeks in the mouse and involves several thousand different gene products [4]. Three of the major aspects of spermiogenesis are acrosome development, head shaping, and growth of the sperm flagellum [5,6]. These events are critically reliant upon complex microtubule structures, including the manchette and the axoneme, and a highly orchestrated series of protein transport mechanisms [7]. The manchette is a transient microtubule structure, which encircles the spermatid nucleus during the initial steps of elongation, and has a role in sculpting the species-specific nucleus shape. Abnormalities in manchette structure result in dysmorphic sperm heads [8–10]. We note that dynamic redistribution of the actin cytoskeletal system is also required for normal manchette function [8].

The axoneme forms the core of the sperm tail. It is composed of nine microtubule doublets surrounding a central microtubule pair (9+2) [11]. In contrast to motile cilia on other cells, the sperm tail axoneme is sheathed by accessory structures required for the production of ATP and protection against shear forces [12]. These structures include the outer dense fibers, fibrous sheath and mitochondrial sheath. Sperm tail axonemal development begins in round spermatids with the maturation of the mother centriole into a basal body, followed by plasma membrane attachment and axoneme extension. Several studies suggest that the extension of the basal body into an axoneme is dependent on the intra-flagella transport (IFT) system [13]. We note, however, that while this process is well defined in primary cilia [14], its role in sperm tail biology requires more in-depth analysis. Defectives in germ cell axoneme formation leads to either an absence of a sperm tail or immotile sperm.

Interestingly, research is increasingly indicating continuity between processes governing the development of the manchette and the sperm tail [13,15]. This may explain the extremely common association between abnormal sperm head morphology and motility in infertile men and mice [16]. Indeed, pioneering research from the Kierszenbaum lab has demonstrated a protein transport highway wherein proteins processed by the Golgi apparatus are transferred to the surface of the sperm acrosome at the cranial pole of the sperm head in a microtubule-dependent manner, then onto a cytoskeletal scaffolding plate known as the axoplaxome that anchors the developing acrosome to the nucleus [17]. Subsequently, proteins then localise to the microtubules of the manchette and ultimately into the growing sperm tail. The latter part of the
highway has been termed intra-manchette transport and it appears to have mechanistic similarities with, and to interface with, the classical IFT pathway required for axoneme extension in most cilia types [13,18].

In an effort to understand these processes, we have undertaken a random mouse-mutagenesis screen to identify genes with critical roles in sperm formation. One of the genes identified encodes the previously uncharacterized gene leucine-rich repeats and guanylate kinase domain contain (Lrguk). Here we have shown that LRGUK isoform 1 (LRGUK-1) has a critical role in both sperm head shaping and basal body attachment to the plasma membrane, and in the early aspects of axoneme development. LRGUK is transported along the acrosome-acroplaxome-manchette-tail axis in a potential complex with the adaptor protein HOOK2 [19,20]. LRGUK-1 dysfunction leads to abnormal manchette formation and movement, and an absence of axoneme extension from the basal body. Collectively these data define LRGUK-1 as a crucial regulator of male germ cell basal body function, microtubule dynamics and fertility.

Results

Lrguk is required for male fertility

In order to identify critical regulators of male germ cell development and fertility, we undertook a random N-ethyl-N-nitrosourea (ENU) mouse mutagenesis screen [9,21]. The ‘Kaos’ mouse line was identified based on male-specific infertility and the presentation of chaotic spermatogenesis (see below). Mapping of the mutation and sequencing of candidate genes revealed a C to T substitution in exon 14 of Lrguk (Fig. 1A). Lrguk is predicted to encode 3 splice variants, Lrguk transcript 1 (Lrguk-1), -2 (Lrguk-2) and -3 (Lrguk-3). Lrguk-1 is the longest transcript and the only isoform affected by the mutation (Fig. 1A). The C→T mutation resulted in the conversion of an arginine (R) at position 528 to a premature termination codon (R528Stop, the LrgukKaos allele) and in turn the truncation of 293 residues from the C-terminal region of LRGUK-1 (Fig. 1B). Orthologues of LRGUK can be observed in many species, and sequence alignment of LRGUK-1 from multiple species revealed that R528 residue is conserved in all (Fig. 1C).

LrgukKaos/Kaos testis showed a 75% reduction in the level of Lrguk-1 expression (8–12 weeks-of-age) compared to LrgukWT/WT siblings (Fig. 1D) and a complete absence of both the native LRGUK-1 protein of 93 kDa and the predicted truncated LRGUK-1Kaos of 60kDa (Fig. 1E). We note that the antibody used in these experiments is directed against an epitope present in both the native and truncated forms of LRGUK-1, but C-terminal to LRGUK-2 and -3 sequences. These results suggested Lrguk-1Kaos mRNA was unstable. Quantitative PCR (qPCR) showed no evidence of a compensatory up-regulation of Lrguk-2 and Lrguk-3 mRNAs in the LrgukKaos/Kaos testis (Fig. 1F-G). Collectively, these data revealed that LrgukKaos/Kaos males were sterile as a consequence of a specific absence of the LRGUK-1 protein.

LrgukKaos/Kaos males sterility was characterised by oligoasthenoteratospermia

All LrgukKaos/Kaos males had chaotic and disorganised spermatogenesis and were sterile (n>50, aged 7–26 weeks), whereas LrgukWT/Kaos males and LrgukKaos/Kaos females had apparently normal fertility. LrgukKaos/Kaos males had a normal body weight (S1A Fig.) and were anatomically indistinguishable from wild type siblings. LrgukKaos/Kaos male sterility was characterized by a 13% reduction in testis weight (Fig. 2A) and an 81% reduction in daily sperm production (Fig. 2B). Collectively these data suggest that the majority of missing germ cells were from the haploid phase of spermatogenesis wherein they contributed relatively little to testis weight. A similar reduction (79%) in sperm content was observed in the epididymal sperm content (Fig. 2C). The loss of germ cells from the seminiferous epithelium via sloughing was indicated...
Fig 1. The Kaos mouse line phenotype was caused by a mutation in the Lrguk gene specifically affecting the LRGUK-1 isoform. (A) The Lrguk gene produces three transcripts in mouse. The Kaos mouse line contained a point mutation (C→T substitution, black arrowhead) located in exon 14 of Lrguk transcripts 1 (Lrguk-1). (B) The Kaos mutation resulted in the conversion of arginine 528 to a stop codon (R528*). LRR = leucine-rich region, LRRCT = leucine-rich repeat C-terminal domain, and GK = guanylate kinase-like domains. (C) A comparison of the relevant LRGUK region across species. The black box indicates the position of arginine 528. (D) Lrguk transcript 1 mRNA expression in LrgukKaos/Kaos (Kaos) versus wild type (WT) testes (n = 3 for each genotype). Expression was normalised against Ppia expression and shown as mean ± SEM. (E) LRGUK-1 content in LrgukKaos/Kaos (Kaos) versus wild type testes, 40μg of testis lysate per lane. Actin was used as a loading control. The predicted molecular weight of native LRGUK-1 was 93kDa and the
by the presence of immature germ cells in Lrguk<sup>Kaos/Kaos</sup> epididymis (Fig. 2F). In contrast, apoptosis levels were similar between genotypes (S1B–C Fig.). Of the elongate spermatids present in the seminiferous epithelium, virtually none appeared to contain a sperm tail (Fig. 2E), and of the very few sperm that were present in the epididymis, all had grossly misshapen heads and shortened tails (Fig. 2D) that displayed no capacity for motility. A closer examination of spermatid structure also revealed the presence of fragmented acrosome in early round spermatids (Fig. 2E and below).

A more quantitative stereological analysis of the dynamics of spermatogenesis confirmed the histological observations. Lrguk<sup>Kaos/Kaos</sup> adult testes contained normal numbers of supporting Sertoli cells, spermatocytes and early round spermatids (S1 Table). They contained, however, a 37% reduction (p < 0.05) in elongated spermatids (steps 13–16). In addition, we noted a pronounced increase in the number of elongated spermatids retained in the basal crypts of Sertoli cells in stage VII–XII tubules that was suggestive of a failure of the early phases of sperm release (spermiation) (S1D Fig.). Consistent with this interpretation, the levels of DNA double strand breaks, as determined by γH2AX staining, in the retained spermatid nuclei was notably elevated (Fig. 2G) in Lrguk<sup>Kaos/Kaos</sup> suggesting the initiation of degeneration [22].

Cumulatively, these data show that Lrguk<sup>Kaos/Kaos</sup> males were sterile as a consequence of germ cell sloughing and degeneration, abnormal sperm development and an inability of those sperm that were produced to ascend the female reproductive tract following mating.

**LRGUK-1 is testis-enriched and associated with the acrosome-acroplaxome-manchette-tail axis**

qPCR analysis revealed that Lrguk-1 mRNA was highly enriched in the testis compared to other adult tissues (Fig. 3A). An analysis of mouse testes taken at defined periods during the establishment of the first wave of spermatogenesis also revealed that Lrguk-1 mRNA was detectable at low levels from birth, up-regulated at day 14 coincident with the appearance of pachytene spermatocytes, then maximal from day 18 coincident with the appearance of haploid germ cells (Fig. 3B). This result was suggestive of Lrguk-1 being predominantly expressed in haploid germ cells. During spermatid development, and using an antibody that should detect all of LRGUK1–3, LRGUK protein was initially localised to a supra-nuclear region of round spermatids, and was particularly evident at the leading edge of the developing acrosome and acroplaxome (Fig. 3C). As maturation proceeded and nuclear elongation initiated, LRGUK moved distally to ultimately reside on the microtubules of the manchette (Fig. 3D and S2 Fig.). LRGUK was also evident in the sperm basal body and the sperm tail (Fig. 3E). These data, and the abnormal sperm head and tail morphology in Lrguk<sup>Kaos/Kaos</sup> germ cells, suggested that LRGUK-1 has a role in acrosome and tail biogenesis. We note that LRGUK protein was not detectable in our hands in spermatocytes. At present it is unknown if this was due to a lack of abundance or a translational delay as is often seen in spermiogenic genes [23].

Lrguk<sup>Kaos/Kaos</sup> germ cells showed reduced LRGUK immunolabelling compared to wild type cells (S2 Fig.), however, residual, presumably LRGUK-2 and/or LRGUK-3 staining could be seen within mutant spermatids (Fig. 3C–D and S2 Fig.). The possibility also remains that a small amount of truncated LRGUK-1 was detected by this antibody. Within these cells the distribution of residual LRGUK was notably perturbed. Specifically, the movement of LRGUK appeared to stall at the leading edge of the acrosome/acroplaxome complex (Fig. 3C–D and
Fig 2. *Lrguk* \(^{\text{Kaos/Kaos}}\) males are oligoasthenoteratospermic and sterile. (A) A comparison of testis weight from *Lrguk* \(^{\text{WT/WT}}\) (WT) and *Lrguk* \(^{\text{Kaos/Kaos}}\) (Kaos) adult mice (age = 8–12, n = 12–13 per genotype ±/− SEM). (B) Daily sperm production (DSP) by the testes from *Lrguk* \(^{\text{WT/WT}}\) and *Lrguk* \(^{\text{Kaos/Kaos}}\) adult mice (n = 7 per genotype ±/− SEM). (C) Total epididymal sperm content from *Lrguk* \(^{\text{WT/WT}}\) and *Lrguk* \(^{\text{Kaos/Kaos}}\) adult mice (n = 7 per genotype). (D) PAS stained testis sections from *Lrguk* \(^{\text{WT/WT}}\) and *Lrguk* \(^{\text{Kaos/Kaos}}\) mice. * denotes the absence of sperm tails in the lumen of *Lrguk* \(^{\text{Kaos/Kaos}}\) males; black arrowheads point to fragmented acrosomes. Scale bar = 50 μm. (E) Haematoxylin and eosin stained sperm smears from *Lrguk* \(^{\text{WT/WT}}\) and *Lrguk* \(^{\text{Kaos/Kaos}}\) animals. Scale bar = 10 μm. (F) Haematoxylin and eosin staining of an epididymis from *Lrguk* \(^{\text{WT/WT}}\) and *Lrguk* \(^{\text{Kaos/Kaos}}\) mice. The green arrow indicates sloughed of immature germ cells. Scale bar = 50 μm. (G) γH2AX foci (green) were visible in retained step I-II elongated spermatids from *Lrguk* \(^{\text{Kaos/Kaos}}\) mice but not *Lrguk* \(^{\text{WT/WT}}\) spermatids. Acrosomes were visualised by PNA (red) and nuclei were visualised with Dapi (blue). Scale bars = 10 μm. *p < 0.05, ****p < 0.0001.

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Fig 3. LRGUK-1 is highly expressed in the testis and localizes to the acrosome-acroplaxoneme-manchette-tail network. (A) Quantitative PCR analysis of Lrguk-1 mRNA expression in various tissues and during the establishment of the first wave of spermatogenesis n = 3. Expression was normalised against Ppia expression and shown as mean ± SEM. (B) The localization of LRGUK (red) in purified round spermatids.
S2 Fig.). LRGUK staining on the microtubules of the LrgukKaos/Kaos manchette was reduced and more variably distributed (Fig. 3D and S2 Fig.). These data are consistent with the acroplaxome operating as a loading dock for cargo proteins prior to loading onto the microtubules of the manchette [15,17] and the C-terminal 293 amino acids of LRGUK-1 having a role in this transition.

LRGUK is involved in sperm head formation

One of the features of the LrgukKaos/Kaos phenotype was the abnormal morphology of sperm heads. A detailed analysis of acrosome formation on periodic acid Schiff’s (PAS) stained sections revealed the presence of fragmented acrosomes in step 2–4 spermatids (Fig. 4A). The acrosome is formed by the sequential fusion of Golgi-derived pro-acrosomal vesicles to form a cap overlying the nucleus. The membrane overlying the acrosome is essential for the initial binding to and penetration through the oocyte complex [15,24,25]. Electron microscopy also revealed ~20% of acrosomes in elongated spermatids were detached from sperm nuclei (Fig. 4B). A close inspection of these cells indicated that both the acrosome and the acroplaxome were detaching from the nuclear membrane suggesting that LRGUK-1 has a role in establishing the integrity of the connection between the acroplaxome and the nuclear membrane. LrgukKaos/Kaos spermatids also contained abnormal manchettes (S2 Fig., Fig. 4D and Fig. 5). The manchette is a grass skirt-like structure that encircles the elongating spermatid nucleus. As indicated above, it has a role in both nuclear shaping and protein transport into the growing sperm tail via a process called intra-manchette transport [13,18]. In normal spermatids, the manchette contains a series of parallel microtubule bundles that extend from a perinuclear ring and lie in close proximity to, and are parallel with, the nuclear membrane into the distal cytoplasm (Fig. 4C and Fig. 5). An analysis of elongating LrgukKaos/Kaos spermatids, using α-tubulin as a microtubule marker, revealed that the manchettes formed at the correct time and that the perinuclear ring began to move caudally along the sperm head during spermiogenesis (Fig. 5). The microtubule bundles of the manchette skirt were, however, unevenly distributed and had a ‘raggedy’ appearance compared to wild type cells. Further, and based on both the light and electron microscopic images the distal movement of the manchette along the nucleus that normally occurs during elongation was also abnormal in LrgukKaos/Kaos spermatids (Fig. 5, S3 Fig.). In contrast the perinuclear ring of the manchette continued to constrict, as it would normally, thus leading to nuclear distortion (Fig. 4C, Fig. 5 and S2 Fig.).

Collectively these data illustrate a role for LRGUK-1 in acrosome attachment to the sperm head and in microtubule organisation within the manchette.

LRGUK binds to HOOK2 and functions in sperm tail axoneme extension

LRGUK-1 contains multiple domains with known roles in protein-protein interactions i.e. a guanylate kinase-like domain (GK), leucine rich repeats (LRRs) domains and a leucine rich repeat C-terminal domain (LRRCT) [26,27] (SMART entry IPR000483). In order to explore this potential, and to define pathways within which LRGUK-1 may be involved during haploid germ cell development, we performed a yeast two-hybrid screen to identify LRGUK-1 binding
partners. One of the binding partners identified was HOOK2 (Fig. 6). The identified Hook2 clone encoded the C-terminal 348 amino acids of HOOK2. Specific transfection of the full length HOOK2 with LRGUK-1 in a separate yeast two hybrid assay confirmed this interaction and the introduction of the R528Stop mutation into the Lrguk-1 sequence completely abolished binding to HOOK2 (Fig. 6A). An interaction between LRGUK-1 and endogenous HOOK2 was confirmed by co-immunoprecipitation, wherein full-length mouse Lrguk-1-GFP was transfected into HEK293T cells then the complex co-precipitated (Fig. 6B).

HOOK2 is a member of the HOOK family of proteins, which are adaptor-like proteins involved in loading cargos (including protein complexes and organelles) onto microtubules for transport [28]. Notably, HOOK2 has recently been found to function in the maintenance of centriole structure and primary cilia assembly [19]. Like sperm flagella, primary cilia contain a central axoneme, but rather than possessing a 9+2 microtubule structure, they lack the central...
microtubule pair (9+0) [16,29]. Specifically, within retinal epithelial cells HOOK2 bound to the essential cilia proteins PCMI and RAB8a and was required to initiate axoneme growth from the basal body [20]. These data raise the possibility that HOOK2 is involved in the delivery of LRGUK-1 to the basal body, or vice versa, and in the initiation of sperm tail growth.

As for LRGUK, HOOK2 was localised to the Golgi-derived spermatid acrosome, the acroplaxome, the manchette and the sperm basal body (Fig. 6C-E). HOOK2 localization was not appreciably altered in the presence of the LrgukKaos mutation, albeit in the presence of abnormal germ cell structure (Fig. 6D), indicating that LRGUK-1 does not define HOOK2 localization. Unfortunately genetically modified Hook2 mouse lines are not available to test the dependence of LRGUK localization on HOOK2. As such, and although the formal possibility remains that HOOK2 may be a LRGUK-1 cargo, it is more likely that HOOK2 functions in its established role to transport LRGUK-1.

The LrgukKaos/Kaos sperm phenotype has some similarities with that observed for HOOK2 depleted retinal cells, suggesting they lie in a common pathway. Greatly reduced axoneme development was evidenced in testis sections stained for the axoneme marker acetylated tubulin and the absence of 9+2 axoneme structure was seen at the electron microscopic level (Fig. 7A-E). An analysis of the early steps of centriole/basal body movement and axoneme development, however, revealed abnormalities in LrgukKaos/Kaos germ cells, strongly suggestive of a critical role for LRGUK-1 beyond that currently documented for HOOK2, specifically in the formation of the centriole appendages. In early wild type spermatids the mature centriole can be distinguished from the adjacent daughter centriole by the possession of accessory structures known as the sub-distal appendages (SAP) and distal appendages (DAPs) [30]. The mature centriole then migrates to the spermatid periphery and attaches to the plasma membrane before attaching to the nuclear pole opposite the acrosome [16]. Recent data has shown the DAPs

Fig 5. Manchette structure in LrgukWT/WT and LrgukKaos/Kaos elongating spermatids. α-tubulin staining (green) of manchette microtubules in elongating spermatids from LrgukWT/WT (A) and LrgukKaos/Kaos mice (B) are shown in progressive steps during head elongation. Nuclei were stained with DAPI (blue). The scale bar = 5 μm. Please see S3 Fig. for negative control images.

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Fig 6. LRGUK-1 binds to HOOK2. (A) Growth of yeast carrying wild type LRGUK-1 (LRGUKWT/pDEST32) and HOOK2 (HOOK2/pDEST22) compared to those carrying LRGUKKaos (LRGUKKaos/pDEST32) and HOOK2 (HOOK2/pDEST22), and empty pDEST32 and pDEST22 vectors. Binding was absent (i.e. no growth) when empty vectors were used as a negative control. (B) Confirmation of the interaction between
LRGUK-1 and HOOK2 by immunoprecipitation from HEK293 cells expressing EGFP-tagged LRGUK-1. EGFP-tagged LRGUK-1 was immunoprecipitated using GFP antibody-conjugated beads (anti-GFP-Trap-A beads), and immunoblotting was performed using HOOK2 antibody to detect the interaction between the EGFP-tagged LRGUK-1 and endogenous HOOK2. pEGFP-C1 represents a negative control which over-expressed EGFP protein only. Input: whole cell lysates of transfected HEK293 cells. 

(C) The localization of HOOK2 (red) in WT round spermatids, co-labelled with PNA (green) to mark the acrosome. HOOK2 was present within the developing acrosome/acroplaxome (arrowhead), at a region consistent with the leading edge of the acrosome and acroplaxome (arrows) and at a site consistent with the basal body (asterisk). 

(D) The localization of HOOK2 (red) onto the microtubules (α-tubulin, green) of the manchette in WT and Lrguk<sup>Kaos/Kaos</sup> elongating spermatids. Scale bar = 5 μm. 

(E) HOOK2 (red) within caudal epididymal sperm. Acetylated (Ac) tubulin was used as an axoneme marker. The asterisk indicates HOOK2 in the sperm basal body. Scale bar = 5 μm. In all images, nuclei are labelled with DAPI (blue). Please see S3 Fig. for negative control images.

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**Fig 7. Axoneme defects in Lrguk<sup>Kaos/Kaos</sup> mice.** (A) The staining of testis sections with the axoneme marker, acetylated tubulin, indicated an almost complete absence of axoneme development in Lrguk<sup>Kaos/Kaos</sup> germ cells. (B) Transmission electron microscopy indicating the absence of axoneme development in wild type and Lrguk<sup>Kaos/Kaos</sup> elongating spermatids. Red arrows indicate outer dense fibers. Scale bar = 0.2 μm. (C-D) Transmission electron microscopy of basal body docking to the plasma and nuclear membrane (C) and subsequent axoneme extension in Lrguk<sup>WT/WT</sup> cells but the absence of plasma membrane attachment and axoneme of extension in Lrguk<sup>Kaos/Kaos</sup> spermatids. Scale bar = 2 μm. (E) A closer examination of the early phases of basal body structure and movement revealed the presence of normal distal appendages (DAPs, green arrow) and sub-distal appendages (SAPs, red arrow) in wild type cells associated with plasma membrane attachment. Lrguk<sup>Kaos/Kaos</sup> germ cells contained enlarged SAP and the basal body failed to associate with the plasma membrane. The scale bar = 2 μm.

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are required for basal body-to-membrane docking [31], and SAPs are believed to be where axoneme microtubules are anchored and are thus required for axoneme extension [32–34]. Electron microscopy of spermatids from Lrguk^{WT/WT} spermatids clearly showed the docking of mature centrioles to the plasma membrane and the associated DAPs and SAPs (Fig. 7E). In contrast, the over-whelming majority of basal bodies in spermatids from Lrguk^{Kaos/Kaos} mice contained SAPs that were overtly enlarged and plasma membrane attachments were very infrequent (<1% of the time) (Fig. 7E). Consistent with abnormal SAP function, microtubule extension from the Lrguk^{Kaos/Kaos} basal body into an axoneme was extremely rare (Fig. 7D). Despite the absence of membrane attachment basal bodies did contain DAP-like structures. Lrguk^{Kaos/Kaos} basal bodies did, however, appeared to attach to the nuclear membrane normally (Fig. 7E). In comparison to the basal body phenotype seen in Lrguk^{Kaos/Kaos} spermatids, the loss of HOOK2 function during retinal cell primary cilia formation had no reported effect on SAP formation [19]. These data suggest that LRGUK-1 is required for the formation / function of the DAPs and SAPs in spermatids. Currently it is unknown if these effects are independent of HOOK2 or if they are spermatid-specific functions and thus, not seen following HOOK2 knockdown in retinal cells.

In contrast to the lack of axoneme development in Lrguk^{Kaos/Kaos} spermatids, we observed considerable evidence of the assembly of outer dense fiber-like structures within the distal cytoplasm of elongated spermatids (Fig. 7B). These data suggest that while the outer dense fibers sit in close apposition to the microtubules of the axoneme in normal germ cells, their development can occur independently.

**Discussion**

The dynamic organisation of microtubules and the ability to transport proteins over long distances are fundamental processes required for many cell types, but perhaps none more so than haploid male germ cells, where sperm head shaping and tail development occurs in the virtual absence of transcription [35]. Here we have demonstrated that LRGUK-1 is a critical component of this process with roles that impact upon multiple facets of sperm structure. LRGUK-1 is required for normal acrosome attachment, manchette function, the initiation of the axoneme extension and ultimately male fertility. In accordance with these roles, we observed LRGUK-1 within a protein transport corridor [13] involving movement from the Golgi complex in round spermatids, to the acrosome/acroplaxome, onto the manchette in elongating spermatids then ultimately into the sperm tail. Collectively our data establish LRGUK-1 as a vital component for haploid germ cell development and male fertility.

The earliest discernable abnormality seen in Lrguk^{Kaos/Kaos} germ cells was in acrosome development. The acrosome is a vesicle-like structure at the caudal pole of the sperm head. The plasma membrane immediately above the acrosome is the first point of binding between the sperm and the cumulus oocyte complex prior to fertilisation, and the contents of the acrosome is required for sperm penetration through the outer vestments of the oocyte [36]. Data presented here suggests that LRGUK-1 is required for the appropriate attachment of Golgi-derived pro-acrosomal vesicles onto the nucleus and the full structural integrity of the acrosome-acroplaxome attachment to the nuclear membrane [15,17]. LRGUK-1 dysfunction resulted in the fragmentation and loss of some pro-acrosomal vesicles in early round spermatids, and the detachment of the acrosome-acroplaxome from the sperm head during late spermiogenesis. Clearly however, acrosome-like structures were formed in spermatids (albeit with apparently reduced efficiency in early round spermatids) indicating that the enhanced adhesion inferred by LRGUK-1 may only become critical when shear forces are applied, for example during stages I through to VIII wherein elongated spermatids are sequentially dragged down into
Sertoli cell crypts then pushed up to the luminal aspect of the seminiferous epithelium [37]. The apparent absence of LRGUK-1 protein from the acroplaxome region in mature sperm suggests that this defect maybe mediated by an LRGUK-1 binding partner rather than LRGUK-1 itself, or that LRGUK-1 is part of a stepwise process resulting in firm acrosome-acroplaxome-nuclear attachment. Taken together our results suggest a role for LRGUK-1 in the trafficking of pro-acrosomal vesicles from the Golgi to the acroplaxome and in the attachment of acrosome-acroplaxome to the nuclear membrane.

A second major defect, and almost certainly the cause of the head abnormalities in \(Lrguk^{Kaos/Kaos}\) sperm, was a defect in the structure and movement of the manchette. The manchette is a complex microtubule array that forms around the spermatid nucleus concurrent with the initiation of nuclear compaction. It is composed of a perinuclear ring and a fringe of microtubule bundles that extend into the distal cytoplasm. As spermiogenesis proceeds the manchette moves distally, and in the case of species such as the mouse that contain falciform shaped sperm heads, it pivots in a manner dependent on microtubule severing [22]. In parallel, the perinuclear ring constricts, likely contributing to the tapered shape of the post-acrosomal region of the sperm head. Our data show that LRGUK-1 is required for the distal movement of the perinuclear ring. In the presence of the \(Lrguk^{Kaos/Kaos}\) mutation the perinuclear ring constricted more proximally than normal leading to abnormal sperm head shape. Interestingly the remaining LRGUK present within elongating spermatids (LRGUK-2 and/or -3 or small amounts of truncated LRGUK-1) remained at the leading edge of the acroplaxome, suggesting that there are motifs C-terminal 293 amino acids of LRGUK-1 that are required for the transition to the manchette.

Of relevance, a second member of the HOOK family is required for manchette function and ultimately male fertility. The \(azh\) mouse phenotype was caused by the deletion of exons 10–11 from the \(Hook1\) gene [38]. Homozygous \(azh\) males displayed OAT characterised by severe head abnormalities and the frequent decapitation from the sperm tail. The latter is suggestive of a weakened connection between the nuclear membrane and the basal body-derived axoneme. This phenocopying and data suggesting that HOOK family members frequently function as heterodimers [39] raise the possibility that LRGUK-1 exists in a complex with both HOOK2 and HOOK1. This possibility will be tested in future experiments.

The most striking aspect of the Kaos phenotype, and the ultimate cause of male sterility, was the almost complete block of sperm tail development. LRGUK-1 dysfunction lead to abnormal SAP formation on basal bodies and an absence of plasma membrane attachment, likely associated with DAP dysfunction. As a consequence, basal bodies failed to nucleate axoneme microtubules. Within retinal primary cilia, HOOK2 is involved in the assembly of a complex containing PCM1 and RAB8a, and the initiation of axoneme microtubule extension [19]. In contrast to HOOK2 action in primary cilia, however, our data revealed that LRGUK-1 also has a critical role in the formation of basal body-plasma membrane attachment and SAP formation. Currently it is unknown if these effects are independent of HOOK2 or if they are spermatid-specific functions and thus, not seen following HOOK2 knockdown in retinal cells. Our data do not, however, support a role for LRGUK-1 in cilia broadly. Notably, the Kaos mouse line showed none of the morphometric features characteristic of primary cilia disorders such as polydactyly and craniofacial abnormalities [40], although the presence of \(Lrgk-1\) mRNA in tissues containing motile cilia (9+2 axonemes) raises the possibility of an age-related pathology associated with motile cilia dysfunction.

The phenotypic similarity observed in \(Lrguk^{Kaos/Kaos}\) and mice containing mutations in core proteins of the IFT pathway including \(Ift88\) and \(Kif3a\) [41,42] highlights the continuity of the protein transport pathway underlying the movement of proteins involved in sperm head shaping and axoneme extension. Our data suggest that LRGUK-1 functions upstream/before the
IFT pathway i.e. spermatids with defective LRGUK-1 will undergo a developmental arrest prior to the requirement for the IFT pathway. These data also highlight the exquisite value of using the testis as a model to define processes of likely fundamental importance to cell biology broadly.

Collectively, we have identified LRGUK-1 as a protein critically involved in multiple aspects of sperm assembly and function. LRGUK-1 is required for appropriate acrosome attachment to the sperm head, sperm head shaping via the manchette and tail growth from the basal body. During the initiation of sperm tail axoneme extension from the basal body, LRGUK-1 functions in plasma membrane attachment and the formation of SAP and ultimately microtubule extension. The specific loss of LRGUK-1 function results in OAT in mice and raises the possibility that LRGUK dysfunction leads to human OAT.

Materials and Methods

Identification of the Kaos line and the causal mutation

Animal procedures were performed in accordance with Australian NHMRC Guidelines on Ethics in Animal Experimentation and approved by the Australian National University and Monash University Animal Experimentation Ethics Committees. Point mutant mice were generated on a C57BL/6 background and outbred to CBA and individual lines screened for sterility causing mutations as described previously [9,21]. Mating behaviour, as indicated by the presence of copulatory plugs was normal.

The sterility causing mutation in the Kaos line was initially linked to a region on chromosome 6 using a SNP-based microarray approach. The region was subsequently narrowed using additional mice and SNPs to a 12.9 Mb region (30872499–43776812 bp) that contained 74 genes. Candidate genes were selected based on expression in haploid germ cells, i.e. the site of the phenotype, in EST expression databases. 28 genes were expressed within round spermatids. The full coding region of eight of these genes was sequenced and a single C to T point mutation within the Lrguk gene was identified.

Following the identification of the causal mutation, mice were genotyped using the Amplifluor system (Chemicon) using a wild type-specific reverse primer 5’-GAAGTGGGAGTTCAACGGATTCCATAGGCACCACACAAAGATATC-3’, a mutant allele specific reverse primer 5’-GAAGGTGACCAAGTTCATGCTCAGGCACCACAAAGATATC-3 and a conserved forward primer 5’-CAGCCTTGGACTATTTATAGGGAGTGTG-3’ as described previously [9].

LRGUK orthologues were identified as described previously [7].

Characterization of the Kaos sterility phenotype

Sterility in the Kaos mouse line was characterized using the regime outlined in Borg et al [43]. Daily sperm production (DSP) in the testis and total epididymal sperm content was determined using the Triton X100 nuclear solubilization method as described previously [44]. Sperm ultra-structure was assessed using electron microscopy as described previously [45]. Sperm motility was assessed visually using sperm back-flushed from the cauda epididymis [46]. Cauda epididymal sperm morphology was visualized using haematoxylin and eosin staining. The stages of the epithelium tubule were judged by Periodic Acid Schiff (PAS) staining [47]. Apoptosis was evaluated by TUNEL Apoptosis Detection Kit (Millipore) according to the manufacturer’s instructions; n = 5 mice per genotype and the positive cells in 100 seminiferous tubules were counted for each mouse.

Germ and Sertoli cell numbers were counted in 25µm-thick PAS stained methacrylate embedded testis sections using the optical dissector method as described previously [48]. Basal
retained elongated spermatids were counted in stage VII-XII [22,48]. N = 5 mice per genotype and 240 counting frames for each mouse.

Quantitative real time PCR

Lrguk in adult tissues and at different time points during the establishment of the first wave of spermatogenesis was defined as outlined previously [49]. Lrguk expression was detected using the Taqman assay (Mm01166701-m1). Lrguk-2 expression was detected using a custom designed assay (Forward primer 5’-CCCCAAAAATCTCAAGGTATACTTATCAT-3’; reverse primer 5’-CCGCAGCTGAAGCAAAAACGC-3’; probe 5’-CAAAGCACACAATGGTT-3’). A custom designed assay was also used to detect both Lrguk2 and Lrguk3 transcripts (Forward primer 5’-CTGGCTATCTGCTGTACATC-3’; reverse primer 5’-CCGCAGCTGAAGCAAAAACGC-3’; probe 5’-CAAAGCACACAATGGTT-3’). All expression data was normalized to the Ppia house-keeping gene (Mm02342429-g1).

Germ cell purification and immunochemistry

Germ cells were isolated using the Staput method described previously [50]. For immunofluorescence staining, the purified cells were fixed with 4% paraformaldehyde (PFA) and stained as described previously [9]. Primary antibodies used included: anti-LRGUK (1:200 dilution, Novus Biologicals, raised against amino acids 32–181); anti-HOOK2 (1:200 dilution, GeneTex), anti-acetylated tubulin (1:1000 dilution, Sigma), anti-α-tubulin (1:5000 dilution, Sigma). All primary antibodies were diluted in 10% non-immune horse serum (NHS) in PBS and incubated at 4°C overnight. Secondary antibodies included: Alexa Fluor 555 donkey anti-rabbit IgG and Alexa Fluor 488 donkey anti-mouse IgG, 1:500 dilution in 10% NHS in PBS for 45 minutes at room temperature. Nuclei were labelled with DAPI. The acrosome was visualized using FITC-PNA (1:2000 dilution, Sapphire Bioscience). Images were taken with a SP8 confocal microscope (Leica Microsystems). For consistence α-tubulin was artificially coloured green in all images.

Paraffin-embedded testis sections were stained for acetylated tubulin as described previously [45].

Yeast two hybrid screen

The adult mouse testis cDNA library (pDEST22 prey vector) used for yeast two hybrid screen was as described previously [51]. The full length LrgukWT and LrgukKaos cDNAs were cloned into pDEST32 vector using the Gateway cloning kit (Invitrogen). For the initial identification of LRGUK binding partners, the LRGUKWT-pDEST32 vector was used as bait in a yeast two hybrid screen with the ProQuest Two-Hybrid System (Invitrogen). Briefly, the LRGUKWT-pDEST32 vector and pDEST22 testis cDNA library constructs were co-transformed into Mav203 yeast strain. Putative interacting clones were isolated from the resulting yeast colonies and re-transformed into E. coli to obtain high purity plasmids for sequencing cDNA inserts. Sequencing of the longest HOOK2 clone revealed an open reading frame of 1078 bp that corresponded to the C-terminal amino acids 369–716 of the HOOK2 protein. To determine the effect of the LRGUKKaos mutation on the binding to HOOK2, the identified HOOK2-pDEST22 prey vector was co-transformed into the Mav203 yeast strain along with either LRGUKKaos-pDEST32, LRGUKWT-pDEST32 or pDEST32 empty vector only and plated on selection media.
Cloning and co-immunoprecipitation

Full length mouse Lrguk-1 was amplified from wild type C57BL/6J testis cDNA using Lrguk-Fw: 5’-ATATAAGATCTGGGCTTTGAGCAAAT-3’ and Lrguk-Rev: 5’-ATAAAGGGCCCTATGCGGCGATGCGGGAT-3’ primers than cloned into the pEGFP-C1 expression vector (Clontech, Gen Bank Accession number U55763). The LRGUK-1/pEGFP-C1 construct was transfected into HEK293 cells using Lipofectamine 3000 Reagent (Life Technologies, Cat. No. L3000008). Co-immunoprecipitation was carried out using anti-GFP-Trap-A beads as per the manufacturer’s instructions (Chromotek, Cat. No. gta-100). Empty pEGFP-C1 vector was used as a negative control. The presence of LRGUK-GFP and endogenous HOOK2 proteins were determined using anti-GFP (Roche, Cat. No. 11814460001 and HOOK2 antibodies (GeneTex, Cat. No. GTX115898, corresponding with amino acids 197–380 of HOOK2).

Western blotting

For LRGUK-1 expression, protein was extracted from both wild type and mutant testes using 1% NP-40/PBS lysis buffer with 1:200 protein inhibitor cocktail (Promega). Protein (40 μg) was separated on a 12% SDS-PAGE gel. Non-specific antibody binding was minimized by blocking the membrane with 5% skim milk for 1 hour room temperature and antibodies were diluted in 1% skim milk. The membrane was probed with a rabbit LRGUK-1 antibody (Sigma, raised against amino acids 336–427 of LRGUK-1 and thus will not bind to LRGUK-2 or -3). Bound antibody was detected using a goat anti-rabbit IgG HRP (Dako) secondary antibody. Antibody binding was detected using the enhanced chemiluminescence (ECL Plus) detection kit (Amer-sham Biosciences). Protein loading was normalized against actin.

Supporting Information

S1 Table. The results of a stereological analysis of germ cell content in the Lrguk<sub>Kaos/Kaos</sub> mouse compared to Lrguk<sub>WT/WT</sub> littermates. † Data is expressed as number per Sertoli cell, n = 5 per group, mean±SEM. * denotes p<0.05 compared to Lrguk<sub>WT/WT</sub>.

S1 Fig. (A) Body weigh of Lrguk<sub>WT/WT</sub> males versus males (n = 12–13, age 8–12 weeks). (B–C) The levels of apoptosis were unchanged in Lrguk<sub>Kaos/Kaos</sub> testes compared to Lrguk<sub>WT/WT</sub> n = 3 per genotype. (D) Stage VIII tubules showing the retention of elongating spermatids in the basal region of the seminiferous tubule in Lrguk<sub>Kaos/Kaos</sub> males: Note elongating spermatid numbers were also reduced in number and lacked tails. Scale bar = 100 μm.

S2 Fig. The localization of LRGUK in Lrguk<sub>WT/WT</sub> and Lrguk<sub>Kaos/Kaos</sub> elongating spermatids. The localization of LRGUK (red) in elongating spermatids from Lrguk<sub>WT/WT</sub> and Lrguk<sub>Kaos/Kaos</sub> mice. Scale bar = 5 μm. Manchettes were visualised by α-tubulin (green) and nuclei were stained with DAPI (blue). Manchettes were abnormal in Lrguk<sub>Kaos/Kaos</sub> mice, as indicated by patchy α-tubulin signal. Abnormal manchette movement was indicated by the constriction of the spermatid head. LRGUK immunostaining was more obvious in the marginal ring of Lrguk<sub>Kaos/Kaos</sub> mice (row 3), compared to Lrguk<sub>WT/WT</sub> mice. Please see S2 Fig. for negative control images.

S3 Fig. Negative controls of immunofluorescent cell labelling. (A) A negative control image for Figs. 3C and 6C- isolated round spermatids. Negative control for secondary antibody
donkey anti-rabbit Alexa 488. (B) A negative control image for Figs. 3D and 6D (D) and S2 Fig. (F)—isolated elongating spermatids. Negative control for secondary antibodies donkey anti-rabbit Alexa 488 and donkey anti-mouse Alexa 555. (C) A negative control for Figs. 3E and 6E—mouse sperm. Negative control for secondary antibodies donkey anti-rabbit Alexa 488 and donkey anti-mouse Alexa 555. (E) A negative control image fro Fig. 5—isolated elongating spermatids. Negative control for secondary antibody donkey anti-mouse Alexa 555.

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Author Contributions

Conceived and designed the experiments: MKO CJO CCG YL KDB DJ. Performed the experiments: YL KD DJM AEO AE DAJ DJM BW HO. Analyzed the data: MKO CJO CCG DJ RIM DMdK YL KDB AEO DJM LO BW HO. Contributed reagents/materials/analysis tools: DAJ AE. Wrote the paper: MKO LO YL KD DJ.

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