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The landscape of RNA-binding proteins in mammalian spermatogenesis

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Despite continuous expansion of the RNA-binding protein (RBP) world, there is a lack of systematic understanding of RBPs in mammalian testis, which harbors one of the most complex tissue transcriptomes. We adapted RNA interactome capture to mouse male germ cells, building an RBP atlas characterized by multiple layers of dynamics along spermatogenesis. Trapping of RNA-crosslinked peptides showed that the glutamic acid-arginine (ER) patch, a residue-coevolved polyampholytic element present in coiled-coils, enhances RNA binding of its host RBPs. Deletion of this element in NONO (non-POU domain-containing octamer-binding protein) led to a defective mitosis-to-meiosis transition due to compromised NONO-RNA interactions. Whole-exome sequencing of over 1000 infertile men revealed a prominent role of RBPs in the human genetic architecture of male infertility and identified risk ER patch variants.

Mammalian spermatogenesis spans three prime phases: mitosis, meiosis, and spermiogenesis, during which germ cells differentiate in turn as spermatogonia, spermatocytes, and spermatids, ultimately producing spermatozoa. During this multi-step, dynamic process germ cells undergo drastic morphological alterations accompanied by a cascade of molecular events encompassing chromatin remodeling (1), transcriptional transition (2), and posttranscriptional regulation (3), that give rise to the remarkably complex testicular transcriptome (4, 5). Testis expresses an abundance of RNA-binding proteins (RBPs) (6), involved in processing, transporting, and modulating RNA, as well as coordinating transcription (7). Testis-specific RBPs orchestrate the storage and translational activation of spermiogenic mRNAs that are transcribed several days before they are needed due to the spermiogenic

nucleus undergoing compaction and transcriptional silencing (8, 9). Despite our growing knowledge of individual RBPs in spermatogenesis, a comprehensive atlas of authentic RBPs is still lacking.

RNA interactome capture (RIC) is a high-throughput RBP-profiling approach (10, 11) that has advanced our understanding of structural and functional features of RBPs through its application in various cell lines and somatic tissues (12–17). However, capturing the RBPome in specific types of primary cells derived from a tissue is technically challenging. In mammalian testis, germ-cell-specific RIC is valuable for mining RBPs and studying their spermatogenic dynamics.

RBDmap and related methodologies allow systems-level detection of RNA-crosslinked peptides (18, 19). These

methods verify well-defined canonical RNA-binding domains (RBDs) and reveal noncanonical RBDs and RNA-binding nondomain elements (short motifs not in annotated domain) especially from intrinsically disordered regions (IDRs). IDRs drive protein-protein and protein-RNA interactions independently or in cooperation with other globular RBDs (12, 20). While IDR elements are increasingly recognized to widely function in cellular processes and diseases, they are largely undefined and only a few studies have used animal models to pinpoint their roles (21, 22).

Human mutations in RBP-encoding genes are critical determinants for both mendelian and somatic pathogenesis (23). Intersecting the known RBP repertoire with public disease association data showed heterogeneity of RBP mutation properties among different diseases (23). Male infertility is a growing major reproductive problem worldwide, and its etiology remains unknown in about 40% of patients, with genetic factors being recognized as crucial contributors (24). By far, scaled genomic sequencing of male infertile cohort has been largely restricted to patients with Western ancestry (25, 26). Despite many efforts to decipher its monogenetic causes (27, 28), our understanding of the genetic architecture underlying male infertility is rather limited (29), lagging behind other diseases.

Here, we performed RIC-based proteome-wide mapping of endogenous RBPs in mouse male germ cells (mMGCs) and further delved into a systematic profiling of RBDs and RNA-binding nondomain elements. Our approach led to the discovery of a polyampholytic nondomain element, named glutamic acid-arginine (ER) patch, that is widely present in coiled-coil (CC) motifs of RBPs and promotes RNA-binding activity of its host RBPs and male germ cell development. Intersecting these datasets with our whole-exome sequencing profiles from a large cohort of infertile men created an opportunity to explore the genetics of male infertility with a focus on and full coverage of RBPs.

Results

mMGC RIC identifies highly specialized, germline-specific RBPs

To systematically identify RBPs in mMGCs, we used STA-PUT velocity sedimentation to purify sufficient numbers of spermatogonia (SG), pachytene spermatocytes (pSC), and round spermatids (rST), which were subjected to a modified RIC procedure separately (Fig. 1A and fig. S1A). Several known RBPs were validated by Western blot assays, confirming they were captured only when UV (254 nm)-crosslinked in RIC (fig. S1B). Liquid chromatograph-mass spectrometry (LC-MS/MS) identified highly reproducible amounts of proteins between two biological replicates of RIC (fig. S1C). Under a stringent enrichment criterion of crosslink (CL)/noncrosslink (noCL) ≥ 10 for iBAQ intensity in mass spectrometry (fig.

S1C), we defined 918 proteins from SG, 955 from pSC, and 1048 from rST (Fig. 1B and table S1), resulting in a total of 1744 nonredundant proteins referred to as “mMGC RBPome/RBPs”, of which 388 proteins are present in all three cell types, and 299, 275, and 381 are specific to SG, pSC, and rST, respectively (Fig. 1B). As expected, GO analysis of identified proteins displays their predominant involvement in RNA-related cellular processes and molecular functions (fig. S1D), and mMGC RBPome exhibits physicochemical properties similar to other somatic RBPomes (fig. S1E) (10), supporting the validity of our approach.

Taking into account only orthologous genes across species, 302 genes are common to our mMGC RBPome and other published RBPomes [data collected by (30)], whereas hundreds of proteins are dataset-specific (Fig. 1C), including 345 genes particular to our mMGC RBPome. At the protein level, comparative analysis of mMGC RBPome and combined mouse somatic (mSC) RBPomes (30) revealed distinct processes enriched in mMGC RBPome or mSC RBPomes (Fig. 1D). Excluding proteins with annotated RBDs (10, 30) and those common to both mMGC RBPome and mSC RBPomes defined a group of 515 “mMGC novel” candidate RBPs, specifically enriched in spermatogenesis-associated processes such as reproduction, meiosis, and fertilization (Fig. 1, D and E, and table S1), and phenotypically related to male infertility and abnormal spermatogenesis (fig. S1F). A large proportion ($n = 397$) of these candidate RBPs lack RNA relevance (Fig. 1F, fig. S1G, and table S1). Relative to a minority (18.7%) of SG-specific RBPs, more (31.6%) of pSC-specific and even half (49.9%) rST-specific RBPs were novel and not previously related to RNA (Fig. 1, F and G, and table S1).

To confirm that the identified candidate RBPs are bona fide factors that act on RNA, we applied the Protein-RNA Complex Capture (2C) technique (31, 32) using single cell suspension of whole adult testis and 293T cells (fig. S2A). UV crosslinking to capture ribonucleoprotein (RNP) in 2C was manifested effective by silver staining (fig. S2A) and 16 mMGC RBPs were verified by Western blot (fig. S2, B to E). PDHA2 and TEX30, two 2C-validated novel RBPs, were orthogonally verified by CLIP-autoradiography (figs. S3A and S4A), eCLIP-seq (fig. S3, B to D; fig. S4, B to D; and table S2), and EMSA (figs. S3E and S4E), and were both testis-specific and required for spermatogenesis (fig. S3, F to K, and fig. S4, F to L), detailed in supplementary text. Collectively, our results suggest that RIC faithfully captures testis-specific RBPs, which have likely evolved specialized features adapted to the complex meiotic and post-meiotic mechanisms of RNA regulation.

Temporal expression and RNA-binding dynamics of mMGC RBPs

To delineate expression patterns of RBPs during

spermatogenesis, we leveraged our previously published datasets of tandem mass tag (TMT) based quantitative proteomes of whole cell extract of four types of mMGCs (SG, pSC, rST, and elongated spermatids (eST)) (33) (fig. S5A), and clustered all proteins based on their temporal expression correlation (Fig. 2A and table S3). Classical and nonclassical RBPs constitute the majority (80-94%) of RBPs that are primarily expressed in spermatogonia (Clusters 1-5), while factors that first appear in meiotic or post-meiotic stages are enriched (41-59%) for novel RBPs (Clusters 6-10) (Fig. 2B). Functional analysis revealed distinct roles of RBPs across clusters (Fig. 2C), in concert with the distinctive nature of the cellular processes in the four mMGC types. Protein-protein interaction (PPI) modules within several clusters (Clusters 2, 6, and 9) hint that multiple RBPs may jointly engage in certain pathways (fig. S5B and supplementary text).

Next, we tracked mRNA-protein concordance/discordance along spermatogenesis through integration of the TMT-MS data with RNA-seq data (34) (fig. S5A and table S4). Novel RBPs showed lower concordance than classical and nonclassical RBPs, and lower than DNA-binding proteins (DBPs) (fig. S5C). Moreover, rST-specific RBPs showed the lowest concordance among all subsets of mMGC RBPs and nonRBPs (fig. S5C). We then grouped genes with distinct mRNA-protein discordance patterns across four stages (SG, pSC, rST, and eST) into four clusters (Clusters A to D) (fig. S5D). Cluster B was enriched for known translation-delayed genes and chromatoid body genes (35, 36) (fig. S5E and table S4). From the proportions of RBPs relative to nonRBPs, the Cluster-B-type mRNA-protein discordance of RBPs is likely a consequence of widespread translational delay regulation (fig. S5F).

A recent study assessed relative RNA-binding activity of RBPs by dividing RNA association degree (RIC) by protein abundance of whole cell extract (WCE), and demonstrated that proteins in direct contact with RNA have a high RIC/WCE ratio, which was leveraged to model concordant behavior of RBPs as ribonucleoprotein subunits (37). Following this lead, we calculated relative RNA-binding activity of 1218 RBPs in spermatogenesis (Fig. 2D and table S5). Classical RBPs showed consistently strong RNA binding, while novel RBPs exhibited overall lower but more volatile RNA binding - inactive in SG but relatively active in pSC and rST (Fig. 2D). The latter could be explained by increased access to target RNAs (38) for novel RBPs, as transcriptome evolves extensively with developmental transitions and these RBPs primarily localize at germ-cell-specialized entities where RNAs might be selectively associated and compartmentalized, especially those in rST (Fig. 2E and fig. S6, A and B).

We then classified our mMGC RBPs of different stages into three groups (Fig. 2F, table S5, and supplementary text): Strong ($\text{Log2(RIC/WCE)} \geq 2$, $\text{Log2(WCE)} > 0$), Weak

($\text{Log2(RIC/WCE)} < -2$), and Moderate ($\text{Log2(RIC/WCE)} < 2$ and ≥ -2). To test if relative RNA-binding activity analysis could reflect RNP assembly, MIWI-ELAVL1 associated RBPs (9) were examined, and these proteins exhibited a convergent rise of relative RNA-binding activity across stages (pink module; fig. S6, C to E), a phenomenon compatible with the MIWI-ELAVL1 complex formation in rST (9). As expected, housekeeping ribosome proteins stably act as Strong RBPs across three stages (green module; fig. S6, F to H). We showed three additional modules of RNP assembly (Supplementary text): ribosome biogenesis-associated proteins (yellow module; Fig. 2, F and G, and fig. S6I), mainly composed of nonclassical RBPs, emerged as a Strong module specifically in SG; conversely, some mRNA splicing and transport factors (red module; Fig. 2, F and G, and fig. S6J), mainly classical RBPs, were more active in pSC and rST than in SG; cilium movement and acrosome assembly proteins, mainly novel RBPs, exhibited a marked increase of relative RNA-binding activity from SG to pSC and rST (blue module; Fig. 2, F and G, and fig. S6K). These examples suggest stage specificity of the RNP network in spermatogenesis.

RBDmap identifies testis-enriched RNA-binding domains

According to annotations of RBDs from InterPro database, we found that canonical RBDs, such as RRM and KH, were overrepresented in mMGC-mSC common classical RBPs (fig. S7A and table S6). WD40 and Tudor domains frequently emerged in mMGC-mSC common nonclassical RBPs (fig. S7A). Domains enriched in mMGC novel RBPs generally lacked evidence for RNA binding, such as PKINASE and ANK2 domains (fig. S7A and supplementary text). These two domains were confirmed for their RNA-binding capacity by 2C assay (fig. S7B). We also observed distinct sequence patterns, as RS, polyQ, and polyG (39) were enriched in IDRs of classical and nonclassical RBPs, while the GPP motif was specifically presented in IDRs of mMGC novel RBPs (fig. S7C).

To experimentally profile RBDs, we conducted RBDmap (18, 19) using adult and postnatal day 28 mouse testes separately. LC-MS/MS identified total 721 non-redundant peptides (hereafter named RBDpeps) of RNA-bound regions of 393 RBPs under a Moderate criterion (Fig. 3A, fig. S8A, table S7, and supplementary text). 511 RBDpeps overlap with documented domains (domain RBDpeps) and 210 RBDpeps do not (nondomain RBDpeps) (Fig. 3B). The specificity of these RBDpeps was proven by their considerable overlapping with RRM domains of classical proteins such as DAZL (fig. S8, B and C, and supplementary text). As expected, both domain and nondomain RBDpeps were enriched with glycine (G) and arginine (R) amino acids (fig. S8D), which are known hotspots of RNA interaction (20). Our RBDmap identified 89 RBPs shared with mouse HL-1 cells, in which 64 RBPs

harbored overlapping RBDpeps (fig. S8E, left). Moreover, 307 and 155 domain RBDpeps (90% in total) mapped to classical and nonclassical RBPs, respectively (fig. S8E, right). Examples of these 462 RBDpeps included classical RBPs (YTHDF1 and PUM1) and nonclassical RBPs of the Tudor family (fig. S8, F and G; fig. S9, A and B; and supplementary text). We also identified 49 nonredundant domain RBDpeps mapped to 37 novel RBPs, of which 15 are testis-specific proteins (fig. S8E, right, and table S7). 5 core histones and 2 linker histones were identified in both our mMGC RBPome (table S1) and testis RBDmap (table S7). Representative examples are testis-specific histone variants, such as HIST1H1T and H2AFB1 (Fig. 3, C and D, and fig. S10A). Additionally, 6 RBDpeps were found in the protamine-P2 domain of PRM2, and 2 in the TP2 domain of TNP2 (Fig. 3D). Additional instances of RBDpeps overlaying domains of novel RBPs were illustrated (fig. S10, B and C, and supplementary text). Four of these testis-specific RBDs were 2C-validated (fig. S10D), confirming their bona fide RNA-binding capacity.

RNA-binding elements identified in nondomain areas

Based on their physicochemical properties, motifs within the 210 nondomain RBDpeps (Fig. 3B) could be clustered into three major archetypes (40): polar tracts, polyelectrolytes, and polyampholytes (Fig. 3E and supplementary text). One polyampholytic RNA-binding element identified herein is the ER patch that features a mixed-charged sequence composed of E and R amino acids (Fig. 3E and table S8). Furthermore, nondomain RBDpeps are much more enriched in alpha helices forming coiled-coils (CCs) ($n = 23$; all CCs are annotated based on Pfam database here and after) than in disordered regions and low complexity regions (Fisher's exact test; CCs: odds ratio (OR) = 9.62, $P = 1.1 \times 10^{-4}$; disordered regions: OR = 2.43, $P = 1.1 \times 10^{-3}$; low complexity regions: OR = 2.46, $P = 9.8 \times 10^{-4}$) (fig. S11A). All the ER hits ($n = 4$) are evolutionarily conserved and located in CCs of NONO (non-POU domain-containing octamer-binding protein), SFPQ (splicing factor proline- and glutamine-rich), and SAFB (scaffold attachment factor B) (Pfam database), as well as PSPC1 (paraspeckle component 1) [suggested by (41)] (Fig. 3F). An additional conserved ER patch located in the CC of RBP (RBP-CC) was also identified in two somatic RBDmap datasets (18, 42) (Fig. 3F). Since these five ER-containing CCs overlapped with RBDpeps, we speculated that they themselves possess RNA-binding capacity. We tested two of them by 2C assay, and found that both indeed bound RNA when overexpressed in 293T cells (fig. S11B). We then used EMSA to further examine NONO CC, and found it can bind RNA substrates with various sequences, preferentially polyA and G-rich RNAs (fig. S11C). Collectively, RBDmap identified these RNA-binding CCs harboring ER patch elements.

The ER patch is a frequent element coevolving in local in RBP-CCs

We explored the prevalence of the ER patch at a genome-wide level, especially those in CCs, as CCs were more widespread in mMGC RBPs than in nonRBPs and DBPs (Fig. 3G). Consensus sequence analysis further revealed a specific enrichment of the ER patch in mMGC RBP-CCs but not in nonRBP-CCs (fig. S12A). We extended the pool of ER patch by searching for any 10-mer peptide containing at least 4 Glu and 4 Arg (4E4R) or 3 Glu and 3 Arg (3E3R) throughout mMGC RBPome, DBPome, and nonRBPome (table S9). Both CC-3E3R and CC-4E4R were more frequently present in RBPs than in DBPs and nonRBPs (fig. S12B). The in silico identified ER patches (4E4R) were classified into three groups (fig. S12B and table S9): The first group has 21 ER patches in nonCCs of 19 RBPs; the second group has 92 ER patches in CCs of 74 nonRBPs; the third group has 51 ER patches localized in CCs of 34 RBPs, in which 4 of these ER-containing CCs were capable of binding RNA (fig. S11B). Notably, some RBPs restricted to somatic RBPomes also contain 4E4R patches (table S9), suggesting a potentially wider spread of the ER patch in RBPomes.

To infer potential structural and functional interaction between ER patches and other elements of the same proteins, we conducted sequence covariation analysis commonly used to predict inter-residue contacts/distance or functional relevance (43–45). We found no long distance contacts. Instead, inter-residue interdependence were confined within ER patches (Fig. 3H). Amino acids in the RBP-CC-contained ER patch (RBP-CC-ER) were predicted by EVcouplings (46) to be evolutionarily more co-variated than the other two groups (T-test; RBP-CC-4E4R vs RBP-nonCC-4E4R: $P = 0.046$; RBP-CC-4E4R vs nonRBP-CC-4E4R: $P = 0.027$) (Fig. 3, H and I, and fig. S12C), suggesting that a mutual interdependent relation (45) of E and R pairing within the RBP-CC-ER is critical for maintaining function. For example, residues of LUC7L3 were intensively co-variated within amino acids 238–297 where ER patches are located (Fig. 3H). Similar local evolutionary covariation was also observed in the ER patches of INCENP, UPF3B, and NONO (Fig. 3H). These results hint the functional relevance of the ER patch from a co-evolutionary perspective.

The ER patch can enhance RNA binding of its host RBP

Next, we asked whether the ER patch has a role for its host protein in RNA binding. To address this, we deleted the sequence EEERRRRE in NONO (NONO^{ΔER}) and conducted eCLIP-seq to compare RNA targets of NONO and NONO^{ΔER} proteins in 3T3-L1 cells. NONO^{ΔER} showed a decrease in the number of RNA-binding peaks and genes and also a slight change in target RNA profiles (Fig. 4, A and B, and fig. S13, A

to C). We also observed a global decrease in binding intensity for ~75% of target genes (Fig. 4C). To directly visualize radio-labeled protein-bound RNA levels, we used conventional CLIP and found that RNA-binding capacity of NONO^{ΔER} was much weaker (Fig. 4D). This result was reinforced by in cellulo quantitative-2C (q-2C) assays comparing overexpressed NONO and NONO^{ΔER} and in vitro q-2C comparing purified recombinant NONO (rNONO) and NONO^{ΔER} (rNONO^{ΔER}) proteins as well as mRNP capture assay in 293T cells (Fig. 4, E and F, and fig. S13, D and E). Similar impairment of RNA-binding capacity was also found by conventional CLIP for PSPC1^{ΔER} (fig. S13F), indicating that the ER patch enhances RNA binding of these two RBPs.

To further assess whether E or R amino acid contributes to enhanced RNA binding, we produced a panel of NONO-CC-ER mutants with either substitution or deletion of certain amino acids on the ER patch (Fig. 4E). NONO^{AA}, NONO^{EA}, and NONO^{RA} (substitution of all E and/or all R by Alanine (A)) all exhibited impaired RNA binding (Fig. 4F and fig. S13D). Nonetheless, NONO^{RE} (replacement of EEERRRRE by ERERERER) did not alter RNA-binding capacity of NONO (Fig. 4F and fig. S13D). These data demonstrate that both E and R amino acids in the ER patch are functionally indispensable.

Phase separation drives intermolecular interaction-mediated NONO recruitment for granule assembly (47), for which we reasoned that the diminished RNA binding of the NONO mutants might be coupled with a negative effect on their phase-separated condensate properties. We therefore examined these GFP-fused NONO proteins in 293T cells. In cells expressing high levels of GFP-NONO, ER deletion or substitution induced a conversion of NONO from a droplet state to a fiber-like state (Fig. 4, G and H). In cells expressing GFP-NONO at low levels as endogenous NONO, the foci of the mutant proteins remained in a droplet state and these foci were examined for fluorescence recovery after photobleaching (FRAP) (Fig. 4I). We found that the ER-mutated NONO except NONO^{RE} exhibited a concordant reduction of the mobility of NONO foci (Fig. 4, I to K). Nonetheless, the protein interactome of NONO versus NONO^{ΔER} in 293T cells and in mouse testes (see next paragraph) showed that ER patch deletion in NONO did not affect its major partners SFPQ and PSPC1 (fig. S14, A to D, and table S10) nor its co-localization with PSPC1 (fig. S14E).

The ER patch in RBP-CCs is functional during spermatogenesis

Nono and *Safb* are the two RBDmap-identified RBP genes that not only harbor ER patch sequences but also have knockout-based evidence demonstrating their physiological requirement in the testis. *Nono* knockout mice exhibited a decrease in body weight, testis weight, and sperm counts (48).

Safb knockout mice exhibited male infertility with postnatal germ cell deficiency (49). To further assess the biological relevance of the ER patch, we created two knock-in mouse mutants with the ER patch precisely deleted from CCs of *Nono* and *Safb* genes (Fig. 5A and fig. S15A). ER patch-truncated RNA and protein in both *Nono*^{ΔER/Y} and *Safb*^{ΔER/ΔER} mice were expressed at similar levels compared with wild-type littermates (Fig. 5B and fig. S15, B and C). *Nono*^{ΔER/Y} mice exhibited reduced body weights, sperm counts, and sperm motility, as well as increased numbers of apoptotic germ cells (Fig. 5C). Overall, in several measurements focusing on the testis, *Nono*^{ΔER/Y} mice exhibited phenotypic defects resembling, albeit quantitatively less severe than, *Nono* knockout (48) (Supplementary text). *Safb* knock-in mice also exhibited defects in germ cell development, with a milder phenotype than the *Safb* knockout (49). Whereas *Safb*^{ΔER/ΔER} mice exhibited normal testis weight, sperm count, and sperm motility, there was an increase in the number of apoptotic germ cells and abnormal elongated spermatids in the testis (fig. S15, D and E). *SAFB*^{ΔER} showed a trend for reduced RNA-binding capacity in 293T cells, which, however, did not reach statistical significance (fig. S15F), possibly because *SAFB* harbors additional ER residues flanking the deleted ER patch (Supplementary text). These results indicate that the ER patch should be indispensable for its host RBPs to fully function in spermatogenesis.

Co-immunostaining NONO with SOX9 (Sertoli cell marker) or with LIN28A (SG marker (50)) showed that NONO protein is predominantly expressed in Sertoli cells and spermatogonia, and its signal was detectable in spermatocytes (Fig. 5D and fig. S16). *Nono*^{ΔER/Y} mice displayed increased numbers of Sertoli cells (differentiating to mitotic arrest in adulthood) and spermatogonia but decreased spermatocytes and spermatids (Fig. 5E and fig. S17, A to C). The partial loss of leptotene spermatocytes was observed at stage IX-X seminiferous tubules in *Nono*^{ΔER/Y} mice (fig. S17D). Spermatocyte chromosome spread analysis showed a normal progression of meiotic prophase after leptotene stage (fig. S17E). These results suggest an abnormal expansion of Sertoli cell and spermatogonia populations and a subsequent defect in cell transition into meiotic leptotene spermatocytes.

The ER patch enhances endogenous NONO-RNA interactions and regulates mitotic transcripts ensuring germ cell transition into meiosis

Repressed RNA-binding activity of NONO protein was also found in *Nono*^{ΔER/Y} testes (Fig. 5F). We then conducted eCLIP-seq to assess transcriptome-wide alteration of RNA binding profiles of NONO^{ΔER} proteins in the testis. While NONO^{ΔER} shared most of the bound targets with wild-type NONO in the testis (Fig. 5G and fig. S18, A to C), its binding capacity was compromised in that ~70% of bound targets showed

decreased binding intensity in NONO^{ΔER} testes (Fig. 5H and table S11), consistent with the aforementioned findings in cell lines (Fig. 4, A to C), thereby corroborating the role of the NONO ER patch as an RNA-binding enhancer.

To further investigate whether the diminished RNA-binding capacity of NONO^{ΔER} alters cellular gene expression across the spermatogenic lineage, we performed single-cell RNA sequencing (scRNA-seq) on testicular cells of adult *Nono*^{+Y} and *Nono*^{ΔER/Y} mice. Cell types identified in our study corresponded well with previous testicular scRNA-seq datasets (51, 52) (Fig. 5I and fig. S18, D and E). The higher ratio of mitotic germ cells (type A spermatogonia, type B spermatogonia, and preleptotene spermatocyte) to meiotic ones (leptotene, zygotene, pachytene, and diplotene spermatocyte) in *Nono*^{ΔER/Y} versus *Nono*^{+Y} single testicular cells (fig. S18D) is in keeping with our cytological data (Fig. 5E and fig. S17, B to D), confirming the defect at the mitosis-to-meiotic transition in *Nono*^{ΔER/Y} mice.

To ascertain whether and how NONO binds and regulates RNAs that are responsible for this transition (53), we intersected scRNA-seq datasets with eCLIP-seq profiles to dissect RBP-mediated cell-state-specific RNA regulation (54). NONO CLIP targets are predominantly expressed in mitotic germ cells and Sertoli cells (Fig. 5J, top), the two cell types where NONO is mainly expressed (Fig. 5D and fig. S16), supporting the specificity of our CLIP data. Furthermore, most CLIP targets are up-regulated specifically in these two cell types in *Nono*^{ΔER/Y} mice (Fig. 5J, bottom, and table S11). These results suggest the cell-type-specific target RNA binding and regulation mediated by the NONO ER patch in the testis.

Given that NONO tends to bind and up-regulate RNAs of transcription factors at mitotic phase in *Nono*^{ΔER/Y} germ cells (Fig. 5J and fig. S18A), we assessed the transcriptome alterations between *Nono*^{ΔER/Y} and *Nono*^{+Y} globally. We defined two gene sets with their expression levels enriched separately in the mitotic and meiotic phases based on our wild-type scRNA-seq data (table S12). NONO bound 40.9% (247/604) mitosis-enriched genes and 2.1% (4/191) meiosis-enriched genes (fig. S19A), indicating that NONO regulates mitosis-enriched genes more directly. *Nono*^{ΔER/Y} mitotic germ cells, but not meiotic germ cells, showed overall up-regulation of mitosis-enriched genes and overall down-regulation of meiosis-enriched genes compared to *Nono*^{+Y} (Fig. 5K and table S12). Among 6 representative mitosis-enriched genes illustrated by trajectory analysis were 5 NONO direct targets, and deletion of the NONO ER patch increased their expression (fig. S19, B and C, and supplementary text). Immunofluorescence analysis further revealed the protein levels of two NONO targets were increased (fig. S19D). These results point to a role of the NONO ER patch in programming the mitosis-to-meiosis transition at the RNA level.

Homozygous RBP variants prioritized for human infertility

We performed a large whole-exome sequencing study of male infertility, including samples from 516 infertile men with idiopathic non-obstructive azoospermia (NOA) and 530 infertile men with oligozoospermia that passed our quality control procedures (table S13). Additional 300 fertile men with normal childbearing history were enrolled as controls (table S13). Population genomics analysis confirmed Asian origin of our cohort and no notable difference in population structure between patients with male infertility and controls (fig. S20). We first screened for loss of function (LOF) and damaging nonsynonymous (DNS) variants, and then applied a bioinformatic pipeline to identify potential pathogenic genes and prioritize the genetic components (types of genes, variants, and elements) more relevant to the pathogenesis of infertility (fig. S21). Both homozygous LOF (Fisher's exact test; OR = 2.52, $P = 1.2 \times 10^{-4}$) and DNS (Fisher's exact test; OR = 1.24, $P = 3 \times 10^{-5}$) variants were significantly enriched in patients (Fig. 6A), indicating that homozygous variants are more accountable for the pathogenesis of infertility.

Further, we observed significant overrepresentation of homozygous DNS in RBPs in patients (Fisher's exact test; OR = 1.62, $P = 0.0484$), suggesting a prominent role of RBPs in the genetic architecture of male infertility (Fig. 6A), though there was no significant difference in the odds of having homozygous LOF in RBPs between patients and controls (Fisher's exact test; OR = 2.24, $P = 0.3590$) (Fig. 6A). Using a stringent criterion (fig. S21), we compiled a list of candidate RBPs with homozygous variants exclusively in patients, leading to 22 LOF variants (candidate LOF) in 18 genes and 137 DNS variants (candidate DNS) in 105 genes, with their expression specificities, categorical and functional annotations, and patient origin in terms of infertility subtypes plotted (Fig. 6B, table S14, and supplementary text). 156 out of 159 total variants (~98%) were confirmed by Sanger sequencing (table S14). This genetic landscape of RBP variants associated with male infertility led to four observations (Fig. 6B): (1) Homozygous LOF variants were frequent in novel RBPs; (2) Homozygous DNS variants were overrepresented in nonclassical RBPs that are mostly ubiquitously expressed; (3) Variants were enriched in genes involved in four pathways (supplementary text), which were also overrepresented in mMGC RBPome relative to mSC RBPome (Fig. 1D); (4) Variants were in an array of genes with known (above moderate-level evidence) and unknown (limited or no evidence) pathogenicity for human infertility, based on the online IMIGC database (<http://www.imigc.org/data-sharing/>).

RBPs with heterozygous damaging variants tend to be mutation-intolerant

Sporadic studies have revealed pathogenic effects of

individual heterozygous RBP variants in human infertility (55). At the genome-wide scale, we next examined if heterozygous variants could contribute to the pathogenesis by prioritizing human disease genes on the basis of fitness consequences of disrupting variants, notably pLI (the probability of being LOF-intolerant) and Z-scores (the probability of being missense-intolerant) (56). Candidate genes (RBPs and nonRBPs) with patient-exclusive heterozygous LOF and DNS variants are more prone to be mutation-intolerant than genes with heterozygous LOF and DNS variants in controls (Wilcoxon test; $P = 2.21 \times 10^{-8}$ for RBPs with LOF; $P = 8.95 \times 10^{-19}$ for nonRBPs with LOF; $P = 3.27 \times 10^{-5}$ for RBPs with DNS; $P = 2.03 \times 10^{-8}$ for nonRBPs with DNS) (fig. S22, A and B). Moreover, candidate RBPs are more prone to be mutation-intolerant than candidate nonRBPs (Wilcoxon test; $P = 5.53 \times 10^{-6}$ for LOF; $P = 7.75 \times 10^{-8}$ for DNS) (fig. S22, A and B). These observations suggest that heterozygous damaging variants in RBPs are more likely to have a pathogenic impact for human infertility. In addition, candidate RBPs with heterozygous LOF or with heterozygous DNS are enriched for RNA splicing, translation, and cytoskeleton associated functions (fig. S22, C and D, and table S15), in concert with mMGC RBPome (Fig. 1D) and homozygous candidate gene analysis (Fig. 6B), reflecting a convergence of the biological and etiological roles of RBPs underlying infertility.

Homozygous DNS variants identified in RBP-CCs and RBP-CC-ERs

Widespread DNS variants in nondomain areas including IDRs were shown in our (Fig. 6B and supplementary text) and others' (23) work, though they were not significantly overrepresented in patients (Fisher's exact test; OR = 0.96, $P = 0.3767$) (Fig. 6A). Nevertheless, we found a significant enrichment of DNS variants within RBP-CCs in patients (Fisher's exact test; OR = 1.24, $P = 0.0313$) (Fig. 6A). We did not detect significant enrichment of DNS variants in ER patches (3E3R) among patients (Fisher's exact test; OR = 1.45, $P = 0.1387$) (Fig. 6A), perhaps due to the sparsity of ER patches genome-wide. All but one homozygous DNS variants in ER patches were found in infertile men (Fig. 6B, table S14, and supplementary text). All these patient-specific variants were located in the second CC-contained ER (3E3R) patch of UPF3B, including one highly recurrent homozygous variant -c.G1073A:p.R358H in eight patients and one non-recurrent homozygous variants -c.C1135T:p.R379C in one patient (fig. S23, A and B, and table S14). UPF3B is known as a nonsense-mediated decay (NMD)-related and ubiquitously expressed nonclassical RBP (57). No report directly links UPF3B with male infertility, yet several clues hint its functional role in spermatogenesis (58). The ER patch in the first CC of UPF3B is essential for its RNA affinity (59) and in its second was evolutionarily co-varied (Fig. 3H). The recurrent homozygous

variant in its second CC was validated for its impact on RNA binding (fig. S23C).

Discussion

We adapted RIC to three typical types of mMGCs (mitotic, meiotic, and post-meiotic), leading to the establishment of an atlas of mMGC RBPs, including proteins with previously unknown RNA-related activities that are specialized for meiotic and post-meiotic functions. This implies that meiotic and post-meiotic germ cells may provide an environment for proteins that harbor noncanonical or unknown RBDs to moonlight as RBPs, a trove for future studies. Some DBPs and chromatin-associated factors were identified as novel RBPs by our mMGC RIC. The concept of DNA/RNA binding proteins (DRBPs) has raised much interest, as DRBPs could coordinate gene expression at multiple levels more efficiently (60, 61). The highly ordered and dynamic chromatin-associated events during spermatogenesis may require DBPs or RBPs to evolve a dual role as DRBPs (62). We note that the possibility cannot be excluded that residual DNA and its crosslinked DBPs could be captured via RNA-DNA hybrids during the polyA pulldown even with Dnase treatment (61). Some structural proteins are present in the mMGC interactome; they could be components or structural constituents of large RNP complexes and their interactions with RNAs could be indirect. In addition to mMGC RBPs that bind polyA RNAs, those exclusively binding non-polyA RNAs (such as many noncoding RNAs) could be missed by RIC (63). We then showed that mMGC RBPs were involved in multi-layered dynamics covering proteomic expression patterns, proteo-transcriptomic discordancy, relative RNA-binding activity, and RNP concordant behavior. Our results highlighted developmental transition of expression and RNA-binding activity of RBPs, and illustrated RNP modules with distinct dynamic patterns of concordant behavior in spermatogenesis. With future availability of the global germline PPI network, we prospect to create a comprehensive atlas of RNP dynamics during spermatogenesis. Combined, our study presents the initial effort to systematically understand RBPs during mammalian spermatogenesis.

We used RBDmap to screen for testicular RBDs and RNA-binding nondomain elements to look into RNA-binding mechanisms, and discovered that the ER patch is a polyampholytic element present in RBP-CC and contains coevolutionarily coupled residue pairs. A recent perspective showed that the CC abounds in human RBPome, implicating its potential role in RNA regulation (64). The positional coupling of ER patches and CCs could be explained by the canonical alpha helices formation with polyampholytic sequences (40). The ER patches we identified are relatively small in length (18 amino acids on average), the negatively and positively charged residues are alternating and not in long blocks of the

same charge, and they prefer to reside in alpha helices. These features differ from what others have found about mixed charged stretches, which are relatively long (50 residues or more) and often contain blocks of negatively and positively charged residues within IDRs resulting in high local charge density for phase separation (65, 66). The well-mixed linear sequence of oppositely charged residues of the ER patch may adopt random-coil conformations, which promotes the flexibility and the high local concentration of connected functional elements (40, 67). Although a few polyampholytes are recently emerging to play a role other than regulating RNA such as condensates assembly (66), little is known about their role as an RNA-binding element. The high positional concordance of ER patches and CCs suggests that they are functionally coupled to regulate RNA.

The ER patches of three (out of four) tested RBPs (NONO, PSPC1, and UPF3B; ER mutation of UPF3B was found in infertile men) were verified biochemically to enhance RNA binding. By creating knock-in animal models, the ER patches of both tested RBPs (NONO and SAFB) were verified as a requirement for securing normal spermatogenesis. With a focus on NONO, we further found that its ER patch regulates the mitosis-to-meiosis transition of male germ cells, which can be explained by an impairment of RNA-binding activity of NONO and a concomitant up-regulation of NONO target mRNA transcripts. Previous studies show that NONO predominantly binds to the 5' ends of pre-mRNAs, and NONO knockdown induces a processing defect at the 5' end of NONO-bound pre-mRNAs (68). The depth of our scRNA-seq is not sufficient for analysis of sub-transcripts. Thus, we cannot exclude the possibility that some NONO targets are actually decreased in *Nono*^{ΔER/Y} testes due to the potentially altered pre-mRNA 5' end processing. On the whole, the NONO ER patch enhances NONO protein-RNA interactions, maintains target mRNAs expression, and ensures mitosis transition into meiosis. Loss of the NONO ER patch results in decreased NONO binding to its target mRNAs, up-regulated levels of mitotic transcripts, abnormal accumulations of Sertoli cells and mitotic spermatogonia, and impaired progression of mitotic germ cells into meiotic spermatocytes.

NONO seems to play a critical role under stress (69). External stimuli such as UV irradiation up-regulate NONO expression in the testis (48). Feeding increases the number of NONO-containing speckles in the liver (70). NONO knockout mice often exhibit a more severe phenotype under specific stimuli (48, 70). In our study, ER deletion caused severe functional consequence, with a droplet-like state being transformed into a fiber-like state, in high-level NONO expression in cell lines. We therefore speculate that the NONO ER patch may augment its role in response to stress-induced higher expression of NONO.

There are some noteworthy points about the RBPs

harboring CC-ER. First, CCs are especially enriched in RBPs and ER patches are especially enriched in RBP-CCs, implying that the ER patch may play a role in CC-mediated RBP-RNA interactions, rather than mediating CC formation or oligomerization, which is a general function of CCs in other kinds of nonRBPs (71). Second, RBPs containing CC-ER always form homo- or hetero-dimers, which are critical for the assembly of RNP granules. For example, DBHS family proteins SFPQ, NONO, and PSPC1 share structural similarities, with a common ER-patch-embedded CC, and their dimers could form via CC interactions (41, 72, 73). However, the ER patch seems dispensable for dimer formation as it can be absent from the available structures of these dimers (41). This notion is supported by the stable intracellular interaction of NONO^{ΔER} with SFPQ and PSPC1. Third, mutations in a CC of SFPQ affect SFPQ localization and polymerization and reduce SFPQ-nucleic acid interactions, and specifically, the C-terminal part of the CC that contains the ER patch mediates higher-order homotypic associations (73). Based on the above and our results, it is probable that such higher-order associations are linked to enhancing RNA binding.

Linkages between a wide spectrum of RBPs and specific diseases have not yet been established. Genetic etiology for human idiopathic infertility remains far from clear. Homozygous mutations are generally believed to be more relevant to infertility, and hence the intense focus of current research. Our whole-exome sequencing analysis strengthened this argument by uncovering enrichment of homozygous LOF and DNS variants in infertile patients, and prioritized these mutations in RBPs for future studies. Note that most homozygous LOF variants were found in novel RBPs, implying classical and nonclassical RBPs that have crucial roles in maintaining human development homeostasis are generally intolerant to mutations of large effects (74). However, the low frequencies of homozygous variants are not sufficient to explain a considerable proportion of idiopathic infertility patients. As a general scenario, idiopathic infertility could be of polygenic origin and caused by the cooperative effects of deleterious heterozygous variants (24). Our genome-wide heterozygous analysis supports a pathological role of heterozygous variants of both RBPs and nonRBPs in human infertility, with RBPs being more likely to be mutation-intolerant than nonRBPs, at both the LOF and DNS levels. It suggests that heterozygous variants constitute an unignorable genetic component to the pathogenesis of infertility, in concert with a recent finding demonstrating that single heterozygous DNS variant in *HIWI* is sufficient to cause human infertility (55). Moreover, the pathogenic role of compound heterozygous DNS or LOF variants of RBPs was documented (25, 75, 76). We also note that the odds of finding a DNS variant in CCs and ER patches (3E3R) of RBPs are higher in patients compared to controls, at least statistically significant for the CCs,

whereas the ER patches did not show a significant difference and should be revisited with a larger cohort in the future. These findings are beginning to link nondomain elements with human infertility.

Materials and methods

mMGC purification

We isolated each type of spermatogenic cells from C57BL/6J mouse testes by STA-PUT velocity sedimentation (77, 78). Pachytene spermatocytes (pSC) and round spermatids (rST) were isolated simultaneously from 16 adult testes per round of STA-PUT; spermatogonia (SG) were purified from 80 pairs of P6-8 testes per round. Testes were decapsulated, washed with DMEM twice, and incubated with 1 mg/ml collagenase IV in DMEM in 37°C water bath for 15-20 min. Seminiferous tubules were then gently pipetted to disperse into the solution, washed twice with DMEM, and collected after sedimentation by gravity. The sedimented tubules were further digested with 0.25% trypsin containing 1 mg/ml DNase I and gently pipetted up and down for 5-15 min before adding 10% FBS. Cells were pelleted by centrifugation at 500×g for 5 min and then resuspended with wash buffer (10% FBS, 0.5% BSA, 200 µg/µl DNase I), and filtered with 40 µm cell strainer to obtain a monodisperse suspension. The single-cell suspension was loaded into a cell separation apparatus (ProScience, Canada), layered over a linear gradient (2%-4%) of bovine serum albumin (BSA) in DMEM, and then allowed to sediment by gravity for 2.5-3 hours. Finally, the cells were fractionated, type-determined, and pooled into quasi-homogeneous populations. Cell purity was checked under DIC light microscope based on distinctive diameters and morphological features, and selectively validated under immunofluorescence microscope based on staining patterns of DAPI and/or protein marker antibodies. Note that the spermatogonial preparations contain no spermatocytes, which are not supposed to appear in P6-8 testes. pSC may be contaminated by other similar-sized meiotic cells such as zygotene and diplotene spermatocytes, and rST preparations may be contaminated by elongated spermatid. To achieve sufficient cell numbers qualified for RIC and proteomics, the STA-PUT procedure was repeated. Specifically, for each replicate of RIC, 20 rounds of STA-PUT were performed to collect $\sim 2.5 \times 10^8$ SG, 14 rounds to collect $\sim 2.0 \times 10^9$ rST, and 28 rounds (14 rounds synchronous with rST, plus 14 extra rounds) to collect $\sim 5.0 \times 10^8$ pSC, respectively. A total of 96 rounds of STA-PUT provided two replicates of RIC for each cell type.

mMGC RIC

RIC was carried out according to a previously described protocol (79), with modifications for freshly purified mMGCs. Single types of cells were pooled and washed with ice-cold PBS 3 times to thoroughly remove DMEM. Before ultraviolet

(UV) crosslink, we sought to remove excess liquid buffer that could not be easily pipetted out in order to minimize the loss of UV energy, considering that we used suspended cells rather than regularly cultured cells adherent tightly to the plate bottom. To this end, we spread 1 ml buffer with suspended cells through 4 or 5 10 cm dishes sequentially. The cells were allowed to settle for a while in each dish and then transferred with most of the buffer to the next dish, with a considerable portion of cells left in a tiny amount of the remaining buffer. Microscopy inspection showed that most settled cells dispersed well and adhered to the dish bottom. Each dish without its lid was placed on ice at a distance of 10-15 cm from the UV lamp and irradiated with 0.15 J/cm² at 254 nm UV light in a UV crosslinker device (CL-1000). Every time, half the number of cells were irradiated with UV, and in parallel, the other half were not UV-crosslinked. The cells were resuspended in ice-cold PBS, pelleted in a conical tube by centrifugation at 500×g for 5 min, snap-frozen in liquid nitrogen, and stored at -80°C until use. After collection for RIC, the frozen pellets were thawed and combined using ice-cold lysis buffer (20 mM Tris pH 7.5, 500 mM LiCl, 0.5% lithium dodecyl sulfate, 1 mM EDTA, 5 mM DTT). Oligo dT₂₅ beads (NEB, S1419S) were used to capture polyA RNA. After 8 stringent washes, covalently crosslinked RNA-protein complexes were released from beads by 55°C incubation. Protein eluates obtained from two RIC replicates for each cell type were preserved for mass spectrometry.

Testis RBDmap

RBDmap was performed following a previously described protocol (19), with adaptations for fresh testicular cells. For each RBDmap replicate, 6 adult testes and 24 P28 testes were decapsulated and separately subjected to collagenase IV and trypsin enzyme digestion to obtain a mixed population of single cells. As detailed for RIC, cells were resuspended with minimal ice-cold PBS and settled to the dish bottom to facilitate UV crosslinking. RNA-protein complexes were first captured on oligo(dT) beads following standard RIC procedure. The RNA-protein complexes were then digested with Lys-C (WAKO, 125-05061) in RBP elution buffer: 20 mM Tris-HCl (pH 7.5), at 200 rpm in thermo mixer at 37°C for 8 hours, generating primary peptides which harbor all the RNA-bound and release peptides. New Oligo dT₂₅ beads were incubated with the bound groups at 4°C for 1 hour. The beads were then washed with 1 × hybridization buffer (200 mM Tris-HCl pH 7.5, 5 M LiCl, 2 mM EDTA, 50 mM DTT), and the supernatant was saved as the release-peptide sample. The beads were sequentially washed by lysis buffer (20 mM Tris-HCl pH 7.5, 500 mM LiCl, 0.5% LiDS, 1 mM EDTA, 5 mM DTT), Wash-1 buffer (20 mM Tris-HCl pH 7.5, 500 mM LiCl, 0.1% LiDS, 1 mM EDTA, 5 mM DTT), Wash-2 buffer (20 mM Tris-HCl pH 7.5, 500 mM LiCl, 1 mM EDTA, 5 mM DTT), and

Low salt buffer (20 mM Tris-HCl pH 7.5, 200 mM LiCl, 1 mM EDTA, 5 mM DTT). The beads were resuspended in an elution buffer to elute the RNA-peptide complex with 500 rpm in a thermomixer at 55°C for 10 min, and the supernatant was saved as the bound-peptide sample. New Oligo dT₂₅ beads were added in primary peptide samples for another two rounds of bound and release peptide capture. All the generated bound and release peptide samples were pooled separately for mass spectrometry.

LC-MS/MS analysis

The RIC and RBDmap proteins were concentrated using Amicon Ultra Centrifugal Filters (10 kDa cutoff, Millipore) and subsequently reconstituted in lysis buffer (8 M urea, 75 mM NaCl, 50 mM Tris pH 8.2, 1% (v/v) EDTA-free protease inhibitor cocktail, 1 mM NaF, 1 mM β -glycerophosphate, 1 mM sodium orthovanadate, 10 mM sodium pyrophosphate, 1 mM PMSF). The RIC and RBDmap protein lysates were then subjected to protein digestion, followed by peptide desalination. Briefly, cysteine residues were reduced by 5 mM DTT for 25 min at 56°C followed by alkylated using 14 mM iodoacetamide for 30 min at room temperature in the dark, and quenched with DTT for an additional 15 min. Samples were then diluted to 1.6 M urea with 25 mM Tris-HCl, pH 8.2, and digested overnight at 37°C with the addition of 2 μ g trypsin. Following digestion, peptide mixtures were acidified by trifluoroacetic acid (TFA) (0.4% vol/vol) (Sigma-Aldrich), desalted using a SepPak 1cc tC18 cartridge (Waters), and then lyophilized. For RIC analysis, purified peptide samples were resuspended in Buffer A (20 mM ammonium formate, pH 10), fractionated using an XBridge BEH C18 column (Waters) to generate a total of 8 fractions, and dried in vacuum.

For analyses of mMGC RIC peptides, the peptide samples were resuspended in 0.1% formic acid (FA) and analyzed using a LTQ Orbitrap Velos mass spectrometer (Thermo Fisher Scientific) coupled online to a Proxeon Easy-nLC 1000. Peptides were loaded onto a trap column (75 μ m $\times$ 2 cm, Acclaim PepMap100 C18 column, 3 μ m, 100 Å; DIONEX, Sunnyvale, CA) and separated by a reverse-phase microcapillary column (75 μ m $\times$ 25 cm, Acclaim PepMap RSLC C18 column, 2 μ m, 100 Å; DIONEX, Sunnyvale, CA) at a flow rate of 300 nl/min. The HPLC solvent A was 0.1% FA, and the solvent B was 100% acetonitrile (ACN), 0.1% FA. A 95-min linear gradient (3% to 8% buffer B for 3 min, 8% to 29% buffer B for 71 min, 29% to 41% buffer B for 12 min, 41% to 90% buffer B for 1 min, 90% buffer B for 8 min) was used for peptide analysis in the data-dependent acquisition (DDA) mode. An MS survey scan was obtained for the m/z range 350-1800 at a resolution of 60,000, and MS/MS spectra were acquired from the survey scan for the 20 most intense ions.

For analyses of testis RBDmap peptides, the peptide samples were resuspended in 0.1% FA and separated on an

analytical column (75 μ m $\times$ 160 mm, 1.9 μ m, Dr. Maisch) at a flow rate of 300 nl/min in a 95 min linear gradient (3% to 5% buffer B for 5 s, 5% to 15% buffer B for 40 min, 15% to 28% buffer B for 34 min 50 s, 28% to 38% buffer B for 12 min, 30% buffer to 100% buffer B for 5 s, and 100% buffer B for 8 min) using buffers of 0.1% FA (buffer A) and 80% ACN, 0.1% FA (buffer B). The effluents from LC system were then analyzed using Orbitrap Fusion Lumos mass spectrometer (Thermo Fisher Scientific) operated in DDA mode. A full survey scan was obtained for the m/z range of 350-1500 with 60,000 resolution, and MS/MS was performed by high energy dissociation (HCD) in the Orbitrap with a resolution of 15,000.

Antibodies, Western blots, and immunofluorescence

Commercial antibodies: NONO (Abcam, Ab70335; Abclonal, A3800; Sangong, D199144); SAFB (Abclonal, A7927); MSH6 (Abclonal, A8795); HA (MBL, M180-3; CST, 3724S); CEP72 (Proteintech, 19928-1-AP); CENPV (Sangon, D162083); SYCP3 (Abcam, ab97672); PNA (Vector Labs, RL-1072); HNRNPQ (Abcam, Ab189405); MOV10 (Proteintech, 10370-1-AP); MILI (Abcam, Ab36764); MIWI (Abcam, Ab181056); GAPDH (Abclonal, AC002); ACTIN (Sigma, SAB5701411); TUBULIN (Abclonal, AC008); FYCO1 (Sangong, D126224); TEK5 (Proteintech, 26108-1-AP); LIN28A (Abcam, ab279647); SOX9 (Millipore, AB5535); DNMT3A (Huabio, ET1609-31); ZBTB16 (R&D, AF2944). BOULE and PUM1 antibodies were gifts from Eugene Yujun Xu Lab (Nanjing Medical University). TEX30 and PDHA2 antibodies were generated by Abclonal.

For Western blot analysis, testes or 293T cells were homogenized in RIPA buffer (50 mM Tris-HCl pH 8.0, 150 mM NaCl, 5 mM MgCl₂, 0.5% sodium deoxycholate, 1% Triton X-100, 2% SDS, 10% glycerol) added with protease inhibitor. The lysates were centrifuged at 12,800 rpm at 4°C for 30 min, and the supernatant was collected. The proteins were denatured by adding SDS loading buffer, heated for 10 min at 95°C, and loaded on SDS-PAGE. Immunofluorescence analysis of testis sections was performed as described previously (80). TUNEL analysis of testis sections was performed with the In situ Cell Death Detection Kit (Roche, 11684817910) according to the manufacturer's protocol.

Radioactive CLIP

Conventional CLIP of RBPs (PDHA2, TEX30, NONO, and PSPC1) was performed as previously described (80). For CLIP using testicular tissues, 100 mg juvenile testes from P21 (for PDHA2) or P28 (for TEX30) mice were detunicated and mildly pipetted to form discrete single seminiferous tubules in ice-cold PBS, and immediately UV-crosslinked with 0.4 J/cm² at 254 nm for 3 times. For CLIP using cell lines, HA-tagged NONO or PSPC1 was transfected into 293T cells in 10 cm dish. The dish was washed by cold PBS and UV-crosslinked with 0.12 J/cm² at 254 nm for one time.

Crosslinked samples were washed with cold PBS and pelleted in a conical tube by centrifugation at 500×g for 5 min. Pellets were snap-frozen in liquid nitrogen and stored at -80°C until use. To begin CLIP, briefly, pellets were dissolved with 1× PMPG (1× PBS, 2% Empigen) containing protease inhibitors (Roche, 11873580001) and RNA inhibitors (Promega, N2518), and then treated with DNase I. Then protein-RNA complexes were immunoprecipitated by 10 µg antibody using protein A Dynabeads (Invitrogen, 10002D) for rabbit antibody or protein G Dynabeads (Invitrogen, 10003D) for mouse antibody. Then RNA was labeled with [γ -³²P] ATP and run on NuPAGE 4-12% Bis-Tris (Life Technology, NP0323BOX). The isotopic crosslinked RNA-protein complexes were transferred onto nitrocellulose membrane (Invitrogen, LC2001) and detected by a phosphor screen system (Typhoon).

eCLIP-seq

eCLIP-seq was performed as previously described (81, 82). Crosslinked testicular samples for eCLIP of RBPs (PDHA2, TEX30, NONO, and NONO^{ΔER}) were prepared as for conventional CLIP. HA-tagged NONO and NONO^{ΔER} were transfected into 3T3-L1 cells in 10 cm dish and crosslinked as conventional CLIP. To begin eCLIP, briefly, samples were dissociated in eCLIP lysis buffer containing protease inhibitor (Roche, 04693159001), RNase inhibitor, RNase I (Life Technology, AM2295), and Turbo DNase (Invitrogen, AM2239). Crosslinked target RNPs were captured on Dynabeads using 10 µg antibodies. Protein-bound RNAs were then dephosphorylated and ligated with 3' RNA adapter. eCLIP and input samples (collected before immunoprecipitation) were then washed stringently, boiled from beads, run on NuPAGE 4-12% Bis-Tris (Life Technology, NP0323BOX), and transferred onto nitrocellulose membrane (Amersham, 10600008). Size-matched subregion of the membrane was collected and proteins were degraded by Urea and proteinase K (NEB, P8107S). Harvested RNAs were subsequently reverse-transcribed into cDNA with Affinity Script (Agilent, 600107), and purified by Exo-Sap-IT (Affymetrix, 78201). The 3' end of cDNA was ligated with a DNA adapter by T4 RNA Ligase (NEB, M0437M). Libraries were then PCR-amplified using Q5 Polymerase and run on 3% low-melting temperature agarose gel. Sizeable libraries were purified with a MinElute gel extraction kit (Qiagen, 28604) and run on the Illumina platform.

scRNA-seq libraries

Mouse testes were washed in ice-cold RPMI1640 and dissociated using Collagenase II (Sigma, V900892-100MG) and DNase I (Sigma 9003-98-9). Cell count and viability was estimated using fluorescence Cell Analyzer (Countstar Rigel S2) with AO/PI reagent after removing erythrocytes (Solarbio R1010), debris, and dead cells (Miltenyi 130-109-398/130-090-

101). Finally fresh cells were washed twice in RPMI1640 and then resuspended at 1 × 10⁶ cells per ml in 1× PBS with 0.04% bovine serum albumin. scRNA-seq libraries were prepared using SeekOne Single Cell Full-length RNA sequence Transcriptome-seq Kit according to the manufacturer's instructions (SeekGene, Catalog No.K00801). Briefly, an appropriate number of cells were mixed with reverse transcription reagents and added to the sample wells of the SeekOne DD Chip S3. Then, Barcoded Hydrogel Beads (BHBs) and partitioning oil were dispensed into corresponding wells separately in Chip S3. Subsequently, cell-containing reverse transcription reagents and BHBs were encapsulated into emulsion droplets using SeekOne Digital Droplet System. Immediately after transferring emulsion droplets into PCR tubes, 15 cycles of annealing (ramping from 8°C to 42°C) followed by a 5 min heat inactivation at 85°C were performed to obtain barcoded cDNA. Next, the barcoded cDNA was purified from broken droplet and PCR reactions were performed twice to remove the majority of ribosomal and mitochondrial cDNA. AMPure beads were used to purify cDNA from the post-PCR reaction mixture. Finally, one fourth volume of cDNA was fragmented, end-repaired, A-tailed and ligated into sequencing adaptor. DNA amplified by index PCR contained any part of polyA or non-polyA RNA and Cell Barcode and Unique Molecular Index. The indexed sequencing libraries were purified using AMPure beads and quantified by quantitative PCR (KAPA Biosystems KK4824). The libraries were then sequenced on Illumina NovaSeq 6000 with PE150 read length.

2C assay

2C assay was performed as previously described (31, 32), with modifications. Crosslinked testicular single-cell mixtures were prepared as described for testis RBDmap, and cross-linked 293T cells were prepared as described for conventional CLIP. The primer sequences for cloning are listed in table S16. Samples were then processed using an Allprep DNA/RNA mini Kit (QIAGEN, 80204). Briefly, crosslinked samples were lysed in RLTplus buffer (QIAGEN, 1053393) with 1% beta-mercaptoethanol using an ultrasonic homogenizer for 10 s on/10 s off for about 40 cycles until lysate became transparent. To eliminate DNA, the lysate was loaded on the Allprep DNA mini spin column and centrifuged at 3000×g for 5 min. An equal volume of 70% ethanol was added to the flowthrough, mixed well, loaded onto RNeasy mini spin column, and centrifuged at 3000×g for 5 min. The RNeasy column was then washed in sequential with RW1 and RPE wash buffers by centrifugation at 3000×g for 3 min. 200 µl nuclease-free water was added to RNeasy column and incubated for 3 min. The free RNA and RNP complexes were eluted in nuclease-free water and collected by centrifugation at 3000×g for 3 min. RNA concentration was measured by Nanodrop 2000. For quantitative 2C (q-2C) assays, the RNA

concentration of the sample was required to vary less than 10% of that of the control. Samples were then dried into powder by vacuum centrifugal concentrator (Centivap) and re-suspended in 30 μ l nuclease-free water. RNAs were digested by adding 1 μ l 10 mg/ml RNase A (Thermo, EN0531) or 1 μ l benzonase (Merck, E8263) at 800 rpm in thermomixer for 12 hours. Samples were then subjected to Western blot analysis.

For in vitro q-2C assay, 500 ng purified NONO or NONO Δ^{ER} proteins and 1 μ g total RNAs isolated from 293T cells were incubated in 2 $\times$ RNA binding buffer (10 mM HEPES pH 7.4, 50 mM KCl, 4 mM MgCl₂, 4 mM DTT, 0.2 mM EDTA, 7.6% glycerol, 3 mM ATP) for 30 min, transferred into a well of a 96-well plate, and UV-crosslinked with 5J at 254 nm for 10 min. 8 wells of protein-RNA mixtures were then pooled and subjected to the same q-2C assay.

mRNP capture assay

mRNP capture assay was performed as a small-scale RIC (10). Briefly, transfected 293T cells in 10 cm dishes were UV-crosslinked and lysed with 50 μ l mRNA interactome lysis buffer as in RIC. 50 μ l Oligo dT₂₅ beads were used to capture polyA RNAs. After 8 rounds of stringent washes, proteins were eluted from Oligo dT₂₅ beads and then subjected to Western blot analysis.

Protein expression and purification

All coding sequences (CDS) were PCR-amplified from mouse testis cDNA. Full-length CDS sequences of PDHA2 and TEX30 were separately cloned into the pET28a expression vector. CDS of mouse NONO (1-473) and NONO Δ^{ER} (1-346, 355-473) were cloned into the pET22b expression vector. The primer sequences for cloning are listed in table S16. All constructed plasmids were transformed into *E. coli* BL21. Different proteins (PDHA2, TEX30, NONO, and NONO Δ^{ER}) were expressed and purified with differential modifications of our previous protocol (83) (Supplementary methods). Protein concentration in the supernatant was measured by Nanodrop 2000 (Thermo) on a 280 nm absorption peak. Proteins were stored at -80°C until use.

Nono Δ^{ER} and Safb Δ^{ER} knock-in mice

Knock-in mice with precise ER truncation were generated by GemPharmatech using the CRISPR-cas9 system. sgRNA sequences were designed to direct Cas9 endonuclease to the front of Exon 9 of *Nono* to generate *Nono* Δ^{ER} allele (Fig. 5A) and to the Exon 15 of *Safb* to generate *Safb* Δ^{ER} allele (fig. S15A), respectively, by homologous recombination. Mouse zygotes were co-injected with donor, Cas9 mRNA, and sgRNA, and transferred into surrogate mothers' endometrium. All animals were handled as stipulated by the Guide for the Care and Use of Laboratory Animals at Nanjing Medical University (ethics approval ID: 2101008 and ID: 2102008).

Human male infertility diagnosis and cohort information

A total of 1046 unrelated infertile men (ranging from 19 to 51 years old) who displayed unexplained (idiopathic) NOA (n = 516) or oligozoospermia (n = 530) were recruited between June 2017 and March 2022 at the Reproductive and Genetic Hospital of CITIC-Xiangya (Changsha, China) (table S13). A group of 300 people (Han Chinese) (ranging from 20 to 73 years old) with normal childbearing history was referred to as the control group (table S13). This study was approved by the ethics committee of CITIC Xiangya Reproductive Genetics Hospital (LL-SC-2017-025 and LL-SC-2019-034).

All patients received an inquiry for detailed history to rule out those male individuals with clear causes, including iatrogenic injury, reproductive tract infection, testicular inflammation, drug exposure and other factors (84). We also excluded other testicular injuries, such as testicular trauma and testicular torsion. Physical examination, including height, weight, hair distribution, mental status, and external genitalia, was conducted and all men in our cohort were normal. Genetic screening for karyotyping (G-bands at the 320-400-band level) and Y chromosome AZF region deletions (sY84 and sY86 for AZFa, sY127 and sY134 for AZFb, sY254 and sY255 for AZFc) was performed according to Reproductive Medicine Professional Committee of Chinese Medical Doctor Association. In our cohort, no patients had Y chromosome AZF microdeletions or chromosomal abnormalities.

Semen analysis was conducted according to the guidelines of the World Health Organization (WHO) Laboratory Manual for the Examination and Processing of Human Semen (85). The individuals were diagnosed with idiopathic male infertility based on the reference (84).

Azoospermia was defined as a condition in which no spermatozoa were detected on three consecutive occasions when ejaculated semen was examined microscopically after centrifugal sedimentation (86). A non-obstructive etiology was established on the basis of the following parameters: hormone values, pH and volume of semen, testicular volume, and testicular biopsy according to guidelines (87, 88) (n = 509). For some patients (n = 7), although other parametric information was insufficient to differentiate between obstructive and non-obstructive etiologies, they were also recruited as the latter based on their normal vas deferens. Any azoospermia patients with previous vasectomy, congenital absence of a vas deferens, vasal agenesis or obstructive vas deferens were excluded from our cohort (table S13).

Oligozoospermia was defined as a condition in which the concentration of spermatozoa in the ejaculated semen was below the lower limit of the reference value for semen analysis in normally fertile men (85). All the oligozoospermia patients in our cohort were further classified into two subtypes based on sperm concentrations: severe oligozoospermia: 0.1-

$5 \times 10^6/\text{mL}$ ($n = 503$); extreme oligozoospermia: $< 0.1 \times 10^6/\text{mL}$ ($n = 27$) (table S13).

Whole-exome sequencing

Genomic DNA (gDNA) from peripheral blood samples was extracted using a QIAamp DNA Blood Midi kit (Qiagen, 51106) according to the manufacturer's protocol. The qualified gDNA samples of the 1046 infertile men and 300 fertile controls were subjected to library construction, followed by WES on Illumina HiSeq 2000 or HiSeq X-TEN platforms (89).

Bioinformatics

RIC proteomic analysis

The mass spectrometric raw data were searched with MaxQuant (version 1.5.2.8), identified tryptic peptides assigned to contaminants were removed, and only peptides uniquely mapped to a gene model were considered. High-confidence RIC RBPs were defined as those with intensity based absolute quantitation (iBAQ) in crosslinked (CL) pull-downs at least 10-fold higher than noncrosslinked (noCL) pull-downs in both RIC replicates.

GO annotation was obtained from Ensembl 75. RNA-associated GO annotation used R package mRNA interactomeHela. More specifically, proteins annotated by the GO-term “RNA-binding” and by other GO-terms related to RNA biology were classified as RNA-binding and RNA-related proteins, respectively (10). The rest of the RBPs were classified as RNA-unrelated proteins.

A manually curated list of canonical and noncanonical RBDs was used to classify RIC RBPs as harboring a canonical or noncanonical RBD (table S1). RBPs were annotated as classical if they contained at least one canonical RBD, or classified as nonclassical if they did not contain canonical RBDs but harbored noncanonical ones or were listed in a manually curated nonclassical RBP table from Liao *et al.* (30), or classified as novel otherwise. With reference to a manually curated list of published mouse somatic cell (mSC) RBPs (30), we classified mMGC RBPs (this study) into three families: mSC-specific, mMGC-specific, and common.

Tissue specificities of mMGC RBPs were determined by the expression specificity score (SPM) estimated from RNA-seq profiling data across 17 mouse tissues (90), as described in TiSGeD (91). Genes with SPM score ≥ 0.9 were classified as testis-specific, genes with SPM score ≥ 0.5 and < 0.9 were classified as testis-enriched, and other genes were classified as widely expressed.

Human and mouse orthologous gene mapping

The mapping of orthologous genes between human and mouse was conducted by orthologsBioMART R package (92).

Gene ontology enrichment/network analysis

The GO enrichment was analyzed by either R package clusterProfiler (with $p.\text{adjust} < 0.05$) (93), or R package GSEA (with $p.\text{adjust} < 0.05$) (94), or StringDB web-based platform (95). GO enrichment network was analyzed by Metascape web-based platform (with $p\text{-value} < 0.05$) (96).

TMT-based quantitative proteomics

TMT-MS data, including four spermatogenic stages of mMGCs (SG, pSC, rST, and eST) with four biological replicates for each, were analyzed using MaxQuant (version 1.2.2.5) software with UniProt proteomes (mouse) protein database, as described previously (33). The normalized protein abundance for each protein was used for further bioinformatics analyses. Hereon, unless specially indicated elsewhere, proteins listed as mMGC TMT-MS proteins and not as mMGC RBPs were defined as nonRBPs.

To determine temporal pattern of protein abundance across four consecutive stages, we softly clustered proteins into ten groups based on the average normalized protein abundance using the Mfuzz package in R (97).

Relative RNA-binding activity analysis

The estimation of relative RNA-binding activity refers to Kilchert *et al.* (37). RBPs not detected by TMT-MS in any single cell type were filtered out. Expression of RBPs with multiple protein isoforms were aggregated by taking the average. In total, there were 1218 unique RBP genes left for subsequent relative RNA-binding activity analyses. Raw WCE and RIC intensity were Log2-transformed and scaled within each sample. Scaled Log2 (RIC/WCE) was processed with the eBayes function in the limma R package to quantify relative RNA-binding activity of each RBP, and moderated *t* test was carried out to examine significant alteration in relative RNA-binding activity between conditions.

Scoring intrinsic disorder of amino acids

For all amino acids in a protein, a score for intrinsic disorder was computed using IUPred (98). Amino acid residues with a score larger than 0.4 were called disordered. For each protein, the proportion of disordered residues was estimated.

RBDmap proteomic analysis

The mass spectrometric raw data were processed using MaxQuant (version 1.6.5.0) (99), and mass spectrometric spectra were searched using the Andromeda search engine against mouse proteins in UniProt. MS-identified tryptic peptides assigned to contaminants were removed, and only peptides uniquely mapped to a gene model were considered. A total of 5754 peptides were retained after quality control (table S7). MS-identified tryptic peptides enriched in the RNA-bound or released fractions, respectively, were mapped back to the whole proteomes of the mouse (UniProt) and extended

to the two adjacent Lys-C cleavage sites to recall the original proteolytic fragment. A peptide with the average LFQ.intensity of three replicates in Log1p ratio (logarithm ratio of the LFQ.intensity of Bound to Release after adding a pseudocount of 1) greater than 10 was considered RBDpeps. Finally, overlapping RBDpeps of each protein were merged. RBDpeps between protein isoforms were combined, and the Uniprot ID of canonical isoform was retained.

We used three criteria to filter RBDpeps of different levels of confidence. Under the strictest criterion, we selected those with Log ratio of the average LFQ.intensity of three replicates in Bound to Release > 0 at an FDR < 1%, which generated 361 non-redundant RBDpeps (from total 663 RBDpeps) as Stringent RBDpeps. Under the less strict criterion we selected those with the criterion Log1p ratio of the average LFQ.intensity of three replicates in Bound to Release > 10, which generated 721 non-redundant RBDpeps (from total 1533 RBDpeps) as Moderate RBDpeps. Under the least strict criterion, we selected those with Log1p ratio of the average LFQ.intensity of three replicates in Bound to Release > 0, which generated 793 non-redundant RBDpeps (from total 1859 RBDpeps) as Loose RBDpeps.

To annotate RBDpeps, each identified RBDpep was examined whether it overlaps with a Pfam domain (<http://ftp.ebi.ac.uk/pub/databases/Pfam/>) (100), disorder regions, or CC motifs. Note that CCs adopt an alpha-helix form and are predicted based on Pfam annotation. RBDpeps that overlap with the Pfam domain were considered domain RBDpep, and RBDpeps without overlap with Pfam domain were considered nondomain RBDpeps. Further, nondomain RBDpeps overlapped with Pfam disorder regions, and CC regions were classified as disorder region RBDpeps and CC RBDpeps, respectively. A manually curated list of published mouse somatic cell RBDpeps (42) was used to compare with testis RBDpeps from this study. For sequence motif search in nondomain RBDpeps, multiple sequence alignment (MSA) was built by Clustal Omega (<https://www.ebi.ac.uk/jdispatcher/msa/clustalo>) with default parameters. Motifs were extracted from align part of MSA and visualized by WebLogo (<https://weblogo.threeplusone.com/>) with amino acids colored according to their side-chain chemistry.

For sequence motif enrichment in CC regions of mMGC RBPs and nonRBPs, motif was searched using MEME software. For a motif to be defined, we required the occurrence of the same pattern to be more than 20 times and the P value to be less than 0.00001.

To test enrichment of bound nondomain RBDpeps within CC, disorder, and low complexity regions, we performed Fisher's exact test using Locus Overlap Enrichment Analysis (LOLA) R package with default setting (Version 1.26.0) (101). All nondomain RBDpeps were the user set of LOLA analysis,

all bound RBDpeps served as the user universe of LOLA analysis, and bound RBDpeps within the disorder regions/CC regions/low complexity regions served as the input file of LOLA analysis.

Scoring evolutionary conservation of protein residues

We obtained the normalized evolutionary conservation score for residues on each RBD-mapped protein using the web-based application ConSurf 2016 (UNIREF-90 protein database with default parameters) (102). Local conservation scores were calculated by sliding a ten amino acid window along each protein and averaging the conservation score.

Screening for ER patches in proteins

We defined two classes of ER patches: at least 4 Glu and 4 Arg (4E4R) or at least 3 Glu and 3 Arg (3E3R) within a ten amino acid window, and thus the former was the subset of the latter. To screen for these two classes of ER patch throughout mMGC RBPome and CC sequences of nonRBPome, we extracted subsequences using seqkit (103) sliding with step size 1 amino acid and window size 10 amino acids. Then we extracted the subsequences and protein coordinates with 3E3R and 4E4R patches, and merged the overlapping subsequences to obtain the final 3E3R and 4E4R patch within mMGC RBPome and CC sequences of nonRBPome. The procedure for screening ER patches in human orthologous proteins of mMGC RBPome and nonRBPome is the same as that in mMGC RBPome and CC sequences of nonRBPome.

Mapping protein-relative coordinates to genomic regions

We used geno2proteo (104) R package to map protein-relative coordinates to genomic regions of human and protein-relative coordinates to genomic regions of mouse. We used UCSC Genome browser software liftover to convert genome coordinates between different species.

Co-evolution analysis

For each protein containing defined ER patches, evolutionary couplings between residues were computed by EVcouplings on local and web servers (<https://evcouplings.org>) (46). Residue pairs with stronger co-constraints or interdependence are assigned higher co-evolution scores. As suggested (105), within each protein we defined strongly evolutionarily coupled pairs (ECs) as the top L ones, where L is the protein sequence length, and then counted the number of ECs within each ER patch. Note that nonRBPs were defined as those in TMT RBPome but out of mMGC RBPome and mSC proteome.

eCLIP-seq data analysis

eCLIP-seq data were analyzed following the ENCODE standard eCLIP pipeline (ENCODE Project Consortium, 2012;

<https://www.encodeproject.org/>) (106). On the basis of trimming adapters and 3' adapter dimers using cutadapt (V1.14), we first dropped out reads aligned to the repeat elements from UCSC with STAR (V2.7.6a) and then mapped the rest to mm10 reference genome (GENCODE vM23). PCR duplicates were removed. CLIPPER was used to call peak regions with fold enrichment ≥ 8 and $\text{P}_{\text{adj}} \leq 0.001$. In the end, we identified significantly enriched peaks as those with irreproducible discovery rate cutoff of 0.01. Binding motif among enriched peaks was predicted by MEME with motif length from 5 to 8. All annotations referred to GTF file from GENCODE version vM23.

To compare the NONO binding intensity in cellulo and in vivo, we used the htseq-count software (107) to quantify gene level count in the eCLIP IP (2 replicates in cellulo and 3 replicates in vivo) and input samples (no replicates). Unlike expression data, eCLIP reads were enriched within a few hundred to two thousand genes, so we normalized the raw count of genes of strong signals (Log of count per million ≥ 10) using the edgeR TMM algorithm to avoid normalization bias introduced by thousands of genes of insufficient coverages. The binding intensity is the Log fold change value between the IP and input normalized count.

scRNA-seq data analysis

Data processing was primarily conducted with Seurat R package (version 4.3.0). Poor quality cells were excluded, such as cells with <200 or $>8,000$ expressed genes (as they are potentially cell doublets). Only genes expressed in 3 or more cells were used for further analysis. Cells were also discarded if their mitochondrial gene percentages were over 5%.

Raw expression data were normalized and scaled with NormalizeData function, and ScaleData function. The top 1,000 genes with the highest standardized variance were identified using FindVariableFeatures function (selection.method = "vst"). Principal component analysis (PCA) was computed using RunPCA function with default parameters.

Shared nearest neighbor (SNN) graph was computed using the FindNeighbors function, taking as input the first 20 PCA dimensions. Cell clusters were defined using Louvain algorithm with the FindCluster function. For visualization uniform manifold approximation and projection (UMAP) was used. Cell-type-specific markers were identified by the Seurat FindAllMarkers command, with Log fold change threshold of 0.25 and minimum fractional expression threshold of 0.1. For accurate differential expression analysis between conditions, we used the pseudobulk method to aggregate expression of each sample (108), because treating each cell as an independent sample can lead to underestimated variance, overly small p-values, and thus misleading results (109). Aggregated expressions were normalized by calcNormFactors function in

the edgeR packages, and differentially expressed analyses were performed using the limma package. Developmental trajectories were predicted by slingshot (110). Mitosis-enriched genes were defined as those specifically expressed in SG and preleptotene spermatocytes, while meiosis-enriched genes were those specifically expressed in meiotic spermatocytes.

Whole-exome sequencing data processing

The raw sequencing data were processed using the following steps: (1) Removing reads containing sequencing adapter; (2) Removing reads whose low-quality base ratio (base quality ≤ 5) is more than 50%; (3) Removing reads whose unknown base ("N" base) ratio is more than 10%. Statistical analysis of data and downstream bioinformatics analysis were performed on this filtered high-quality data, referred to as the clean data. Clean data were aligned to the human reference genome (GRCh37) using Burrows-Wheeler Aligner (BWA) (version: 0.7.17-r1188) (111). Picard was used to remove duplicated sequence reads. Realignment was performed with the Genome Analysis Toolkit (GATK) (112). Single-nucleotide polymorphisms (SNPs) and insertions-deletions (InDels) were called using HaplotypeCaller of GATK. SNPs and InDels within low complexity regions (hg19_blackregion.bed) were removed. ANNOVAR was used to annotate the variants for functional and population frequency information (113). Deleterious missense variants were predicted with combined annotation-dependent depletion (CADD) (v1.3) score (114). CADD_phred > 15 was considered deleterious. The RefSeq gene model was used throughout this study. Genes listed in TMT-MS data were applied in subsequent analyses. For clinical analysis, we classified genes listed in mMGC RBPome and testis RBDmap as RBP genes; genes listed in TMT-MS data but not list in mMGC RBPome or RBDmap as nonRBP genes. To assess tissue specificity of human genes, RNA-seq profiling data across 54 human tissues were extracted (<https://www.proteinatlas.org>, (rna_tissue_consensus.tsv)) to calculate the expression specificity score (SPM) of each RBP gene, as described in TiSGeD (91). Genes with SPM score ≥ 0.9 were classified as testis-specific, genes with SPM score ≥ 0.5 and < 0.9 were classified as testis-enriched, and other genes were classified as ubiquitously expressed.

To examine the population structure and check sample anomaly of the patient and control groups, we performed principal component analysis (PCA) based on the genotypes of the called variants in all samples in this study and in hapmap3 (<https://www.broadinstitute.org/medical-and-population-genetics/hapmap-3>) using plink software (115) and visualized the first two components in a two-dimensional space. We performed relatedness examination by KING(v.2.2.4) (116) and showed all samples have no relatedness to each other in the dataset.

Candidate risk variants and gene filtering

Candidate causative rare loss of function (LOF) variants were defined using the following criteria: (1) filter is “PASS”, read depth ≥ 10 , GQ ≥ 20 ; alternative read depth/ read depth ≥ 0.2 ; (2) allele frequencies in 1000 g2015aug_eas, ExAC_EAS, gnomAD_genome_EAS, and gnomAD_exome_EAS public databases were lower than 0.01 or not documented; (3) frameshifts, premature stop gains or the disruption of core splice sites. Rare LOF pathogenic mutation genes with no LOF and rare DNS variant genes hit in control individual were defined as rare LOF pathogenic mutation genes.

Candidate causative rare damaging nonsynonymous (DNS) variants were defined using the following criteria: (1) filter is “PASS”, read depth ≥ 10 , GQ ≥ 20 ; alternative read depth/ read depth ≥ 0.2 ; (2) allele frequencies in 1000 g2015aug_eas, ExAC_EAS, gnomAD_genome_EAS, and gnomAD_exome_EAS public databases were lower than 0.01 or not documented; (3) “nonsynonymous SNV”. Rare DNS pathogenic mutation genes were defined as Rare DNS variants with a Phred-scaled CADD pathogenicity prediction score > 15 and no rare DNS mutation genes hit in control individual.

Prioritization of the genetic components more relevant to the pathogenesis

We classified candidate variants into the following: homozygote/hemizygote (Hom) LOF variants of RBP, heterozygote (Het) LOF variants of RBP, Hom DNS variants of RBP, Het DNS variants of RBP, domain region DNS of RBP, nondomain region DNS of RBP, CC region DNS of RBP, nonCC region DNS of RBP, ER patch DNS of RBP, CC-nonER region DNS of RBP, and among others belonging to nonRBP.

We performed Fisher's exact test to test if the odds of finding the following type of candidate variants in the patients is significantly higher than in the controls: (1) Hom LOF variants (vs. Het LOF variants); (2) Hom LOF variants in RBPs (vs. Hom LOF variants in nonRBPs); (3) Het LOF variants in RBPs (vs. Het LOF variants in nonRBPs); (4) Hom DNS variants (vs. Het DNS variants); (5) Hom DNS variants in RBPs (vs. Hom DNS variants in nonRBPs); (6) Het DNS variants in RBPs (vs. Het DNS variants in nonRBPs); (7) all DNS variants in domains of RBPs (vs. in nondomain regions of RBPs); (8) all DNS variants in CC regions of RBPs (vs. in nonCC regions of RBPs); (9) all DNS variants in CC-ER regions of RBPs (vs. in CC-nonER regions of RBPs). At the gene level ((1)-(6)), we selected DNS variants predicted “deleterious” by any of the five algorithms aforementioned. At the subelement level ((7)-(9)), since Hom and Het variants were combined, we applied a more stringent criterion to select DNS variants, which should be predicted “deleterious” by at least three of the five algorithms.

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SUPPLEMENTARY MATERIALS

www.sciencemag.org/science.adj8172
Materials and Methods
Supplementary Text
Figs. S1 to S23
Tables S1 to S16
References (117–160)

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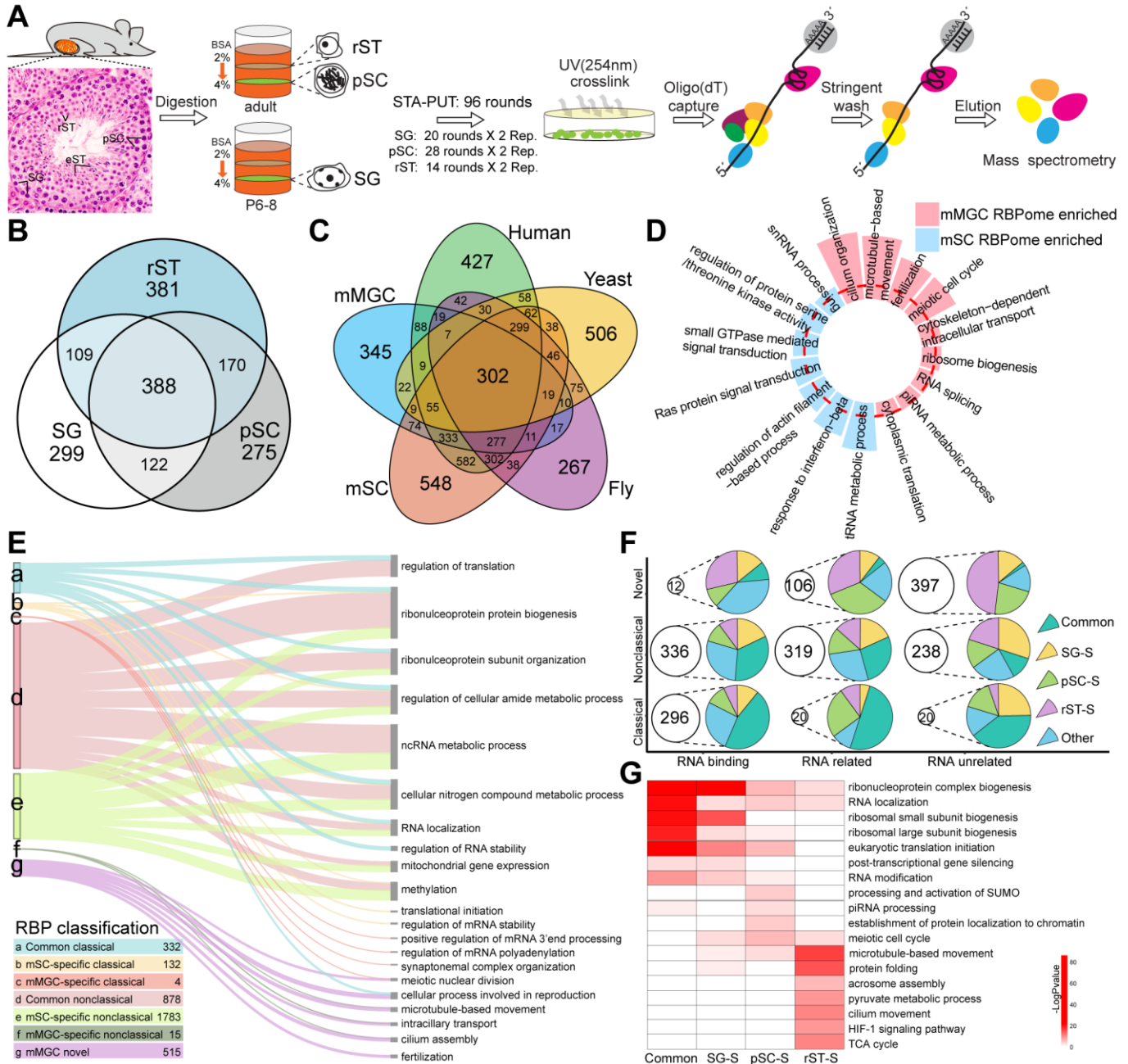


Fig. 1. RIC-based identification and characterization of mMGC RBPs. (A) Schematic of RIC for three types of germ cells (SG, pSC, and rST) purified from fresh mouse testes. SG: spermatogonia; pSC: pachytene spermatocyte; rST: round spermatid. STA-PUT yielded specific populations of SG with >85% purity, rST with >90% purity, and pSC with >85% purity. For each cell type, numbers of RIC replicates (Rep.) and STA-PUT rounds per replicate are indicated. (B) Venn diagram shows intersection of the three subsets of mMGC RBPs. (C) Venn diagram shows intersection of RBPomes from different species and origins. Proteins without mouse homologs are not included. mMGC (mouse male germ cell): RBPome in this study; mSC (mouse somatic cell), human, yeast, and fly: other reported RBPomes. (D) Circular bar plot shows the enriched GO term in mMGC RBPome and mSC RBPome. The bar heights equal to the $-\text{Log}_{10}(P \text{ value})$ of each GO term. Red dashed ring denotes $-\text{Log}_{10}(P \text{ value} = 0.05)$. (E) Sankey plot shows the enriched biological processes GO terms of each RBP subset when intersecting mMGC RBPome with mSC RBPome. Inset table: RBP subsets on the basis of comparison between mMGC and mSC. Common: RBPs shared by mMGC and mSC. Numbers of proteins in each subset are indicated. (F) Balloon plot displays the protein categories of mMGC RBPome based on functional annotations (GO database) and RBP types. Protein number of each category is indicated; each category consists of five subsets. SG-S: RBPs specifically captured in SG; pSC-S: RBPs specifically captured in pSC; rST-S: RBPs specifically captured in rST; Common: RBPs shared by SG, pSC, and rST; Other: RBPs shared by two types of germ cells. (G) Heatmap shows the enriched GO and KEGG terms across four RBP subsets generated from panel F.

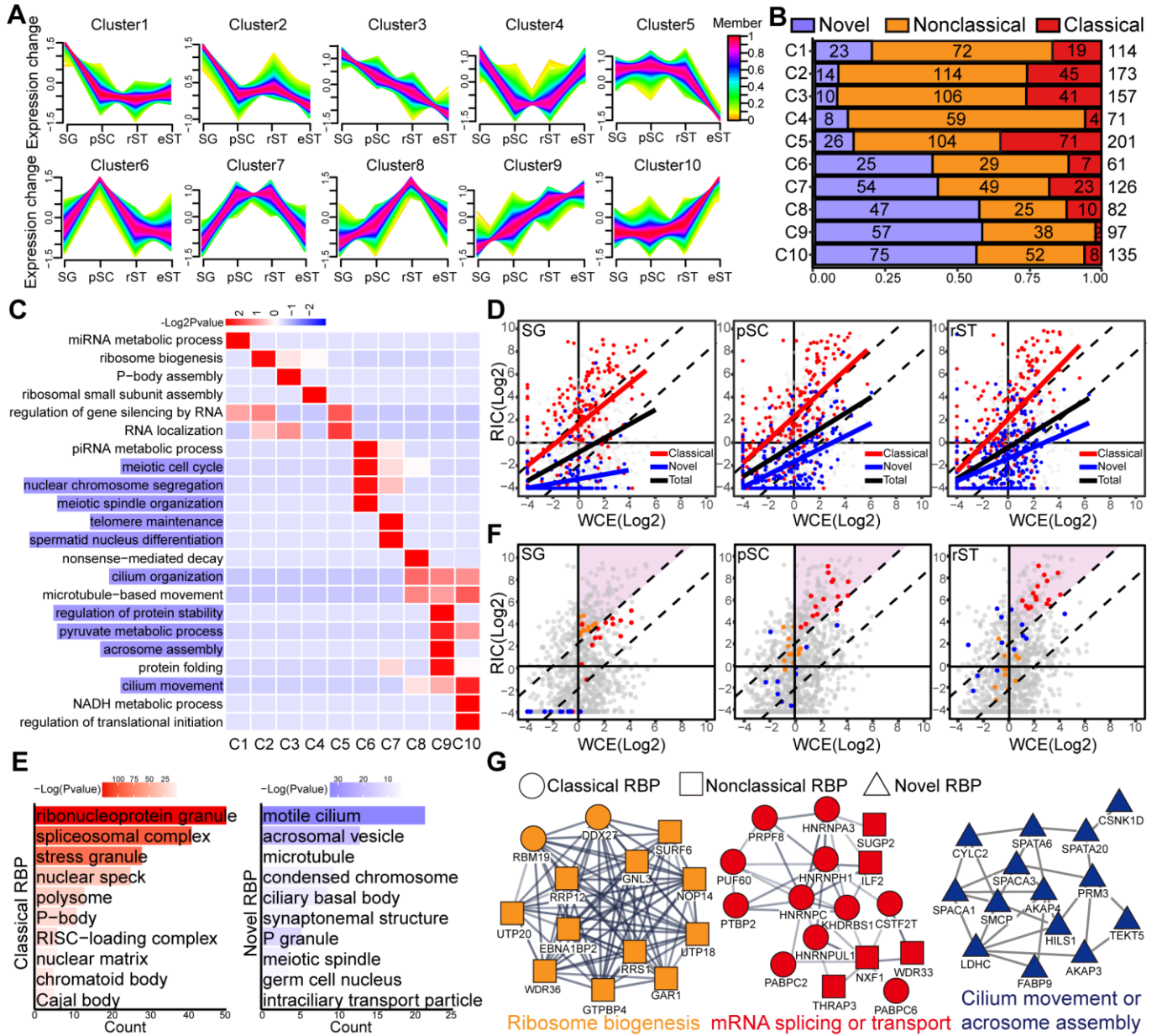


Fig. 2. Multi-hierarchical temporal dynamics of mMGC RBPs. (A) Ten distinct temporal patterns of protein abundance along spermatogenesis. Membership values are color-encoded with red shades denoting high membership values of proteins, whose temporal abundances are highly correlated with the cluster core, and green shades denoting low membership values of proteins. (B) The proportions and counts of each kind of RBP in each cluster are shown. (C) Heatmap shows the significantly enriched GO terms among ten distinct temporal RBP clusters. GO terms with blue background harbor more than half of novel RBPs. (D) Scatter plots show dynamics of relative RNA-binding activity of RBPs measured by $\text{Log}_2(\text{RIC}/\text{WCE})$. Red and blue dots denote classical RBPs and novel RBPs, respectively. Red, blue and black lines show the linear relationship between RIC and WCE to model the relative RNA-binding activity for classical RBPs, novel RBPs, and total RBPs, respectively. (E) Bar plot shows the subcellular localization of classical RBPs and novel RBPs generated by GO cell component (CC) analysis. Top ten GO CC terms based on gene counts are shown. (F) Scatter plots show dynamics of relative RNA-binding activity of RBPs measured by $\text{Log}_2(\text{RIC}/\text{WCE})$. Yellow, red, and blue dots illustrate three representative cases of potentially coordinated RBPs with different extents of inter-stage dynamics. The pink triangle regions define the Strong RBPs with $\text{Log}_2(\text{RIC}/\text{WCE}) \geq 2$ and $\text{Log}_2(\text{WCE}) > 0$ at given stages. (G) Network plots show protein-protein interaction (PPI) clusters of RBPs with colors corresponding to the scatter plots from panel F. Classical, nonclassical, and novel RBPs are shown as circles, squares, and triangles, respectively.

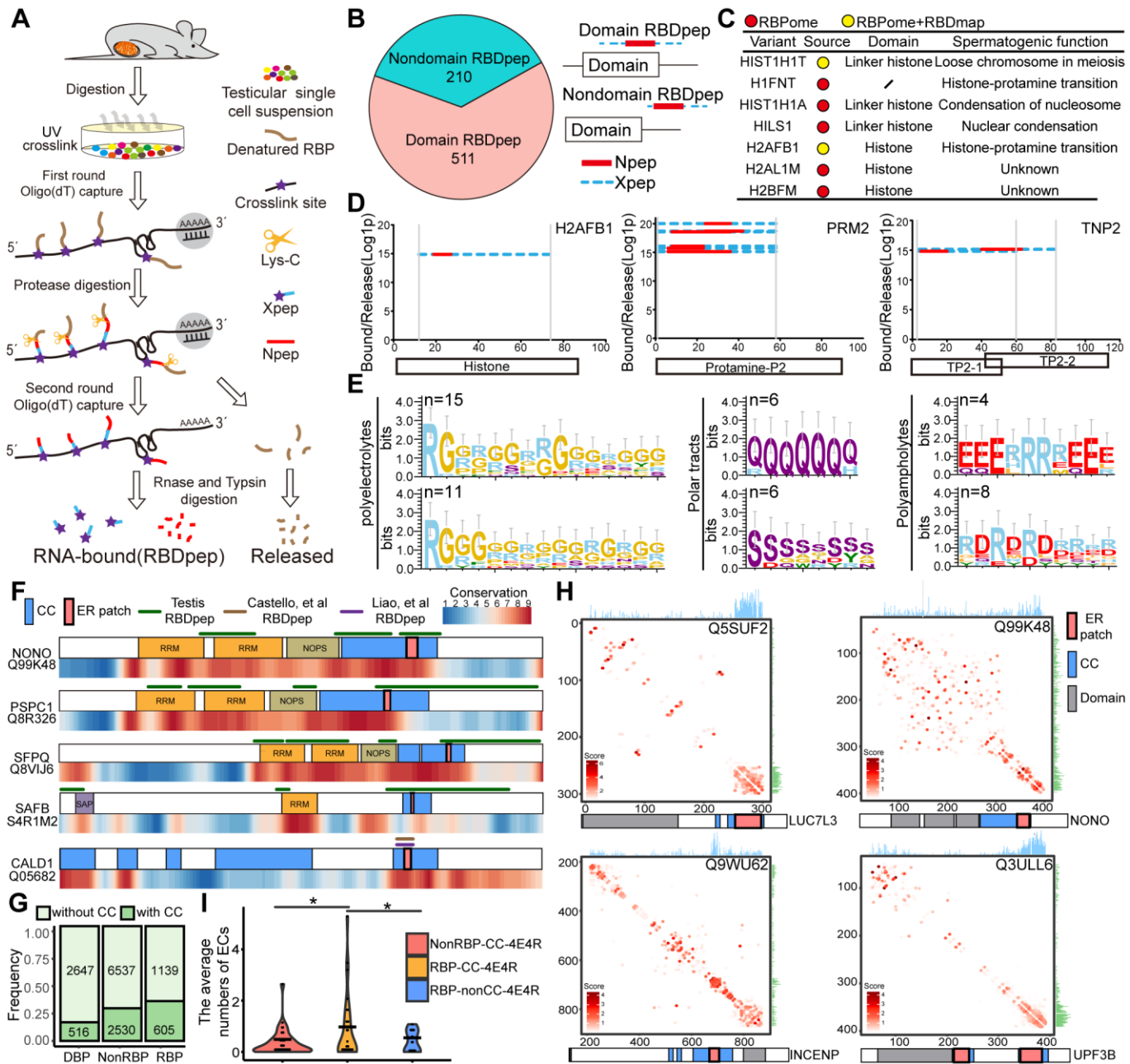


Fig. 3. RBDmap-based discovery of RBDs and RNA-binding nondomain elements. (A) Schematic of RBDmap using fresh mouse testicular cells. Trypsin digestion of RNA-bound peptides generated two kinds of sub-peptides: Npep and Xpep, which combined together are termed RBDpep. (B) Pie plot shows the proportion of domain and nondomain RBDpep in the testis RBDmap dataset. Xpep (blue dashed line) and Npep (red line) constitute RBDpep. Domain RBDpeps are defined overlapping with Pfam domain. Nondomain RBDpeps are defined as no amino acid overlapping with any domain. (C) List shows the testis-specific histone variants identified in mMGC RBPome and/or testis RBDmap dataset. (D) Graphs show all RBDpeps mapping to domain regions of H2AFB1, PRM2, and TNP2 proteins. X axis shows RBDpeps (linear sequences) mapped to Pfam domains at relative positions of RBPs; Y axis shows the enrichment (Log1p) of RNA bound over released fraction for RBDpeps. Blue dashed line: Xpep; red line: Npep. (E) Sequence motifs within nondomain RBDpeps. Multiple sequence alignment of short patches within nondomain RBDpeps used Clustal Omega tool. The sequence logos (motif) were extracted from aligned nondomain fragments. All the motifs are classified to three archetypes based on the physiochemical characteristics of amino acids: polyelectrolytes, polar tracts, and polyampholytes. (F) Schematic representation of RBDpeps mapping to domain and nondomain areas of five ER-containing RBPs. All these ER patches identified by RBDmap are conserved and within coiled-coil (CC) motifs. The locations of our testis RBDpeps (green lines) and other somatic RBDpeps (brown and purple lines) are indicated. CCs and ER patches are shown by blue and pink rectangles, respectively. Conservation score and domain constitution are shown in the bottom color bars. Heatmap shows the evolutionary conservation scores of each residue on proteins from consurf software. The color of the heatmap indicates high (dark red) or low (blue) evolutionary conservation scores of each residue. (G) Bar plot shows the percentage of proteins with at least one CC motif in total proteins classified by its nucleic acid-binding characteristics. Protein list of DBP was generated from cisBP database. RBP: mMGC RBPs; nonRBP: proteins identified by TMT-MS but not by mMGC RIC. (H) Evcoupling's contact maps of several RBPs with co-evolved CC-ER patches. The coevolution scores are color-encoded with a continuous palette of red color. The bar plot on the top X axis (blue) displays the sums of the coevolution scores of each residue with the rest within each RBP, and the bar plot on the right Y axis (green) displays the number of strongly evolutionarily coupled residue pairs (ECs) each residue forms with the rest. (I) Box plot shows the average numbers of ECs within 4E4R patches in different protein groups. * $P < 0.05$, by t test.

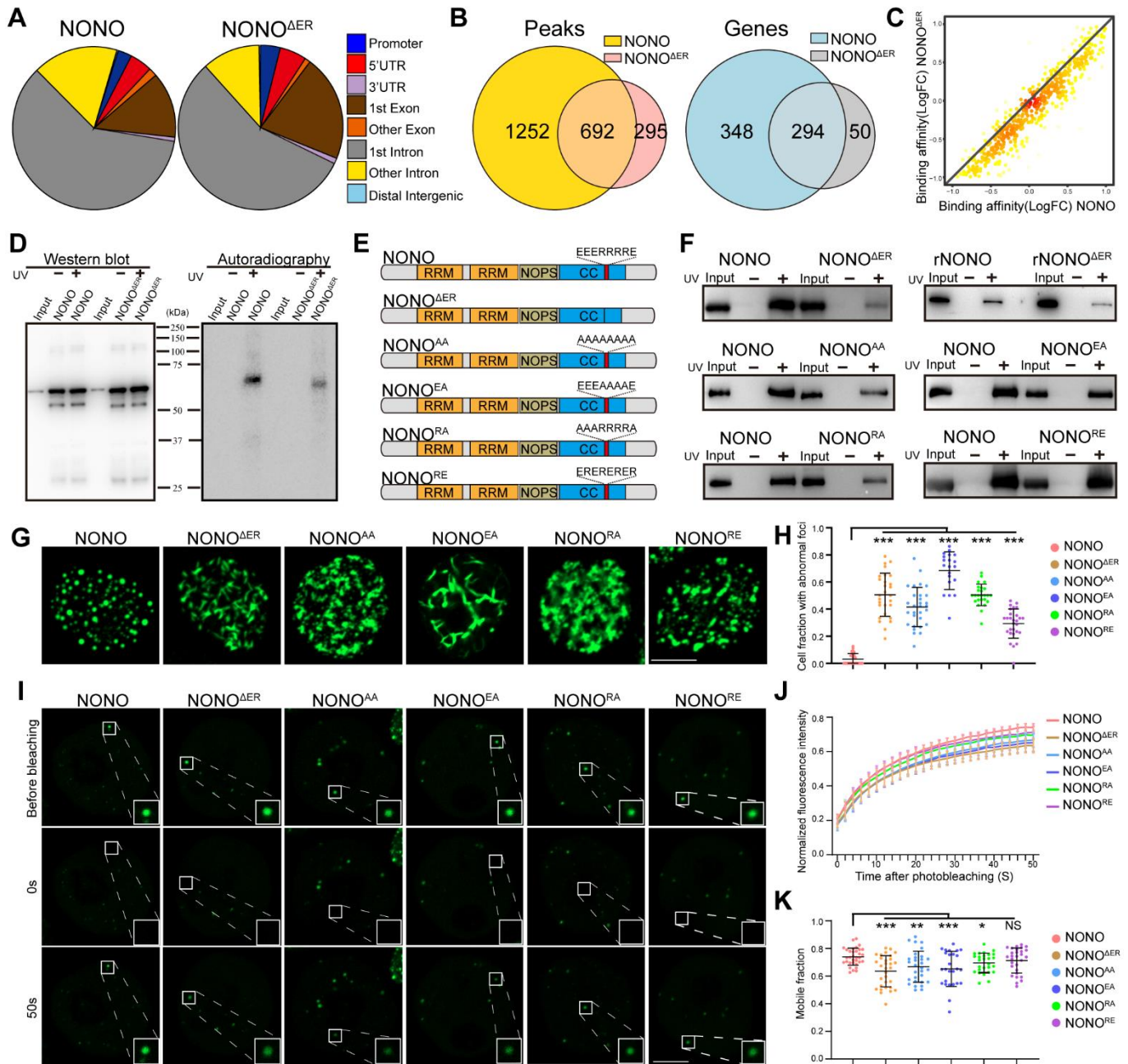


Fig. 4. The NONO ER patch functions in enhancing RNA-binding activity. (A) Pie plots show the genomic distribution of all reproducible eCLIP-seq peaks for HA-NONO-bound RNA and HA-NONO^{ΔER}-bound RNA in 3T3-L1 cells. (B) Venn diagrams show the overlaps of enriched eCLIP-seq binding peaks (left) and target genes (right) between NONO and NONO^{ΔER} in 3T3-L1 cells. (C) Scatter plot compares the relative intensity of each target gene in NONO and NONO^{ΔER} eCLIP-seq in 3T3-L1 cells. Each point shows Log fold change between IP and input in NONO^{ΔER} (Y axis) and NONO (X axis) of one gene. (D) The RNA-binding capacity of NONO and NONO^{ΔER} protein was measured by radioactive CLIP assays. NONO- and NONO^{ΔER}-bound RNAs were radiolabeled by ³²P. Noncrosslinked samples served as negative controls. (E) Schematic diagram of Pfam domains and nondomain areas of wild-type and mutant NONO proteins. ER patches in the CC of NONO protein are shown as red rectangles. (F) Quantitative 2C (q-2C) assays compare the RNA-binding capacity of wild-type and mutant NONO proteins. Total RNPs from 293T cells transfected with HA-NONO, HA-NONO^{ΔER}, HA-NONO^{RA}, HA-NONO^{EA}, HA-NONO^{AA}, and HA-NONO^{RE} were isolated by silica-based matrices after UV crosslink (+) or not (-). Total extracts (1% of input) and purified mRNPs were detected by Western blot with anti-HA antibody. In vitro q-2C assays also compare the RNA-binding capacity of wild-type and mutant NONO proteins. Purified recombinant NONO (rNONO) and NONO^{ΔER} (rNONO^{ΔER}) proteins were UV-crosslinked (+) or not (-) with total RNA extracted from 293T cells. Protein-RNA complexes were isolated by silica-based matrices. 1% of input proteins and purified RNPs were detected by Western blot with anti-NONO antibody. Each experiment was repeated three times and quantitative measurement of each assay is shown in fig. S13D. (G and H) Representative images of foci morphology of overexpressed wild-type and mutant NONO proteins in 293T cells (G). The percentage of the cells with fiber-like foci was shown (H). Only cells with dozens of bright foci, either orbicular or fiber-like, reflective of high expression levels of NONO, were considered. ****P* < 0.001, by *t* test. Scale bar: 10 μm. (I to K) FRAP experiments in 293T cells. Only small foci, reflective of low expression levels of NONO, were picked to do FRAP. (I) Representative images show fluorescence recovery of wild-type and mutant NONO droplets before and after photobleaching a spot of predefined size within the nuclear region outlined in rectangle. (J) The FRAP fluorescence recovery curves record the normalized fluorescence intensity of wild-type and mutant NONO droplets 0-50 s after photobleaching. Mean with 95% CI is shown. (K) Quantification of the mobile fraction of wild-type and mutant NONO droplets at 50 s after photobleaching. Each droplet was picked from one cell. *n* = 33 for NONO, *n* = 31 for NONO^{ΔER}, *n* = 30 for NONO^{AA}, *n* = 26 for NONO^{EA}, *n* = 27 for NONO^{RA}, *n* = 29 for NONO^{RE}; mean ± SD is shown; NS: *P* > 0.05, **P* < 0.05, ***P* < 0.01, ****P* < 0.001, by *t* test. Scale bar: 10 μm.

Fig. 5. The NONO ER patch mediates cell-type-dependent RNA binding and regulation in mitosis-to-meiosis transition. (A) Schematic diagram shows the generation of mice harboring a *Nono*^{ΔER} X-linked allele using CRISPR/Cas9 system. Blue sequences: gRNA; red sequences: PAM. (B) RT-PCR (left) and Western blot (right) validation of NONO expression in *Nono*^{ΔER/Y} testis. (C) Bar plots compare the body weight, testis weight, sperm count, motile sperm, and apoptotic cells between *Nono*^{+Y} and *Nono*^{ΔER/Y} mice. (D) Co-immunofluorescence of NONO with the spermatogonia-specific protein LIN28A in 2-week-old testis sections (top) and with the Sertoli cell marker SOX9 in adult testis sections (bottom). White arrowheads point to NONO-positive spermatogonia and Sertoli cells, respectively. Scale bar: 20 μm. (E) Immunofluorescence analysis of LIN28A-positive cells in 2-week-old *Nono*^{ΔER/Y} versus *Nono*^{+Y} testis sections (top) and SOX9-positive Sertoli cells in adult *Nono*^{ΔER/Y} versus *Nono*^{+Y} testis sections (bottom). Scale bar: 20 μm. ***P* < 0.01, ****P* < 0.001, by *t* test. (F) The RNA-binding capacity of NONO protein in *Nono*^{+Y} and *Nono*^{ΔER/Y} mouse testis was measured by q-2C assays. Each band was measured by ImageJ software. All the intensities of 2C bands are relative to their input. **P* < 0.05, by *t* test. (G) Venn diagrams show the overlap of NONO binding peaks (left) and its target genes (right) between *Nono*^{+Y} and *Nono*^{ΔER/Y} testes. (H) Scatter plots compare the relative intensity of each target gene in NONO and NONO^{ΔER} eCLIP-seq in the testis. Each point shows Log fold change between IP and input in NONO^{ΔER} (Y axis) and NONO (X axis) of one gene. (I) UMAP shows cell type clustering of 46495 single cells from the adult *Nono*^{+Y} and *Nono*^{ΔER/Y} mouse testes, with 3 replicates for each genotype shown in fig. S18D. The 17 cell types (indicated at the bottom) were assigned based on their marker gene expression in fig. S18E. SG: spermatogonia; SC: spermatocyte; ST: spermatid. (J) Top: Heat map shows the relative expression of NONO CLIP target genes (n = 939) in *Nono*^{+Y} testicular cells. Cells were clustered based on panel I. Sertoli: Sertoli cells; mitosis: mitotic cells including type A spermatogonia, type B spermatogonia, and preleptotene spermatocytes; meiosis: meiotic cells including leptotene, zygotene, pachytene, and diplotene spermatocytes; rST: round spermatids; eST: elongated spermatids. Bottom: Density plots show distribution of the expression changes of different gene sets in *Nono*^{+Y} and *Nono*^{ΔER/Y} testicular cells. X axis shows the Log fold change of *Nono*^{ΔER/Y} versus *Nono*^{+Y}. Green line represents NONO CLIP target genes (n = 939), orange line represents total genes (n = 25771). (K) Scatter plots show the average expression of mitosis-enriched genes (red) and meiosis-enriched genes (blue) in mitotic cells and meiotic cells in *Nono*^{+Y} and *Nono*^{ΔER/Y} mice.

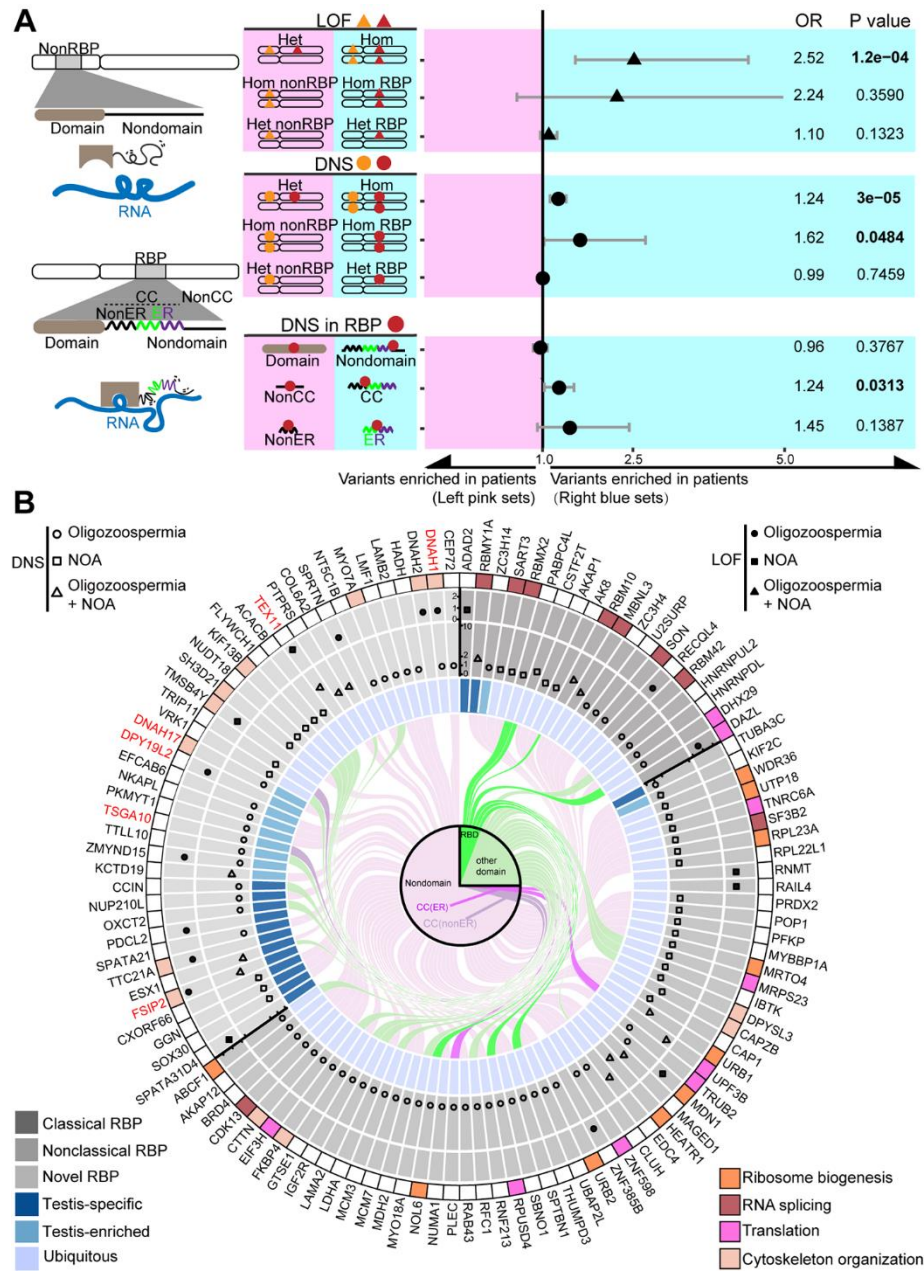


Fig. 6. The genomic architecture of human infertility with a focus on RBPs. (A) Forest plot shows odd ratios for different types of variants in patients with male infertility. Left diagrams demonstrate the various elements of RBP and nonRBP. Middle diagrams show the opposing sides, denoted by left pink sets and right blue sets respectively, of each test. Right numbers displays the actual values of odd ratios and *P* values of Fisher's exact test. LOF variants are marked by triangles; DNS variants are marked by circles. Variants in RBP genes are signed with red and variants in nonRBP genes are signed with yellow. Hom: homozygote; Het: heterozygote. OR: odd ratio; the bold *P* values: significant at $P < 0.05$. (B) Circos plot lists 137 candidate Hom DNS variants in 105 RBP genes and 22 candidate Hom LOF variants in 18 RBP genes. From the inside out: (1) the pie chart clusters variants according to their locations in RBD (dark green lines), other domain (light green lines), 3E3R patch of CC (dark purple lines), non-3E3R of CC (middle purple lines), and nonCC nondomain region (light purple lines); (2) the sankey diagram designates the corresponding RBP of each variant; (3) the blue-color-encoded circle shows the tissue specificity for each RBP: testis-specific (dark blue); testis-enriched (middle blue), and ubiquitous (light blue); (4) the gray-color-encoded circle classifies RBPs into classical (dark gray), nonclassical (middle gray), and novel (light gray). Different shades in this circle indicate the sources of variants for given RBPs found in: oligozoospermia infertile men only (dot), NOA infertile men only (square), and both sets of infertile men (triangle). The radial height denotes the time of recurrence of variants in patients. DNS (hollow) and LOF (solid) variants are represented separately; (5) the outmost circle shows the four selected enriched GO terms among RBPs with candidate variants. According to the IMIGC database, RBPs having been reported for human infertility are highlighted in red.