Linking transcriptomics and proteomics in spermatogenesis

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Abstract

Spermatogenesis is a complex and tightly regulated process leading to the continuous production of male gametes, the spermatozoa. This developmental process requires the sequential and coordinated expression of thousands of genes, including many that are testis-specific. The molecular networks underlying normal and pathological spermatogenesis have been widely investigated in recent decades, and many high-throughput expression studies have studied genes and proteins involved in male fertility. In this review, we focus on studies that have attempted to correlate transcription and translation during spermatogenesis by comparing the testicular transcriptome and proteome. We also discuss the recent development and use of new transcriptomic approaches that provide a better proxy for the proteome, from both qualitative and quantitative perspectives. Finally, we provide illustrations of how testis-derived transcriptomic and proteomic data can be integrated to address new questions and how the ‘proteomics informed by transcriptomics’ technique, by combining RNA-seq and MS-based proteomics, can contribute significantly to the discovery of new protein-coding genes or new protein isoforms expressed during spermatogenesis.

Introduction: unraveling testis specificities with omics technologies

From a genomist’s point of view, spermatogenesis, especially in mammals, is arguably one of the most exciting objects of study available. Not only does this developmental process offer incredible molecular dynamics, but it also embodies several expression specificities and striking genomic features.

To make motile sperm capable of fertilization, germ cells must undergo unique processes, such as meiosis, and develop specific organelles and cell structures, including the acrosome, flagellum, and a highly condensed nucleus. This extreme differentiation process involves the functions of specific molecular factors, many of them expressed only during spermatogenesis. High-throughput tissue-profiling experiments thus regularly identify the testis as the organ that expresses the greatest number of tissue-specific genes and proteins (Chalmel et al. 2007, 2012, Kouadjo et al. 2007, Fagerberg et al. 2014, Uhlen et al. 2015). Similarly, the finding that the testis contains the highest number of alternative splicings (Xu et al. 2002, Yeo et al. 2004, Kan et al. 2005, De la Grange et al. 2010) indicates that what is true for genes and proteins also applies to isoforms.

Evolutionarily speaking, genes involved in male germ cell development are also quite remarkable. For instance, testis-expressed genes show the highest divergence rate between species for both sequence and expression (Khaitovich et al. 2005, Voolstra et al. 2007). Additionally, testicular transcripts have, yet again, the highest number of diverged alternative splicings (Kan et al. 2005). This fast evolution of male fertility-related factors is thought to result from sexual selection, a specific pressure selection that enables mutations providing a reproductive advantage to be transmitted more easily to progeny and thus fixed within a species relatively quickly.

Another striking genomic feature observed during germ cell development concerns sexual chromosomes and especially X-linked genes (for review, see Hu & Namekawa (2015)). Throughout the evolution of eutherian species, the Y chromosome has become progressively shorter, thus preventing the X and Y chromosomes from aligning/pairing with any precision during meiotic prophase I. Accordingly, to prevent misalignments and recombinations, sexual chromosomes condense into a specific nuclear structure named the sex- or XY-body (Solari 1974, McKee & Handel 1993). The strong condensation of X and Y chromosomes within this structure leads to their complete transcriptional silencing (Turner 2007). This so-called meiotic sex chromosome inactivation (MSCI) has been demonstrated clearly at the genome-wide
level: several high-throughput expression studies have failed to find the expression of a single X-linked gene during the meiotic phase of spermatogenesis (Namekawa et al. 2006, Chalmel et al. 2007). Additionally, to compensate for the cessation of transcription of crucial genes for any cell, a large number of X-linked genes have, over the course of evolution, been transposed onto autosomes and acquired specific meiotic and post-meiotic expression (Potrzebowski et al. 2008, 2010). Finally, because the heterozygous nature of sexual chromosomes in males allows advantageous reproductive traits to be fixed quickly, X-linked genes are overrepresented among those preferentially expressed in testicular somatic cells, spermatogonia, and post-meiotic spermatids (Khil et al. 2004, Chalmel et al. 2007, Mueller et al. 2008).

Taken together, the specificities of the male germ cell expression program provide a rich environment for studying regulatory mechanisms of gene expression at various levels as well as for the discovery of new genes and protein isoforms. As many past studies investigating spermatogenesis with high-throughput approaches have been reviewed elsewhere (Rolland et al. 2008, Calvel et al. 2010, Chocu et al. 2012), here we focus on studies that have attempted to link the transcriptome and proteome in spermatogenesis or have combined transcriptomic and proteomic data to gain insight into testicular functions and germ cell biology (Table 1).

**Integrative omics strategies to study spermatogenesis**

The integration of various types of omics data, e.g. epigenomic, transcriptomic, proteomic, interactomic, or regulomic, represents a powerful tool for going far beyond basic descriptive analyses. Combining information from different samples and technologies makes it possible to improve data consistency by refining candidate selection, as well as to address more specific questions and to build new hypotheses (Fig. 1A).

For example, microarray and proteomic data from mouse testes lacking DICER in the Sertoli cells (DCR<sup>fx/fx</sup>;MisCre) were compared to investigate the miRNA-mediated post-transcriptional regulation in these cells (Papaioannou et al. 2011). This approach allowed the authors to identify miRNA-targets within Sertoli cells, i.e. proteins whose abundance increases in KO mice, even

### Table 1 Overview of studies linking transcriptomics and proteomics in spermatogenesis.

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CLIP, cross-linking immuno-precipitation; ICPL, isotope-coded protein labeling; iTRAQ, isobaric tags for relative and absolute quantitation; RNPs, ribonucleoprotein particles; MS/MS, tandem mass spectrometry; MudPIT, multidimensional protein identification technology.
Linking transcriptomics and proteomics

Integrative omics approaches

- Transcriptomic analysis
  - Transcript candidates
- Proteomic analysis
  - Protein candidates

Proteomics informed by transcriptomics

- Sample #A
  - RNA-seq
  - Transcription isoforms
  - Quantification comparison
  - Protein candidates
- Sample #B
  - MS/MS data
  - Derived protein DB
- Sample #C
  - Sample #C'
  - Transcript isoforms
  - Protein identification

Overlaps, discrepancies...

Figure 1 When transcriptomics meets proteomics. (A) Typical integrative omics approaches involve the combination of datasets originating from various technologies, most notably transcriptomics and proteomics. Such strategies are often used to identify more reliable candidates (i.e., factors evidenced at both the RNA and protein levels), but they can also be useful in order to compare and correlate transcription and translation rates. More recently, the combination of RNA-seq and mass spectrometry (MS)-based proteomics has led to the development of the so-called proteomics informed by transcriptomics (PIT) approach. In this approach, the protein sequence database (DB) queried for protein identification purpose is directly derived from transcript sequences obtained following RNA-seq analysis of the same or equivalent sample as that used for MS/MS analysis. (B) The characterization of the transcriptome has long been used as a proxy for the proteome. However, depending on whether nuclear, total cytoplasmic, or ribosome-bound RNAs are analysed, the captured picture will reflect either more the transcriptional rate or the translational rate. Additionally, while both approaches perform equivalently from a quantitative point of view, RNA-seq overcomes microarray technology from a qualitative point of view as it allows full-length transcript reconstruction and can thus discriminate between distinct protein-coding isoforms.

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though the expression of their corresponding mRNAs does not change. Subsequently, they performed 3'UTR luciferase assays to validate SOD-1 as a likely direct target of miR-125a-3p, miR-872 and miR-24.

Many studies have also combined proteomic and transcriptomic data to improve the characterization of the expression landscape during spermatogenesis. Chalmel et al. (2012) using biopsies from infertile patients with spermatogenesis arrested at various stages of germ cell development, first identified genes preferentially expressed in each type of testicular cell. Gene expression data from various tissues and antibody-based profiling data from the Human Protein Atlas (http://www.proteinatlas.org; Uhlen et al. 2010) were then used to identify, respectively, the genes and gene products showing testis-specific expression; most of them were found to be expressed in meiotic and post-meiotic germ cells. Finally, taking advantage of available information on protein-gene interactions (i.e., regulomic data), these authors filtered for a core network of transcription factors and DNA-binding proteins that are likely to drive the germ cell-specific expression program. Conversely, Djureinovic et al. (2014) sought to determine the human testis-specific proteome, beginning with the RNA-sequencing (RNA-seq) analysis of 27 tissues. They then interrogated the Human Protein Atlas about the testis-enriched transcripts they identified to confirm the testis-specificity of corresponding protein products and to identify the testicular cell type(s) in which they were expressed.

The combination of transcriptomic and proteomic approaches has also been very successful in helping to interpret the content of biological fluids or transcriptionally inactive cells, such as spermatozoa. For instance, Rolland et al. (2013) compiled several human seminal plasma proteomic studies and compared the resulting proteome to gene expression data for the organs contributing to this biological fluid, i.e. the testis, epididymis, seminal vesicle, and prostate. This allowed the identification of protein biomarkers for each of the male genital tract organs. Importantly, these biomarkers, including germ cell markers, can thus be monitored in semen.

A recent investigation of the intricate question of Sertoli–germ cell communication used another such integrative approach (Chalmel et al. 2014a). It analyzed the testicular fluid proteome of rats and rams and then combined it with transcriptomic data from isolated testicular cells (Chalmel et al. 2007) and with protein–protein interaction data. The authors were therefore able to identify testicular fluid proteins likely to be secreted by Sertoli cells and to interact with germ cell membrane proteins and, conversely, proteins secreted by germ cells that might interact with Sertoli cell membrane proteins. Among these, the interactions of APOH and CDC42 as well as APP and NGFR were further investigated and validated in situ. Finally, Wang et al. (2013) compiled...
several proteomic studies to determine the proteome of human spermatozoa and used gene expression tissue-profiling data to identify the sperm proteins specifically expressed in the testsis. With this candidate list, they next queried the drug target information available in Drugbank (Wishart et al. 2008) to identify potential male contraceptive molecules. Disulfiram and propofol, two molecules thought to target cilia proteins, were then shown to inhibit sperm motility.

**Correlating transcription and translation rates during spermatogenesis**

Transcriptomic studies often assume that the abundance of mRNAs and that of their corresponding proteins are well correlated. This hypothesis is considered to justify extrapolation from changes in gene expression to changes in protein expression and, ultimately, to their potential functional impact. The rationale of this hypothesis may appear quite reasonable: typical microarray or RNA-seq protocols involve an oligo-dT selection of polyadenylated mRNAs, which are thought to be actively translated, unlike those that are not polyadenylated.

The fate of an mRNA, however, is tightly regulated by a complex interplay of modification, processing, storage, decay, and translation, all involving protein-RNA interactions through messenger ribonucleoprotein (mRNP) complexes. Some of these assembled complexes are conducted directly to translation while others are diverted towards storage and translational repression (for review, see Müller-McNicol & Neugebauer (2013)).

While post-transcriptional and translational gene regulation is a common mechanism in all cell types, it is especially striking during spermatogenesis. As spermatids start to elongate, nuclear histones are sequentially replaced by transition proteins and protamines (for review, see Rathke et al. (2014)). This substitution allows the progressive condensation of chromatin and thus leads to the complete cessation of transcription from mid-spermiogenesis onwards (Kierszenbaum & Tres 1975). As a consequence, many genes that are required for the development and/or functioning of spermatozoa are transcribed much earlier during germ cell differentiation, then translationally repressed, and finally translated several days after the mRNA production, thanks to a complex interplay of RNA-binding proteins and non-coding RNA (for review, see Kleene (2013)). In this context, it is interesting to note the existence of the chromatid body, a germ cell-specific RNA processing center suggested to be involved in the sequestration and translation repression of several mRNAs during spermiogenesis (Kotaja & Sassone-Corsi 2007) and whose RNA and protein content was recently analyzed (Meikar et al. 2014). However, the direct contribution of this organelle to translational regulation remains to be clearly demonstrated (for review, see Kleene & Cullinan (2011)).

More importantly, the use of cross-linking immuno-precipitation (CLIP) together with microarray analysis (CLIP-chip) or high-throughput sequencing (HITS-CLIP or CLIP-seq) has allowed some potential direct targets of RNA-binding proteins to be identified in male germ cells (Reynolds et al. 2005, Grellscheid et al. 2011, Vourekas et al. 2012, Zhang et al. 2015). The combination of such approaches with proteomic analyses of mutant mice would in turn help identify which mRNAs are actually translationally regulated by these specific factors during spermatogenesis.

Because of this prominent uncoupling between transcription and translation, the testsis is often seen as an organ in which transcriptome and proteome are not necessarily linked. This low correlation between mRNA and protein concentrations within the testsis was clearly evidenced in a tissue-profiling experiment that used multidimensional protein identification technology (MudPIT) for human tissue (Cagney et al. 2005). In this study, the abundance of 683 proteins and their corresponding transcripts were measured and compared in eight organs. Interestingly, the gene profiles of all organs clustered together, as did their protein profiles. This finding suggests that transcriptomes or proteomes from different organs are more similar than the transcriptome and proteome of the same organ. Importantly, the correlation between transcriptome and proteome data was weakest for the testsis and highest for the liver (correlation coefficients of 0.138 and 0.432 respectively). To investigate the relation between proteins and mRNA levels during spermatogenesis in more detail, Gan et al. (2013a) used isolated type A spermatogonia, pachytene spermatocytes, round spermatids, and elongated spermatids for an iTRAQ-based proteomic analysis of male germ cell differentiation and compared their results with those of a previously published microarray dataset (Namekawa et al. 2006). While they found a consistent match for a subset of transcriptomic and proteomic profiles, they also observed that several regulation mechanisms – including transcript degradation, translation repression, translation de-repression, and protein degradation – affected most genes and may account for the low correlation between mRNAs and proteins, at both the mitosis/meiosis transition (Pearson correlation of 0.55) and the meiosis/post-meiosis transition (Pearson correlation of 0.41).

**Investigating the translatome of testicular cells**

Another exciting possibility for bridging the gap between gene expression and protein abundance lies in methods that allow investigation of the translatome, i.e. the measurement of transcripts that are actively processed by translational machinery (Fig. 1B). These analyses usually involve the purification of ribosomes or polysomes and the subsequent measurement of associated transcripts.
Iguchi et al. (2006) first applied such an approach to address the question of translational regulation during the meiotic and post-meiotic phases of male germ cell development. The authors monitored the mRNAs associated with free RNPs and polysomes in the testes from mice at various postnatal stages and identified translationally up- and down-regulated transcripts, i.e. mRNAs significantly redistributed between free RNPs and the polysomal fraction during testis development. Not surprisingly, translation increased for most of these mRNAs, in elongating spermatids. This increase reflects a common mechanism compensating for the cessation of transcription from mid-spermiogenesis onwards. Nonetheless, they also identified a small cluster of meiotically-induced mRNAs that were actively translated only in post-meiotic stages.

More recently, the development of genetically modified organisms that express an affinity-tagged ribosomal protein has provided a straightforward means of isolating ribosomes along with their bound mRNAs. Interestingly, the expression of these tagged proteins can be driven by a tissue/cell-specific promoter, such as the Cre-lox system in mice, which enables the capture of tagged ribosomes from an entire organ or tissue without the need to isolate the cells of interest (for review, see King & Gerber (2014)). These methodologies, initially called translating ribosome affinity purification (TRAP) in the mouse (Doyle et al. 2008, Heiman et al. 2008), have been used several times to capture the translatome of various testicular cell types. For instance, Sanz et al. (2013) took advantage of Cyp171Cre and Amh-Cre mice to investigate the translatome of adult Leydig cells and Sertoli cells respectively. They also used this approach to investigate the regulation of Leydig cells by LH and that of Sertoli cells by FSH and testosterone in gonadotropin-depleted mice. They notably found that the early response to LH (within 1 h) was characterized by the induction of several transcription factors and genes involved in cell cycle while the secondary response to LH (after 4 h) involved the up-regulation of genes involved in steroid metabolism and FGF signaling and the down-regulation of several transcription repressors. The regulatory role of testosterone was also successfully examined by De Gendt et al. (2014) who combined a TRAP approach that used Amh-Cre mice, a mouse model lacking a functional androgen receptor (AR) in Sertoli cells, and RNA-seq analysis. After determining the Sertoli cell translatomes of prepubertal and adult mice, which they found to be very similar, the authors compared these WT Sertoli cell translatomes to that of Sertoli cells lacking the AR and identified androgen-regulated genes at the onset of meiosis, which included many plasma membrane and cytoskeleton factors involved in cell junction and adhesion. Finally, another study took advantage of this method to investigate the translatome of neonatal testicular germ cells at the onset of meiosis (Evans et al. 2014). Using a synchronized spermatogenesis model, the authors identified the changes in ribosome-bound mRNAs taking place in both differentiating spermatogonial cells (with Ngn3-Cre and Stra8-Cre mice) and maturing Sertoli cells (with Amh-Cre mice) after retinoic acid restoration.

Note that these ribosomal profiling analyses not only help to evaluate translation efficiency and estimate corresponding protein abundance more accurately than classical transcriptomic approaches, but, when coupled with RNA-seq, they can also provide information about ribosome occupancy, translation initiation, elongation, and termination at near-nucleotide resolution (for review, see Ingolia (2014)).

From gene expression measurement to new testicular protein isoform prediction

As mentioned previously, the measurement of steady-state gene expression does not necessarily estimate the actual proteome well. This is true from both the quantitative and qualitative points of view and especially for microarray experiments. Specifically, because microarrays measure gene expression through the sequence-specific hybridization of RNAs to DNA probes, they cannot gather information about transcript structure outside the sequence targeted by the probes. Furthermore, because most probes recognize several transcript isoforms, they report average gene expression and fail to identify the specific isoforms actually expressed in a given sample.

In this regard, the recent advance of RNA-seq technologies, together with the development of associated analysis pipelines, has revolutionized the field of transcriptomics. RNA-seq is an efficient and cost-effective way to obtain large amounts of transcriptome data and identify both new genes and new isoforms, by the sequencing of novel exons and/or novel exonic junctions. RNA-seq thus makes it simultaneously possible to determine the structure of thousands of transcripts and to measure their abundance. Therefore, it provides a more accurate prediction of all corresponding protein isoforms (Fig. 1B). Several RNA-seq analyses have already been conducted to investigate spermatogenesis in rodents, with either isolated cells (Gan et al. 2013b, Soumillon et al. 2013, Chalmel et al. 2014b) or testes at various stages of the first wave of spermatogenesis (Laiho et al. 2013, Schmid et al. 2013, Margolin et al. 2014). All these studies have led to the reconstruction of a plethora of transcripts, including known isoforms but also thousands of new isoforms of known genes and hundreds of uncharacterized transcripts that correspond to either new coding or non-coding genes. The amount of information generated in such RNA-seq studies is so huge that a single study cannot undertake to report on all the exploration possibilities. For example, Chalmel et al. (2014b) restricted their analysis to novel unannotated loci and
used four different bioinformatics tools to distinguish between transcripts with high and low coding potential. Margolin et al. (2014) initiated a broad analysis of splicing events by identifying transcripts that contained novel splice junctions in which the open reading frame (ORF) was maintained. Finally, Schmid et al. (2013) focused their study on the splicing dynamics of mRNAs in male germ cells, with a special emphasis on the newly identified splicing events that might affect protein isoform production during mouse meiosis, and identified significantly enriched motifs for PTB, TRA2B and STAR proteins in and around meiotically-regulated cassette exons. Therefore, although all these RNA-seq studies have highlighted many potential new protein-coding transcripts expressed in male germ cells, a thorough analysis aimed specifically at identifying the variants that actually code for specific proteoforms with distinct biological functions is still needed. Most importantly, the functional relevance and validity of these findings still require experimental validation at the protein level.

Proteogenomic approaches applied to spermatogenesis

Compared with microarrays, which are intrinsically limited to studying the expression of genes for which probes are spotted on their surface, MS-based proteomics has long been considered to be more powerful in the sense that theoretically it can detect and quantify any protein entity within a given sample. Protein identification, however, typically involves the comparison of experimental masses obtained by mass spectrometry to that of in silico-digested protein databases. Therefore proteomic studies are also limited to the sequence content of the database that is used for identification purposes. Several methods, called proteogenomics, have emerged to overcome this limitation and help identify novel peptides not present in reference protein sequence databases (for review, see Hernandez et al. (2014)). These methods rely on the construction of customized protein sequence databases that include, for example, products resulting from the six-frame translation of the reference genome or the three-frame translation of transcripts, or both, regardless of whether these correspond to known mRNAs, non-coding RNAs, or pseudogenes. Large consortia seeking to decipher the complete human proteome through the analysis of several human tissues, including the testis (Kim et al. 2014, Wilhelm et al. 2014), have recently used such approaches.

Among the most promising proteogenomic methods is RNA-seq-based proteogenomics, also called ‘proteomics informed by transcriptomics’ (PIT). The PIT strategy derives the customized protein sequence database queried for protein identification directly from RNA-seq data of the same or a similar sample (Evans et al. 2012). It therefore limits protein products in the custom database to those resulting from the three- or six-frame translation of the assembled transcripts that are indeed expressed in the organ, tissue, or cell type of interest. Compared with other proteogenomic approaches, PIT offers the advantage of a smaller database, which in turn reduces the number of false positives and increases sensitivity (Fig. 1A).

Recently, this strategy was applied to the identification of new proteins expressed during late stages of rat spermatogenesis (Chocu et al. 2014). In this study, protein extracts from isolated rat pachytene spermatocytes and round spermatids were first trypsin-digested and analyzed by nano LC–MS/MS. Next, MS/MS spectra were queried against a customized database derived from a previous RNA-seq analysis of rat testicular cells (Chalmel et al. 2014b), which had identified almost 12 000 new transcript isoforms. It also reported the existence of more than 1400 completely new unannotated loci, most of them preferentially expressed in spermatocytes and/or spermatids. Because of this high gene discovery potential, Chocu et al. voluntarily restricted their PIT approach to meiotic and post-meiotic germ cells. This experiment led to the identification of 44 novel coding genes expressed during rat spermatogenesis, including 14 that were initially thought to correspond to non-coding RNAs. This approach has also been used to study the testicular proteome of the red abalone, Haliotis rufescens (Palmer et al. 2013) and allowed the identification of almost 1000 proteins. This number of proteins is especially remarkable when we consider that the number of UniProt entries still does not exceed 140 for this non-model species (Release 2014_11).

There is no doubt that the increasing performance of mass spectrometers and decreasing cost of RNA-seq will allow for rapid democratization of PIT studies and of proteogenomics in general. These approaches will be critical to the full characterization of both the transcriptome and the proteome of model organisms in various biological contexts, which in turn will help to annotate the corresponding genomes. PIT strategies are also a unique opportunity for non-model species, for which reference genome sequences are not available. Their transcriptomes and proteomes can be thoroughly examined without requiring the use of nucleic or protein sequence databases from phylogenetically distant species. Finally, regardless of the model of interest, the combination of RNA-seq and mass spectrometry into a PIT study offers a straightforward method of investigating the correlation of transcriptomes and proteomes, because protein profiles can be directly compared to transcript profiles on which protein identifications are also performed (for review, see Wang et al. 2014).

Conclusion

The recent progress in next-generation sequencing technologies, together with the improved performance of mass spectrometers, has made possible a fruitful revisit of the testis genomic landscape. While we are
now getting close to the complete identification of the molecular factors involved in spermatogenesis, an understanding of all the regulatory mechanisms that drive gene and protein expression during germ cell development and the identification of the key factors for male fertility both require additional work. This will notably imply the combination of all types of available data, i.e., from epigenomic, regulomic, transcriptomic, proteomic, and interactomic studies, in order to link the flow of information from DNA to functional proteins and non-coding RNAs. The success of this integrative work will also depend on the development of new types of web servers, such as the ReproGenomics Viewer (http://rgv.genouest.org; Darde et al. 2015), which allows the visualization, mining, and comparison of various types of omics data (e.g., ChIP-seq, RNA-seq, MS-based proteomics) in a multi- and cross-species manner. Finally, a current challenge in biology resides in the development of methods to investigate single cells at the genomic, transcriptomic, proteomic, and metabolomic level (for review, see Tsioris et al. 2014). The use of these so-called single-cell approaches will mandate the more detailed study of the kinetics of germ cell differentiation and, most importantly, enable us to gain insight into the biology of discrete cell populations within the testis, such as the spermatogonial stem cell.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the review.

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