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# NHR-23 and SPE-44 regulate distinct sets of genes during Caenorhabditis elegans spermatogenesis

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#### **Abstract**

Spermatogenesis is the process through which mature male gametes are formed and is necessary for the transmission of genetic information. While much work has established how sperm fate is promoted and maintained, less is known about how the sperm morphogenesis program is executed. We previously identified a novel role for the nuclear hormone receptor transcription factor, NHR-23, in promoting *Caenorhabditis elegans* spermatogenesis. The depletion of NHR-23 along with SPE-44, another transcription factor that promotes spermatogenesis, caused additive phenotypes. Through RNA-seq, we determined that NHR-23 and SPE-44 regulate distinct sets of genes. The depletion of both NHR-23 and SPE-44 produced yet another set of differentially regulated genes. NHR-23-regulated genes are enriched in phosphatases, consistent with the switch from genome quiescence to post-translational regulation in spermatids. In the parasitic nematode *Ascaris suum*, MFP1 and MFP2 control the polymerization of Major Sperm Protein, the molecule that drives sperm motility and serves as a signal to promote ovulation. NHR-23 and SPE-44 regulate several MFP2 paralogs, and NHR-23 depletion from the male germline caused defective localization of MSD/MFP1 and NSPH-2/MFP2. Although NHR-23 and SPE-44 do not transcriptionally regulate the casein kinase gene *spe-6*, a key regulator of sperm development, SPE-6 protein is lost following NHR-23+SPE-44 depletion. Together, these experiments provide the first mechanistic insight into how NHR-23 promotes spermatogenesis and an entry point to understanding the synthetic genetic interaction between *nhr-23* and *spe-44*.

Keywords: Caenorhabditis elegans; spermatogenesis; NHR-23; SPE-44; nuclear hormone receptor; transcription factor; gene regulation

#### Introduction

Spermatogenesis involves a cascade of morphogenetic events to produce the highly specialized, haploid, motile gametes essential for sexual reproduction (Fabian and Brill 2012; Ellis and Stanfield 2014; Nishimura and L'Hernault 2017). Universal features of spermatogenesis include the biosynthesis of sperm-specific proteins and assembly of sperm-specific complexes, preparing for and progressing through the meiotic divisions, and a streamlining step during which cells discard cytoplasmic components that are no longer needed (Nishimura and L'Hernault 2017). At speciesspecific points during sperm development, transcription ceases; the final steps of differentiation and mature sperm function occur in the absence of transcription (D'Occhio et al. 2007; Rathke et al. 2007; Shakes et al. 2009). The sperm's genome is protected from environmental and metabolic damage through a spermatogenesis-specific process of chromatin remodeling and hypercompaction (Wu and Chu 2008). These processes are all essential for the faithful transmission of the sperm genome to offspring.

Caenorhabditis elegans is a powerful model to study spermatogenesis. Within the self-fertile hermaphrodites, uncommitted germ cells have the capacity to undergo either spermatogenesis or oogenesis; but in the normal process, the hermaphroditic germline makes a one-time switch from spermatogenesis to oogenesis as animals enter adulthood. The analysis of the accompanying changes provides insight into how sperm and egg fates are promoted and maintained. Alternatively, the analysis of males allows the study of continual spermatogenesis. Both hermaphrodite and male germlines are polarized organs; the uncommitted germ cells proliferate mitotically at the distal end before committing to entering the meiotic prophase and commitment to the sperm fate. The events of meiosis occur in a linear fashion along the length of the gonad, resulting in the production of haploid spermatids. These include homologous chromosome pairing, DNA breaks that promote recombination, and 2 meiotic divisions (Shakes et al. 2009; Chu and Shakes 2013; Ellis and Stanfield 2014; Griswold 2016). During C. elegans spermatogenesis, general transcription ceases near the end of the meiotic prophase, and there is no burst of postmeiotic transcription as occurs in humans and Drosophila (Bettegowda and Wilkinson 2010; Chu and Shakes 2013). The translation ceases shortly after anaphase II, as not only ribosomes but also tubulin, actin, and endoplasmic reticulum are partitioned away from the budding spermatids and into a structure known as the residual body (Chu

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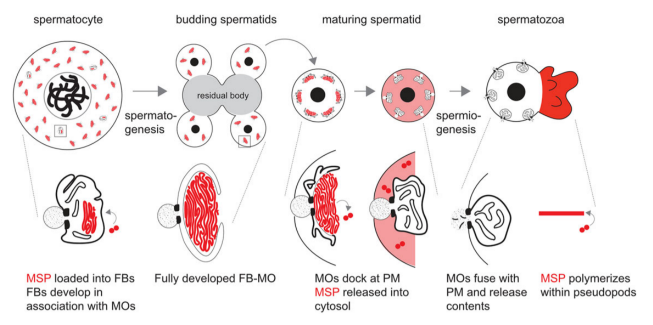


Fig. 1. Overview of MSP dynamics during spermatogenesis. (Top) Spermatocytes undergo reductive cell division during spermatogenesis. During this process, MSP is packaged and sequestered in FB-MOs. Following anaphase II, components which are no longer needed partition to central residual bodies, which are discarded. Spermatids bud off of the residual bodies, mature, and upon activation, undergo morphological changes during spermiogenesis. The differentiated spermatozoa contain cell bodies and pseudopodia. (Bottom) Within spermatocytes, FBs develop on the cytosolic face of the Golgi-derived MO. During the budding division, FB-MO complexes partition to the spermatids. In maturing spermatids, MOs dock with the plasma membrane as the FBs disassemble and release MSP dimers into the cytosol. During sperm activation, MSP localizes to the pseudopod, where it polymerizes, and MOs fuse with the plasma membrane.

and Shakes 2013; Fig. 1). Lacking typical cytoskeletal proteins, nematode sperm motility is driven by the assembly/disassembly dynamics of the 14kDa nematode-specific major sperm protein (MSP) within the pseudopod of the crawling spermatozoa (Smith 2014). The early cessation of both transcription and translation requires all sperm-specific components to be synthesized during meiotic prophase prior to their developmental deployment. During this critical window, newly synthesized MSP assembles into large paracrystalline fibrous bodies (FBs) that form in association with Golgi-derived membranous organelles (MOs; Fig. 1). This form of MSP packaging facilitates the efficient partitioning of MSP to spermatids.

The transcription and translational programs that govern C. elegans spermatogenesis must not only drive differentiation of spermatocytes and oocytes, but they must also create the materials that are needed for sperm differentiation and sperm function. How early-acting gene regulatory networks provision spermatocytes for the extended process of spermatogenesis and sperm morphogenesis remains an open question. To date, 2 transcription factors have been characterized as critical regulators of C. elegans spermatogenesis: SPE-44 and NHR-23. SPE-44 was first identified by its enrichment in sperm-producing germlines. Further investigation revealed that spe-44 null spermatocytes fail to load MSP into FB-MOs, and they are arrested as terminal spermatocytes during the second meiotic division (Kulkarni et al. 2012). spe-44 regulates a broad set of genes that are enriched in spermatogenic germlines (Kulkarni et al. 2012). Consistent with the MSP loading defects, these genes include MSP genes and members of the small sperm-specific protein family, which promote MSP polymerization (Kulkarni et al. 2012). spe-44 also regulates several genes necessary for spermatogenesis, which were previously identified through forward genetic screens for spermatogenesis defective (spe) and fertilization-defective (fer) genes (Kulkarni et al. 2012). However, many other spe and fer genes are

not regulated by spe-44 (Kulkarni et al. 2012). In terms of regulatory hierarchies, spe-44 regulates the transcription factor, elt-1 (Kulkarni et al. 2012), which is a direct regulator of MSP gene expression (del Castillo-Olivares et al. 2009).

We recently discovered that the nuclear hormone receptor NHR-23, a critical regulator of nematode molting, is also expressed in primary spermatocytes and necessary for spermatogenesis (Ragle et al. 2020). NHR-23 and SPE-44 expression patterns almost completely overlap (Ragle et al. 2020). Both proteins label the autosomes but not the X-chromosome from the beginning of the meiotic prophase until global transcription ceases and spermatocytes enter the karyosome stage. Germline-specific NHR-23 depletion causes a terminal spermatocyte arrest during the first meiotic division (Ragle et al. 2020). Spermatocytes fail to load MSP into FBs upon single depletion of either SPE-44 or NHR-23, whereas the Golgi-derived MO portion of the FB-MO complex is specifically perturbed upon SPE-44 depletion and severely disrupted upon double depletion (Ragle et al. 2020). We determined that SPE-44 and NHR-23 are in separate genetic pathways because they do not regulate one another, and because NHR-23 and SPE-44 double depletion produce more severe synthetic phenotypes such as a nearly complete loss of MO differentiation markers and severe germline defects (Ragle et al. 2020). Exactly how NHR-23 promotes spermatogenesis was not clear from our previous work. Here, we use a combination of transcriptomics, genome editing, and cytology to address this question and gain insight into the synthetic genetic interaction between spe-44 and nhr-23.

#### Materials and methods

#### Strains and culture

Caenorhabditis elegans were cultured as originally described (Brenner 1974), except that worms were grown on MYOB media instead of NGM. MYOB agar was made as previously described (Church et al. 1995). Brood sizes were performed picking L4 larvae to individual wells of a 6-well plate seeded with OP50 and incubating the plate at 20°C. Animals were transferred to new plates daily over 4 days. Two days post-transfer, the number of hatched progeny and unfertilized eggs were scored. We scored unfertilized eggs over the first 3 days as on the fourth day animals frequently ran out of sperm.

Table 1 contains a list of all the strains used and their source.

#### RNA-seq and bioinformatics

Adult hermaphrodites of the indicated genotypes were picked onto 6 cm MYOB + 4 mM auxin plates and allowed to lay eggs for 2 h. The hermaphrodites were removed, and the plates were incubated at 20°C for 3 days. All strains carried an him-5 (e1490) allele to increase the number of males in hermaphrodite progeny. Five hundred L4/young adult males were hand-picked into M9 + 0.05% gelatin, washed 1X, and pelleted. Liquid was removed down to 25 µl and 500 µl TRIzol (Invitrogen) was added. The mixture was incubated at room temperature for 5 min with periodic vortexing. Carcasses were pelleted by centrifugation and the supernatant was removed to a clean tube. One hundred microliters chloroform was added to the supernatant, vortexed for 30s and phases separated by centrifugation at 15,000 RPM for 5 min. Two hundred and fifty microliters of the upper aqueous phase was removed to a new tube and  $350\,\mu l$  chloroform was added, vortexed, and separated as in the previous steps. Two hundred and fifty microliters of isopropanol was added to the aqueous phase along with 1 µl Glycoblue (Invitrogen) and vortexed. The solution was centrifuged for 20 min at 15,000 RPM and the supernatant was decanted. The pellet was washed with 1 ml 75% ethanol twice and dried at room temperature for 10 min. The dried pellet was resuspended in  $16\,\mu l$  nuclease-free water. The RNA concentration was determined with a NanoDrop.

Using the NEXTFLEX Rapid Directional RNA-Seq Kit 2.0 (BioO/ PerkinElmer), polyA-selected RNA was isolated from 300 ng of total RNA according to the manufacturer's instructions. The sequencing libraries were prepared using the same kit according to manufacturer's instructions. The concentrations were measured on a Qubit Fluorometer (Invitrogen) and fragment size

determined on a Bioanalyzer 2100 (Agilent). Libraries were sequenced at the University of California Berkeley QB3 NGS facility on a Nova-seq platform. Raw sequences were mapped to transcriptome version WS220 using Tophat2 (Kim et al. 2013). Only reads with one unique mapping were retained, otherwise, default options for paired-end mapping were used. HTSeq was used to obtain read counts per transcript (Anders et al. 2015). All libraries were sequenced twice and for each transcript read counts from both sequencing runs were added together. DESeq2 was used for read count normalization and differential expression analysis (Love et al. 2014). DESeq2's regularized log transformation with option blind=TRUE was used for read count normalization of protein-coding genes prior to principal component analysis (PCA). A Benjamini-Hochberg multiple hypothesis-corrected Pvalue cutoff of 0.05 was used as a cutoff for significant differential expression. PCA was performed using the prcomp function in the stats package of R (R Core Team). Supplementary Table 1 contains the processed data with gene name and location, mean expression, log2 fold-change compared to control, and P-value. The 3D PCA scatter plot was generated with http://www.bioinfor matics.com.cn/srplot (last accessed Sept. 27, 2022), an online platform for data analysis and visualization. Protein and cDNA alignments were produced with Clustal Omega (Goujon et al. 2010; Sievers et al. 2011). Protein alignments were visualized with JalView2 (Waterhouse et al. 2009). Venn diagrams were generated using GeneVenn (http://genevenn.sourceforge.net/; last accessed Sept. 27, 2022) and figures were generated in Affinity Designer.

### CRISPR/Cas9-mediated genome editing

The mutations in candidate NHR-23 target genes were generated using homologous recombination of CRISPR/Cas9-induced double-strand breaks (DSBs). We made Cas9 ribonucleoprotein (RNP) complexes (750 ng/µl house-purified Cas9, 50 ng/µl crRNA, and 100 ng/µl tracrRNA) by incubating at 37°C for 15 min. RNPs were mixed with oligonucleotide repair templates (110 ng/μl; Ghanta and Mello, 2020; Paix et al. 2015; Dokshin et al. 2018) and a coinjection marker (either 50 ng/µl pRF4, 1-10 ng/µl pKA411, or 25 ng/µl pSEM229; El Mouridi et al. 2020) to facilitate isolation of successfully injected progeny. Where possible, we selected "GGNGG" crRNA targets as these have been the most robust in our hand

Table 1. List of strains used.

Strain	Genotype	Source/reference	
N2	Wild type		
JDW85	wrdSi14 mex-5p::TIR1: F2A: mTaqBFP2: tbb-2 3'UTR, I:-5.32 ; him-5(e1490) V.	Ragle et al. (2020)	
JDW86	wrdSi15[mex-5p::TIR1: F2A: mTagBFP2: tbb-2 3'UTR, I:-5.32]; spe-44(fx11ó[spe-44::AID*]) IV; him-5(e1490) V.	Ragle et al. (2020)	
JDW92	wrdSi19 mex-5p::TIR1: F2A: mTagBFP2: tbb-2 3'UTR, I:-5.32], nhr-23(kry61(nhr-23::AID*-TEV-3xFLAG)) I; him-5(e1490) V.	Ragle et al. (2020)	
JDW146	wrdSi19 mex-5p::TIR1: F2A: mTagBFP2: tbb-2 3'UTR, I:-5.32], nhr-23(kry61(nhr-23::AID*-TEV-3xFLAG)) I; spe-44(fx110[spe-44::degron]) IV; him-5(e1490) V.	Ragle et al. (2020)	
JDW307	nsph-3.2(wrd56 C04G2.9::exon 1 STOP ) IV	This study	
JDW308	nsph-2(wrd57[ZK265.3::exon 1 STOP]) Ï	This study	
JDW320	nsph-2(wrd57[ZK265.3::exon 1 STOP]) I; nsph-3.2(wrd56[C04G2.9::exon 1 STOP]) IV	This study	
JDW339	nsph-2(wrd57[ZK265.3::exon 1 STOP]) I; nsph-1.1(wrd67[K08F8.5::exon 2 STOP]) II	This study	
JDW358	ZK596.2(wrd71[exon 1 STOP]) III	This study	
JDW361	ttbk-5(wrd72[exon 1 STOP]) IV	This study	
JDW367	nsph-À.3(wrd74[T04A6.4 exon 1 STOP]) III	This study	
JDW374	ZK757.2(wrd77[exon-1 STOP]) III	This study	
JDW403	nsph-4.1(wrd92(whole gene deletion)) IV	This study	
JDW404	nsph-4.2(wrd93[whole gene deletion]) IV	This study	
JDW427	nsph-4.1(wrd92[whole gene deletion]), nsph-4.2(wrd97[whole gene deletion]) IV	This study	
RV120	spe-44(ok1400) dpy-20 (e1282)/let-92 (s677) unc-22 (s7) IV ok1400 deletion removes 1,577 of the 1,797 base-pair coding region, including the conserved SAND domain. Presumed null	Kulkarni et al. (2012	

and support efficient editing (Farboud and Meyer 2015). All STOP-IN cassettes were designed to be inserted directly at the DNA DSB site. Sequences and descriptions of the crRNAs and repair oligos are provided in Supplementary Tables 2 and 3, respectively. We picked L4 animals, aged them 1 day at 20°C and injected 6-18 animals for each edit. Injected animals were grown at either 20°C or 25°C until progeny were produced. These progenies were isolated to individual plates to lay eggs. We made genomic lysates from parent animals and genotyped them by polymerase chain reaction using primers that anneal to sites flanking the mutations. As our version of the STOP-IN cassette contains a BamHI site, insertions were identified by diagnostic BamHI digestion. Whole gene deletions were identified by PCR product size. Mutations and homozygosity were verified by Sanger sequencing. Genotyping primers are provided in Supplementary Table 4. Knock-in sequences are provided in Supplementary File 1.

#### **Immunocytochemistry**

Embryos were isolated from adult hermaphrodites of the indicated genotypes by standard bleaching methods (dx.doi.org/ 10.17504/protocols.io.j8nlkkyxdl5r/v1; last accessed Sept. 27, 2022) and allowed to hatch overnight in the absence of food. L1 larvae were transferred to multiple OP50 plates for 6-8 h. The treatment groups were washed off the plates and transferred to MYOB+4mM auxin plates. Sibling controls were kept on nonauxin plates. Both groups were raised to adulthood at room temperature. Sperm spreads were obtained by dissecting 10-12 male worms in 7 μl of egg buffer (Edgar 1995) on ColorFrost Plus slides (Fisher Scientific, 12-550) that were additionally coated with poly-L-lysine (Sigma Aldrich, P8290). Worms of different genotypes were dissected on the same slide to ensure that antibody binding conditions were competitive and identical. During analysis, mutants were distinguished by their DAPI-stained nuclear patterns. Separate immunocytology preps were performed with worms of different genotypes on different slides to help identify the phenotypes observed on mixed slides. Samples were freezecracked in liquid nitrogen and fixed overnight in −20°C methanol. Specimen preparation and antibody labeling followed established protocols (Shakes et al. 2009). Primary antibodies included: 1:300 rabbit anti-MSD-1 (F44D1.3) polyclonal (Kosinski et al. 2005), 1:200 rabbit anti-MFP2/NSPH-2 (ZK265.3) polyclonal (Morrison et al. 2021), and 1:250 rabbit anti-SPE-6 (Peterson et al. 2021). 1:400 Alexa Fluor Plus 555 goat anti-rabbit IgG (Invitrogen, A32732) was used as the secondary antibody. Final slides were mounted with Fluoro Gel with DABCO (Electron Microscopy Sciences #17985-02) containing DAPI. Images were acquired with epifluorescence using an Olympus BX60 microscope equipped with a QImaging EXi Aqua CCD camera. Photos were taken, merged, and exported for analysis using the program iVision. Image exposures were optimized for control samples and then all samples were imaged with the same exposures and no further image processing so that images would reflect relative intensities.

#### Results

#### NHR-23 and SPE-44 regulate distinct sets of target genes

Given that NHR-23 is a transcription factor, we hypothesized that loss of NHR-23-regulated gene expression was mediating the phenotypes we observed when NHR-23 was depleted. We therefore depleted NHR-23, SPE-44, and NHR-23+SPE-44 in male germlines using a mex-5p::TIR1 transgene that we developed for an auxininducible degron toolkit (Ashley et al. 2021). We performed bulk RNA-seq on whole male animals to identify differentially regulated genes relative to a mex-5p::TIR1 control strain grown on auxin. With our sequencing data, we performed PCA of the replicates of all conditions. The first 3 principal components captured 63% of the variance and separated the various conditions (Fig. 2, a and b, Supplementary Fig. 1). Using a P < 0.05 cutoff, there were only 70 genes that significantly changed in expression when NHR-23 was depleted (Fig. 2c, Supplementary Table 5). NHR-23 appeared to function primarily to activate gene expression as 55/70 genes were downregulated following NHR-23-depletion (Fig. 2c). SPE-44-depletion resulted in 811 differentially expressed genes, with 523 genes being downregulated (Fig. 2d, Supplementary Table 6). For all 3 datasets, we set the cutoff for differentially regulated genes as a >1.5-fold change in expression relative to the control. NHR-23+SPE-44 double depletion produced 554 differentially expressed genes, with 494 being downregulated (Fig. 2e, Supplementary Table 7). When compared with an RNA-seq study of spermatogenesis vs oogenesis enriched genes (Ortiz et al. 2014), NHR-23-regulated and NHR-23+SPE-44regulated genes had a high percentage [72.9% (51/70) and 84.3% (467/554), respectively] of sperm-enriched genes (Fig. 2, g and h). In contrast, only 59.9% (306/511) of SPE-44-regulated genes were spermatogenesis-enriched (Fig. 2i). A spe-44 mutant microarray experiment found 50.9% (358/703) genes were spermatogenesisenriched, though this percentage went to 67.8% (343/506) if only the downregulated genes were analyzed (Kulkarni et al. 2012). When we performed a similar analysis on our SPE-44downregulated genes we found 53.5% (296/553) were spermatogenesis-enriched (Supplementary Fig. 2a).

We were surprised to find so few differentially regulated genes following NHR-23 depletion in the germline as nhr-23 regulates 265 genes in the soma (Kouns et al. 2011). NHR-23 regulated genes included a few genes previously implicated in spermatogenesis. NHR-23 regulated the MSP Domain containing proteins MSD-1, MSD-2, and MSD-3 which are poorly characterized paralogs of Ascaris MSP fiber protein 1 (MFP1), an MSP polymerization inhibitor (Buttery et al. 2003; Grant et al. 2005; Kosinski et al. 2005; Morrison et al. 2021; Table 2). smz-2 was also NHR-23 regulated (Table 2). smz-2 has a paralog (smz-1), and the proteins SMZ-1/2 localize to chromatin during meiosis (Chu et al. 2006). In addition, RNAi targeting of either SMZ gene causes a metaphase arrest during meiosis (Chu et al. 2006). In contrast to the short list of NHR-23-regulated genes, we found 811 genes that were differentially expressed upon SPE-44 depletion (Fig. 2, Supplementary Table 6) and 554 genes differentially expressed when both NHR-23 and SPE-44 were depleted (Fig. 2, Supplementary Table 7). Surprisingly, our SPE-44-regulated gene set had limited overlap with a set of spe-44-regulated genes identified through a microarray experiment using spe-44 null L4 males (Kulkarni et al. 2012). Only 180 of the 811 SPE-44-regulated genes from our RNA-seq dataset were common to the spe-44 null microarray dataset (Supplementary Fig. 2b). This discrepancy may be due to SPE-44 depletion causing a reduction-of-function while the mutant is a null (Kulkarni et al. 2012). The transcription factor elt-1, a downstream target of spe-44, was found in both the microarray and RNA-seq datasets, as was spe-18, which was validated by promoter reporters in Kulkarni et al. (2012; spe-18 referred to as spe-7 in that paper; Table 2). Two msp genes (msp-49, msp-63) were common to the microarray and RNA-seq datasets (Table 2). There were 5 spe genes that were downregulated in our RNA-seq data, but not in the spe-44 microarray (Table 2). Three genes encode membrane proteins required for fertilization (spe-9, spe-42, spe-49), spe-43 is required for sperm activation, and spe-45 is required

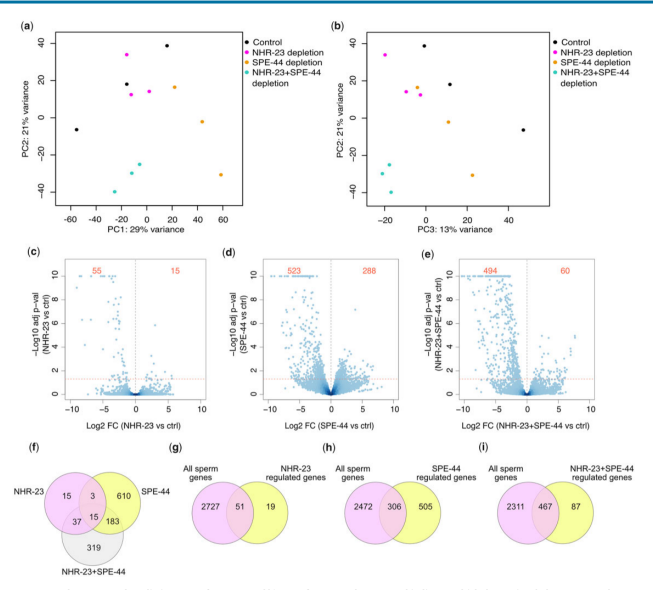


Fig. 2. NHR-23 and SPE-44 regulate distinct sets of genes. a and b) PCA of RNA-seq data. Legend indicates which data points belong to control, NHR-23depleted, SPE-44-depleted, or NHR-23+SPE-44-depleted biological replicates. c-e) Volcano plots of the indicated depletion datasets showing Log10 adjusted P-value against Log2 fold-change (FC). The horizontal dotted line represents our P-value cutoff of 0.05 and the vertical dotted line indicates a fold-change of 0. The red numbers in each figure indicate the number of differentially downregulated (left side) and upregulated (right side) genes in each comparison with a fold-change cutoff >1.5 (log2 = 0.58) and P < 0.05. f) Venn diagram depicting overlap of differentially regulated genes following NHR-23-, SPE-44, and NHR-23+SPE-44 depletion. Venn diagrams depicting overlap between sperm-enriched genes (Ortiz et al. 2014) and differentially regulated genes following NHR-23 depletion (g), SPE-44-depletion (h), and NHR-23+SPE-44 depletion (i).

for sperm to fuse with the oocyte plasma membrane (Singson et al. 1998; Zannoni et al. 2003; Kroft et al. 2005; Nishimura et al. 2015; Krauchunas et al. 2018; Wilson et al. 2018; Takayama et al. 2021). While our current focus in this study is on how NHR-23 promotes spermatogenesis, exploring the commonalities and differences between our RNA-seq data and the microarray data is an important future direction to further elucidate how SPE-44 promotes sperm development.

There were 326 differentially regulated genes unique to NHR-23+SPE-44-depleted males, consistent with the additive phenotype when we deplete both SPE-44 and NHR-23 (Ragle et al. 2020; Fig. 2). These genes included 14 msp genes, as well as smz-1, the paralog of NHR-23-regulated smz-2 (Table 2, Supplementary Table 7). There were also 3 spe genes downregulated when both NHR-23 and SPE-44 were depleted (Table 2). The presenilin paralog SPE-4 localizes to FB-MOs and is required for cytoplasmic

partitioning during spermatogenesis (L'Hernault et al. 1988; Arduengo et al. 1998). A spe-4 promoter reporter was used by Kulkarni et al. to validate their microarray data and confirm that it was not regulated by spe-44. The paternal effect factor SPE-11 prevents polyspermy and is required for eggshell formation (Hill et al. 1989), and SPE-15, a myosin VI homolog expressed in primary spermatocytes, is required for asymmetric partitioning of components such as FB-MOs and mitochondria during residual body formation (Kelleher et al. 2000; Hu et al. 2019).

Fifteen differentially regulated genes were common to NHR-23-, SPE-44-, and NHR-23+SPE-44-depleted animals (Fig. 2). Three of these genes were MFP2 paralogs (nsph-3.2, nsph-4.1, nsph-4.2; Table 2). MFP2 enhances MSP polymerization in Ascaris sperm, countering the activity of MFP1 (Grant et al. 2005), and there are 8 MFP2 paralog genes in C. elegans which have been assigned the nematode-specific peptide family, group H (nsph) gene class

Table 2. Differential regulation of select genes implicated in spermatogenesis.

	Log2 fold change following depletion of:			
Gene	NHR-23	SPE-44	NHR-23+SPE-44	Function
msd-1	-2.51 (1.02E-06)	_	-3.13 (1.62E-11)	MFP1 paralog. Potential role in slowing MSP polymerization
msd-2	-2.42 (6.87E-07)	_	-3.11 (1.51E-12)	MFP1 paralog. Potential role in slowing MSP polymerization
msd-3	-2.14 (4.38E-04)	_	-2.73 (1.51E-12)	MFP1 paralog. Potential role in slowing MSP polymerization
smz-1	_	_	-3.48 (6.40E-08)	Chromatin-bound protein. RNAi causes meiotic metaphase arrest.
smz-2	-5.00 (1.43E-30)	_	-5.16 (6.59E-33)	Chromatin-bound protein. RNAi causes meiotic metaphase arrest.
elt-1	<u> </u>	-1.12 (4.77E-02)	-1.21 (3.78E-02)	Transcription factor that regulates msp gene expression. Regulated by spe-44 in Kulkarni et al. (2012) microarray.
msp-49	_	-1.51 (9.29E-03)	-2.57 (2.55E-08)	MSP gene. Regulated by spe-44 in Kulkarni et al. (2012) microarray.
msp-63	_	-2.17 (8.50E-06)	-1.56 (7.07E-03)	MSP gene. Regulated by spe-44 in Kulkarni et al. (2012) microarray.
spe-4	_	_	-1.74 (2.64E-02)	Required for cytoplasmic partitioning.
spe-9	_	-3.74 (2.23E-04)	-5.71 (2.96E-07)	Membrane protein required for fertilization.
spe-11	_	_	-1.23 (1.31E-02)	Prevents polyspermy and required for eggshell formation
spe-15	_	_	-2.61 (1.81E-07)	Myosin IV homolog required for cytoplasmic partitioning.
spe-18	_	-1.59 (5.97E-03)	-5.50 (8.95E-35)	Promotes MSP assembly within FBs. Regulated by spe-44 in Kulkarni et al. (2012) microarray.
spe-42	_	-2.04 (3.31E-02)	-3.35 (3.13E-05)	Membrane protein required for fertilization.
spe-43	_	-3.78 (3.67E-04)	-6.99 (1.80E-05)	Required for sperm activation.
spe-45	_	-4.44 (1.13E-06)	-4.77 (1.56E-07)	Required for spermatozoa fusion with the oocyte plasma membrane.
spe-49	_	-3.25 (7.53E-05)	-6.25 (3.48E-09)	Membrane protein required for fertilization.
nsph-1.1	_	_	-2.18 (2.90E-03)	MFP1 paralog. Potential role in promoting MSP polymerization
nsph-2	_	-1.70 (1.05E-03)	_	MFP1 paralog. Potential role in promoting MSP polymerization
nsph-3.1	_	-1.26 (3.06E-02)	-1.53 (3.59E-03)	MFP1 paralog. Potential role in promoting MSP polymerization
nsph-3.2	-1.52 (1.50E-02)	-2.26 (4.17E-07)	-4.94 (1.59E-35)	MFP1 paralog. Potential role in promoting MSP polymerization
nsph-4.1	-8.32 (3.71E-67)	-5.97 (1.16E-40)	-8.35 (9.11E-68)	MFP1 paralog. Potential role in promoting MSP polymerization
nsph-4.2	-8.55 (1.00E-72)	-6.11 (6.61E-45)	-8.57 (2.36E-73)	MFP1 paralog. Potential role in promoting MSP polymerization

(Supplementary Fig. 3; Rödelsperger et al. 2021). nsph-2 is downregulated when SPE-44 is depleted, nsph-3.1 is found in both the SPE-44 and NHR-23+SPE-44 depleted differentially regulated genes, and nsph-1.1 is downregulated only when both NHR-23 and SPE-44 are depleted (Table 2). Two nsph genes (1.2, 4.3) are not found in any of the datasets. We note that nsph-4.3 has a large deletion that removes 500 bp of sequence, though a predicted protein product can still be produced (Supplementary Figs. 3 and 4). There is also a pseudogene MFP2 paralog (ZK546.7) downregulated following NHR-23-, SPE-44, and NHR-23+SPE-44 depletion. This pseudogene is related to nsph-4.1 and nsph-4.2 but an insertion causes a frameshift. An initiation codon further downstream could produce a valid coding sequence, so this pseudogene may be worth future exploration (Supplementary Fig. 4).

#### NHR-23 and SPE-44 targets are enriched in phosphatase genes

To gain insight into how NHR-23 and SPE-44 promote spermatogenesis, we performed gene ontology (GO) analysis on the differentially regulated genes to identify enriched biological processes. Phosphatases were enriched in all 3 depletion conditions (Table 3, Supplementary Tables 5-7). Notably, kinase and phosphatase genes are over-represented in sperm-enriched genes (Reinke et al. 2000). The NHR-23-regulated genes were also enriched in genes involved in ATP binding, though these genes overlapped with the phosphatase genes and cytoskeletal protein binding (Supplementary Table 8). SPE-44-regulated genes were enriched in genes involved in DNA replication (origin binding, helicase activity, single-strand DNA-dependent ATPase), carbohydrate-binding, glutamate-ammonia ligase activity, and calcium ion-binding (Supplementary Table 9). In addition to being enriched in phosphatase-related GO terms, the NHR-23+SPE-44-regulated genes were also enriched in kinase genes, NADPH reductase activity, FFAT motif binding, drug binding, and transaminase activity (Supplementary Table 10). Together, these data

implicate NHR-23 and SPE-44 in regulating signal transduction by kinases and phosphatases, which has been shown to be an important regulatory mode following transcriptional quiescence in spermatocytes (Muhlrad and Ward 2002; Bae et al. 2009; Wu et al.

#### NHR-23 regulates uncharacterized kinases and MFP2 paralogs

We focused on the NHR-23-regulated genes as they were few in number so therefore amenable to a candidate-based knock-out approach. In addition to the aforementioned phosphatase and nsph genes, NHR-23 also regulated 4 uncharacterized kinases (Table 3). Three of these kinases were predicted to be Casein Kinase I orthologs. W03G9.5 and C04G2.2 belong to the Tau Tubulin Kinase-Like subgroup, while ZK596.2 belongs to the nematode spe-6 subgroup (Manning 2005). We engineered predicted null mutant alleles through insertion of a "STOP-IN" cassette containing premature termination codons in all 3 reading frames in the first or second exon of targeted genes (Wang et al. 2018). We generated mutants in 1 phosphatase (ZK757.2), 2 kinases (ttbk-5, ZK596.2), and 3 MFP2 paralogs (nsph-2, nsph-3.2, nsph-4.3) all of which were viable with wild-type brood sizes (Table 4). The genes nsph-4.1 and nsph-4.2 are predicted to produce identical proteins, only differ by a single nucleotide in an intron, and have 1,231 bases of identical upstream promoter sequence. This sequence similarity made genotyping a STOP-IN insertion challenging, so we deleted the entire gene for each using crRNAs that bound outside the conserved sequence for each gene. Each mutant had a wild-type brood size and no obvious increase in unfertilized oocytes (Table 4). These data could suggest that these genes are dispensable for fertility, redundant with other genes, or the STOP-IN cassette is not creating a null. We created nsph-2; nsph-1.1 and nsph-4.1, nsph-4.2 double mutants, both of which had wild-type brood sizes.

Table 3. NHR-23-regulated kinases and phosphatases.

Gene	Log2 fold- change	Class	Ortholog
ZK596.2	-9.01	Predicted serine/threonine kinase	Tau-tubulin kinase 2 (human); Casein kinase I delta ortholog
W03G9.5	-8.19	Predicted serine/threonine kinase	Casein kinase delta (human) and Casein kinase I (yeast)
F47B3.2	-4.17	Predicted tyrosine phosphatase	Tyrosine-protein phosphatase nonreceptor type 6 isoform 3 (mammals)
F55H12.5	-4.04	Predicted tyrosine phosphatase	Receptor-type tyrosine-protein phosphatase T (mammals)
F47B3.7	-4	Predicted tyrosine phosphatase	Tyrosine-protein phosphatase nonreceptor type 6 isoform 3 (mammals)
ZK757.2	-3.49	Predicted tyrosine/serine/threonine phosphatase	Dual specificity phosphatase 14 (human)
D2045.7	-3.05	Predicted serine/threonine kinase	Serine/threonine kinase 16 (human)
F52H3.6	-2.7	Predicted serine/threonine phosphatase	Isoform Gamma-1 of Serine/threonine-protein phosphatase PP1-gamma catalytic subunit (humans)
ttbk-5	-2.01	Predicted serine/threonine kinase	Tau-tubulin kinase 2 (human); Casein kinase I delta ortholog

Table 4. Brood size data.

Genotype	Allele class	Predicted gene function	Brood size (±SD)	% unfertilized eggs
N2 (wt)	N/A	N/A	230 ± 91	3.2 (n = 2,891)
nspĥ-2(wrd57)	PTC	MFP2 paralog	$301 \pm 43$	2.75 (n = 2,184)
nsph-3.2(wrd56])	PTC	MFP2 paralog	$277 \pm 39$	2.34 (n = 3,420)
nsph-2(wrd57); nsph-3.2(wrd56)	PTC	MFP2 paralog	$293 \pm 39$	1.5 (n = 3,276)
nsph-2(wrd57; nsph-1.1(wrd67)	PTC	MFP2 paralog	$185 \pm 40$	1.07 (n = 2,241)
nsph-4.3(wrd74)	PTC	MFP2 paralog	$277 \pm 49$	3.08 (n = 3,445)
nsph-4.1 (wrd92)	Whole gene deletion	MFP2 paralog	$228 \pm 45$	5.75 (n = 2,677)
nsph-4.2(wrd93)	Whole gene deletion	MFP2 paralog	$219 \pm 35$	4.15 (n = 2,750)
nsph-4.1 (wrd92), nsph-4.2 (wrd97)	Whole gene deletion	MFP2 paralog	$260 \pm 49$	4.64 (n = 3,296)
ttbk-5(wrd72)	PTC	Kinase	$259 \pm 25$	6.84 (n = 3,373)
ZK596.2(wrd71)	PTC	Kinase	$231 \pm 80$	2.93 (n = 2,931)
ZK757.2(wrd77)	PTC	Phosphatase	$242 \pm 17$	3.31 (n = 2,999)

#### NHR-23-depletion causes NSPH-2 and MSD-1 localization defects

As previously described, both NSPH-2 and MSD-1 antibodies label FBs and colocalize with MSP antibody labeling in wild-type 1° spermatocytes (Morrison et al. 2021; Fig. 3, a and b). In other studies (Kulkarni et al. 2012; Ragle et al. 2020), we found that single and double NHR-23 and SPE-44 germline depletion have distinct effects on FB-MO formation. Single depletions of either NHR-23 or SPE-44 fully disrupt the assembly of MSP into FBs but differentially impact the MO component of the FB-MO complex (Kulkarni et al. 2012; Ragle et al. 2020). When labeled with the MO differentiation marker 1CB4, NHR-23 depleted spermatocytes exhibit distinct labeled structures, whereas 1CB4 labeling is diffuse in SPE-44 depleted spermatocytes and absent in the NHR-23+SPE-44 depleted spermatocytes (Ragle et al. 2020). To further explore these interactions, we tested how depleting NHR-23, SPE-44, and NHR-23+SPE-44 from the germline would affect the localization of other FB components such as NSPH-2 and MSD-1. In each of our depletion strains, germline labeling with anti-NSPH-2 and anti-MSD-1 was dim and diffuse, compared to the bright puncta labeled in control germlines, indicating that assembly of NSPH-2 and MSD-1 into FBs was impaired in the absence of NHR-23, SPE-44, or both. Although our RNA-seq data suggest that SPE-44 and not NHR-23 regulates nsph-2 (Table 2), terminal SPE-44 depleted spermatocytes exhibited NSPH-2 aggregates that were not observed in either NHR-23 or the double-depleted strains (Fig. 3a). Due to this unexpected phenotype, we also investigated spe-44 null (ok1400) mutants (Kulkarni et al. 2012). Auxin-inducible depletion of target proteins is robust, but may not be fully penetrant in some cases, so the null mutant provides additional insight into spe-44's role in FB assembly. Consistent with our RNA-seq results, anti-NSPH did not label germlines in spe-44(ok1400) animals

(Fig. 3a'). As predicted from our RNA-seq data (Table 2), antibody labeling with anti-MSD-1, which labels all MSD isoforms (Kosinski et al. 2005), was most diminished in NHR-23 and NHR-23+SPE-44 depleted spermatocytes, whereas terminal SPE-44 depleted spermatocytes exhibited dim but detectable MSD-1 punctae (Fig. 3b). In both single and double depletions, the MSD-1 antibody labeled centrosomes (arrows), but the significance of this pattern is unclear (Fig. 3b). It may be nonspecific as many preimmune sera label the centrosomes of nematode spermatocytes. Alternatively, in the absence of normal FBs, perhaps centrosomes serve as a secondary localization site for the remaining MSD-1 protein. These data indicate that NHR-23 is either required for the stable expression of NSPH-2 and MSD-1 or for localization of NSPH-2 and MSD-1 to the FB-MO complex, which is consistent with our previous finding that NHR-23 depletion prevents MSP loading into FBs (Ragle et al. 2020).

#### SPE-6 patterns are differentially affected in single and double knockdowns

Given our finding that NHR-23 regulates 3 uncharacterized Casein Kinase I orthologs, W03G9.5, C04G2.2, and ZK596.2, which belong to the nematode spe-6 subgroup, we decided to investigate the localization of SPE-6, the better characterized casein kinase 1 ortholog, in our depletion strains. In C. elegans, spe-6 encodes one of a large family of casein kinase 1 proteins (Muhlrad and Ward 2002). Like NHR-23 and SPE-44 depletion, SPE-6 null mutations result in primary spermatocyte arrest (Varkey et al. 1993). During wild-type spermatogenesis, SPE-6 shifts from a diffuse plus particulate pattern in spermatocytes to being localized around the condensed haploid chromatin mass during the budding division that follows anaphase II (Peterson et al. 2021; Fig. 4a). Although spe-6 transcript levels were not significantly altered in either the

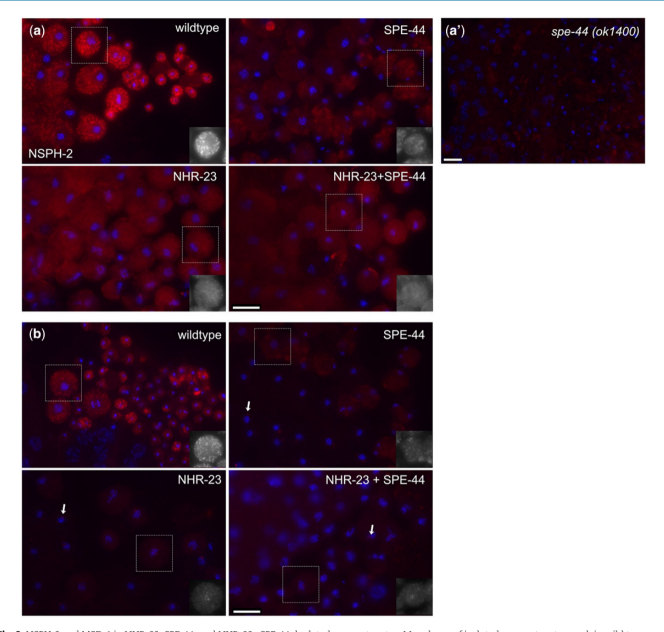


Fig. 3. NSPH-2 and MSD-1 in NHR-23, SPE-44, and NHR-23+SPE-44 depleted spermatocytes. Monolayer of isolated spermatocytes; and, in wild type, budding figures and spermatids labeled with DAPI (blue) and either anti-NSPH-2 antibody (a) or anti-MSD-1 antibody (b) (red). Images are oriented with earlier stages on the left. In the images shown, wild-type spermatocytes are from JDW92 males that contain the nhr-23 degron constructs but were not auxin-treated (a, b). The other images are from males in which the indicated transcription factor has been depleted by auxin-mediated degradation. Images are representative of multiple preparations, but images shown are from the same preparation and taken with the same exposures without additional adjustments. a') Monolayer of isolated spermatocytes from spe-44(ok1400) null mutant labeled with anti-NSPH-2 antibody. b) Arrows show labeled centrosomes. In a and b, black and white inserts of indicated spermatocytes are shown for clarity. Scale bars = 10 µm.

single or double depletion strains (Supplementary Table 1), we tested whether either the SPE-6 localization patterns or levels of abundance would be differentially altered when we used anti-SPE-6 antibody to label the proximal germlines of affected males. In NHR-23 depleted spermatocytes, SPE-6 remains in a diffuse plus particulate pattern, consistent with its early meiosis I arrest (Fig. 4b). In contrast, in SPE-44 depleted spermatocytes, SPE-6 clumps around the chromatin masses in a postmeiotic-like pattern even though the spermatocytes never physically divide to form haploid spermatids (Fig. 4c, Kulkarni et al. 2012). Anti-SPE-6 labeling in NHR-23+SPE-44 depleted germlines exhibited 2 distinct phenotypes: most (51/67) were similar to the NHR-23 single depletion but with overall reduced levels and cell-to-cell variability (Fig. 4d) whereas others (16/67) failed to label with SPE-6

antibodies (Fig. 4e). These data suggest that the developmental program of restructuring the sperm chromatin continues upon SPE-44 depletion but not upon NHR-23 depletion, and that SPE-6 expression is reduced or lost upon NHR-23+SPE-44-depletion.

#### **Discussion**

This work identifies genes differentially regulated following NHR-23-depletion in male germlines, providing insight into how NHR-23 promotes spermatogenesis. We also identify genes that are differentially expressed only when both NHR-23 and SPE-44 are depleted in male germlines. This list provides an entry point to understanding the synthetic phenotype associated with depleting both transcription factors (Ragle et al. 2020). NHR-23 depletion

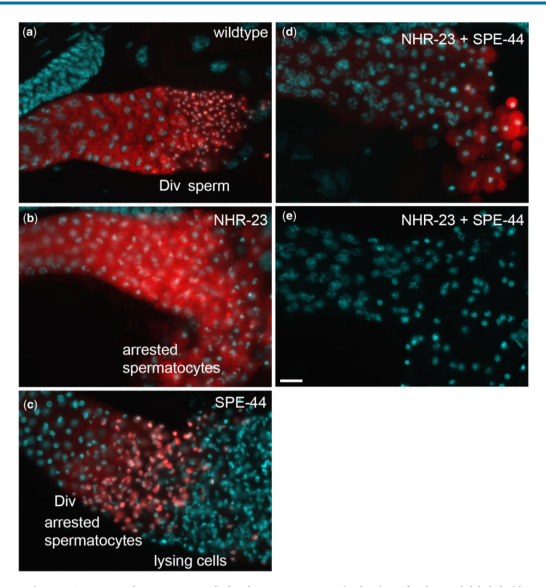


Fig. 4. SPE-6 patterns in NHR-23, SPE-44, and NHR-23+SPE-44 depleted spermatocytes. Proximal regions of male gonads labeled with DAPI (cyan) and anti-SPE-6 antibody (red). Images are oriented with earlier stages on the left. a) Wild-type controls are auxin-treated him-5 males. b-e) The other images are from males in which the indicated transcription factor has been depleted by auxin-mediated degradation. The regions of the gonad are labeled including the meiotic division zone (DIV). Images are representative of multiple preparations, but images shown are from the same preparation and taken with the same exposures without additional adjustments. Scale bars = 10  $\mu m$ .

causes defective localization of NSPH-2 and MSD-1 to FBs, implicating NHR-23 in regulating MSP polymerization dynamics. Although NHR-23 and SPE-44 do not appear to transcriptionally regulate the Casein kinase spe-6, double depletion of these transcription factors causes a partial or severe loss of SPE-6 antibody labeling, suggesting that SPE-6 protein levels are compromised when NHR-23 and SPE-44 are absent from the germline.

#### NHR-23 regulates few genes with known roles in spermatogenesis

Relatively few genes were differentially regulated following NHR-23-depletion (Supplementary Table 5; 70 genes) compared to SPE-44-depletion (Supplementary Table 6; 811 genes), NHR-23+SPE-44-depletion (Supplementary Table 7; 554 genes), or nhr-23(RNAi) in L2 larvae (Kouns et al. 2011). Aside from smz-2, none of the NHR-23-regulated genes have been experimentally confirmed to play roles in spermatogenesis.

Terminally arrested spermatocytes from NHR-23 depleted animals arrest early in meiosis and contain distended MOs, similar to spe-6(lf) and wee-1.3(gf) mutants (Varkey et al. 1993; Lamitina and L'Hernault 2002), yet NHR-23 does not regulate these genes (Supplementary Tables 1 and 5). The dearth of known spermatogenesis regulators is consistent with NHR-23 regulating a novel pathway promoting sperm development. As NHR-23 regulates several uncharacterized kinases and phosphatases, one possibility is that NHR-23 controls meiotic progression and FB-MO biogenesis through kinase-mediated signal transduction. The orthologs of 2 of the kinases identified in our RNA-seq dataset (ttbk-5, ZK596.2) are implicated in mammalian microtubule dynamics and yeast vesicular traffic, meiotic exit, and kinetochore attachment in meiosis II (Petronczki et al. 2006; Goetz et al. 2012; Lee et al. 2017). While our STOP-IN insertions did not have phenotypes, there could be redundancy or genetic compensation, a point which we will expand on in the next section.

#### NHR-23 is necessary for MSD-1 and NSPH-2 localization

Our immunostaining revealed that NHR-23 depletion caused a loss of MSD-1 and NSPH-2 localization in spermatocytes. The orthologs of these factors (MFP1 and MFP2) in Ascaris are specifically known to control MSP polymerization within the pseudopod of crawling sperm (Buttery et al. 2003; Grant et al. 2005) but since they are also present in FBs (Morrison et al. 2021), their downregulation upon NHR-23 depletion might account for the observed defect in MSP loading into FB-MOs (Ragle et al. 2020). The effect of NHR-23 on NSPH-2 localization was surprising as this gene was not differentially regulated following NHR-23 depletion (Table 1, Supplementary Table 5). nsph-2 was downregulated following SPE-44 depletion (Table 1, Supplementary Table 5); and by immunocytology, the protein expression appeared slightly reduced in SPE-44-depleted germlines and significantly reduced in spe-44(ok1400) null mutants (Fig. 3). These data suggest that NHR-23 could promote NSPH localization to FB-MOs. There are 4 uncharacterized MSD1 paralogs in C. elegans, all of which are recognized by the MSD-1 antibody and 3 of which are regulated by NHR-23 (Table 1, Supplementary Table 5). There are 8 MFP2 paralogs, which are also uncharacterized. Three of these paralogs are regulated by NHR-23 while 5 are SPE-44-regulated (Table 2). No phenotype has been previously reported for the C. elegans MFP1 or MFP2 paralogs, though redundancy may be responsible since these paralogs appear to have arisen through duplication. We have inactivated 6 of the MFP2 paralogs either singly or in double mutant combinations and have seen no difference in fertility. Three possible explanations for the lack of phenotype are: (1) that these genes do not play a role in C. elegans spermatogenesis; (2) extensive genetic redundancy exists and we will have to inactivate more paralogs, if not all, to produce a phenotype; or (3) that our knock-out approach caused genetic compensation, where mRNAs containing premature stop codons trigger the upregulation of paralogous genes (Rossi et al. 2015; El-Brolosy et al. 2019). We favor the second and third possibilities. Going forward, it will be important to avoid genetic compensation by systematically deleting MFP1 and MFP2 orthologs or use a multiplexed piRNA-based knockdown approach to determine whether these genes play roles in C. elegans MSP polymerization dynamics (Priyadarshini et al. 2022).

#### A high-confidence set of spe-44-regulated genes

There was a surprising lack of overlap between the SPE-44regulated genes in our RNA-seq dataset and differentially regulated genes in a spe-44 mutant microarray dataset (Kulkarni et al. 2012), though the genes common to both datasets are highconfidence spe-44 targets. The difference could be biological as the spe-44 mutant is a predicted null, while auxin-mediated protein depletion frequently produces hypomorphic phenotypes as we also observed in our NSPH-2 immunocytology. Our data were generated from adult males, while the spe-44 mutant dataset was generated from L4 males, so it is also possible that differences in developmental stage impacted the results. While RNA-seq is reported to have a wider quantitative range in detecting changes in expression levels, a comparison of microarrays vs RNA-seg on rat cDNA reported a 78% overlap in differentially regulated genes (Rao et al. 2019). The genes common to both datasets include elt-1, a transcription factor that acts downstream of SPE-44 to regulate MSP gene expression, and ceh-48 (del Castillo-Olivares et al. 2009; Kulkarni et al. 2012). ceh-48 is an uncharacterized ONECUT homeobox transcription factor ortholog that Kulkarni et al. (2012) identified as sperm-enriched. A deletion mutant is reported to be fertile but displays male tail defects (Kim et al. 2016). spe-18, which promotes MSP assembly within FBs, and the MFP2 paralog nsph-2 were also common to both datasets (Price et al. 2021). Exploring the high confidence target will provide insight into how SPE-44 promotes FB-MO assembly and meiosis II.

#### SPE-6 labeling is reduced or lost in NHR-23+ SPE-44 depleted germlines

SPE-6 is a casein kinase that plays broad roles in spermatogenesis including FB assembly, meiotic progression, and repression of precocious sperm activation (Varkey et al. 1993; Muhlrad and Ward 2002; Peterson et al. 2021; Price et al. 2021). spe-6 was not differentially regulated in any of our datasets, despite the overlap of the NHR-23-depletion and spe-6(lf) meiotic arrest and distended FB-MO phenotype (Varkey et al. 1993; Ragle et al. 2020). SPE-6 labeling in NHR-23-depleted animals appeared in a punctate pattern, as well as diffuse in the cytosol, as expected in metaphase I arrested spermatocytes (Ragle et al. 2020; Fig. 4c). In SPE-44-depleted germlines, SPE-6 labeling appeared similar to a postmeiotic pattern, despite the meiosis II arrest and lack of spermatocytes (Kulkarni et al. 2012; Fig. 4b). These data suggest that progression through metaphase I is required for SPE-6 to localize around chromatin after meiosis, and that this localization and presumably SPE-6 pathway is insensitive to SPE-44 depletion and the meiosis II arrest. Surprisingly, NHR-23+SPE-44 double-depletion caused a partial or complete loss of SPE-6 labeling, another example of a synthetic phenotype caused by depletion of these transcription factors (Fig. 4, d and e). spe-6 transcript levels were not significantly affected by NHR-23+SPE-44 depletion (Supplementary Table 1), indicating that this SPE-6 downregulation is not transcriptional. Possible mechanisms could include impaired translation of spe-6 mRNA or an increase in SPE-6 turnover.

#### Future perspectives

We previously discovered a novel role for NHR-23 in spermatogenesis and a synthetic interaction between NHR-23 and SPE-44 (Ragle et al. 2020). We also proposed that a yet-undetermined pathway promoted spermatogenesis independently of NHR-23 and SPE-44 (Ragle et al. 2020). In agreement with this assertion, there are a large number of sperm-enriched genes not found in our differentially regulated gene lists, and there are genes only differentially regulated following NHR-23+SPE-44 depletion, consistent with synthetic interaction. Numerous experimentally validated spe genes are not regulated by NHR-23 or SPE-44. Our RNAseq data provide an entry point to understanding transcriptional control of C. elegans spermatogenesis and how NHR-23 and SPE-44 interact to promote FB-MO biogenesis.

#### Data availability

The data underlying this article are available in the article and in its online supplementary material. Strains and plasmids are available upon request. Supplementary File 1 contains the genomic sequence of wild-type and genome-edited loci. The RNA-seq data have been deposited in the National Center for Biotechnology Information Gene Expression Omnibus under accession number GSE206676. Supplementary Table 1 contains the processed data with gene name and location, mean expression, log2 fold-change compared to control, and P-value. Sequences and descriptions of the crRNAs and repair oligos are provided in Supplementary Tables 2 and 3, respectively. Genotyping primers are provided in Supplementary Table 4.

Supplemental material is available at G3 online.

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#### **Conflicts of interest**

None declared.

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