Original research

Disrupted spermatogenesis in a metabolic syndrome model: the role of vitamin A metabolism in the gut–testis axis

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ABSTRACT

Objective Effects of the diet-induced gut microbiota dysbiosis reach far beyond the gut. We aim to uncover the direct evidence involving the gut–testis axis in the aetiology of impaired spermatogenesis.

Design An excessive-energy diet-induced metabolic syndrome (MetS) sheep model was established. The testicular samples, host metabolomes and gut microbiome were analysed. Faecal microbiota transplantation (FMT) confirmed the linkage between gut microbiota and spermatogenesis.

Results We demonstrated that the number of arrested spermatogonia was markedly elevated by using 10× single-cell RNA-seq in the MetS model. Furthermore, through using metabolomics profiling and 16S rDNA-seq, we discovered that the absorption of vitamin A in the gut was abolished due to a notable reduction of bile acid levels, which was significantly associated with reduced abundance of Ruminococcaceae_NK4A214_group. Notably, the abnormal metabolic effects of vitamin A were transferable to the testicular cells through the circulating blood, which contributed to abnormal spermatogenesis, as confirmed by FMT.

Conclusion These findings define a starting point for linking the testicular function and regulation of gut microbiota via host metabolomes and will be of potential value for the treatment of male infertility in MetS.

INTRODUCTION

Mounting evidence now suggests that excessive energy intake increases the risk factors for metabolic syndrome (MetS), which is seriously harmful to well-being and has become a major health issue worldwide; alarmingly, it is showing a younger trend, and its increasing prevalence is concomitant with the increased incidence of male infertility.1

Several studies demonstrate that the overconsumption of excessive energy alters male reproductive function and consequently impairs spermatogenesis.2–6 Gut microbiota dysbiosis was observed in excessive energy diet-induced MetS.7,8 Recently, Ding et al9 showed that high-fat diet induced gut microbial dysbiosis with changeable phenotypes induced through faecal microbiome transplantation (FMT) leads to male infertility, confirming a causal relationship between the gut microbiota and spermatogenesis. In another study, Zhang et al10 found that alginite oligosaccharide dosed mice showed an improvement in spermatogenesis following FMT, suggesting the possibility that FMT could potentially be used to treat patients in clinical research. These effects may be mediated through the metabolome, because the gut is ideally situated to sense an

Significance of this study

What is already known on this subject?

► Increasing evidence is showing that an excessive-energy diet-induced obesity is one of the risk factors for metabolic syndrome (MetS).

► Gut microbiota plays a pivotal role in metabolic disorders.

► Recent studies show that gut microbiota can have an impact on spermatogenesis.

What are the new findings?

► A MetS model induced by diets with excessive-energy levels resulted in a notable elevation in the number of abnormal testicular tubules and a sharp decline in spermatogonia differentiation.

► The vitamin A absorption was disrupted, which mainly due to the dramatically reduced abundance of bile acids induced by an imbalanced gut microbiota in the MetS model.

► The abnormal metabolic effects of vitamin A were transferable to the testicular cells through circulating blood, which contributed to the damaged spermatogenesis.

► The abnormal vitamin A metabolism contributed to damaged spermatogenesis in a gut microbiota dependent manner by using faecal microbiota transplantation.

How might it impact on clinical practice in the foreseeable future?

► These findings represent the beginning for linking the testicular function and regulation of gut microbiota via host metabolomes.

► Vitamin A metabolism might be applied to be a potential indicator for male infertility in individuals with MetS.

► The regulation of vitamin A metabolism through restoring the gut microbial ecosystem will be of value for male infertility therapy in individuals with MetS.
RESULTS
Metabolic disorders induced by diets with excessive energy levels resulted in impaired spermatogenesis

To study the influence of diets with excessive energy levels on spermatogenesis, we established an excessive energy diet-fed sheep model (online supplemental tables 1 and 2). As expected, the medium-energy diet (MED) and high-energy diet (HED) groups had significantly increased body weight compared with the normal diet (ND) (standard chow diet) group (online supplemental figure 1A). The MED and HED groups also became glucose intolerant and developed insulin resistance, whereas the ND group did not (online supplemental figure 1B). Meanwhile, the HOMA-IR, QUICKI and Matsuda indexes were dramatically altered in the MED and HED groups (online supplemental figure 1C). Furthermore, fasting blood glucose and insulin concentrations were significantly increased in the HED group (online supplemental figure 1D). Serum low-density lipoprotein cholesterol and total cholesterol (TC) were largely increased in the MED and HED groups, while a significant reduction in high-density lipoprotein cholesterol was observed in the HED group (online supplemental figure 1E). These findings suggested that MetS sheep model was induced by diets containing excessive energy in the MED and HED groups.

Then, we investigated spermatogenesis by analysing H&E stained tissue samples and noted the presence of multinucleated giant cells and disorganised spermatogenic cells in the testes of the MED group (figure 1A). The impaired spermatogenesis became more severe in the HED group, where the number of spermatogenic cells was significantly reduced, and the spermatogenic epithelium became thinner (figure 1A). The percentage of histomorphologically abnormal tubules was significantly increased in the MED and HED groups compared with the ND group (figure 1B). Furthermore, H&E staining of cauda epididymis showed numerous mature sperm in ND group, but several cell debris was observed in the MED and HED groups (figure 1C). The epididymal spermatozoa concentration was lower in the MED and HED groups compared with that in the ND group (figure 1D). The spermatozoa morphology showed greater levels of abnormality in the MED and HED groups compared with the ND group (online supplemental figure 2). Cumulatively, our results suggest that MetS sheep model, induced by diets containing excessive energy, impaired spermatogenesis.

Deep analysis of 10x scRNA sequencing revealed that impaired spermatogenesis was due to the blocked differentiation of spermatogonia (SPG)

To define which spermatogenic cell subtypes were engaged in biological pathways or functions during impaired spermatogenesis in excessive energy diet-induced MetS sheep model, 10x Genomics analysis of testicular cells was performed (online supplemental figure 3A). The ratios of cell viability and aggregation are shown in online supplemental table 3. Filtered data were analysed together from the ND, MED and HED groups by using uniform manifold approximation and projection analysis to identify cell subtypes. Corresponding to known gene markers in different cell types in the testis, we identified nine clusters, including SPG, early spermatocytes (SPCs), late SPC, early round spermatids (Std), mid round Std, late round Std, Sertoli cells, Leydig cells and undefined cells (online supplemental figure 3B). The gene markers for nine clusters were shown in online supplemental figure 3C,D.

To elucidate the molecular nature of germ cell differentiation in impaired spermatogenesis induced by metabolic disorders, we calculated the percentage of each cell subtype, using only the germ cells from the ND, MED and HED group. As shown in figure 1E,F, significant difference was revealed in the SPG cluster (ND: 3.8%, MED: 6.58% and HED: 14.28%), early SPC cluster (ND: 7.86%, MED: 17.60% and HED: 22.99%) and late round Std cluster (ND: 41.00%, MED: 16.12% and HED: 14.36%) among the three treatment groups. Corresponding to the H&E staining results, spermatogenic cells were dramatically lower in the MED and HED groups, which indicated that SPG differentiation was disrupted. Hence, we set out to perform clustering analysis on SPGs (online supplemental figure 4A-E). This revealed four distinct cell clusters, which we defined as: (1) undifferentiated SPG (undiffed SPG), (2) differentiating SPG (diffing SPG), (3) differentiated SPG (diffed SPG), and (4) pre-luteotene SPC (pre-L-SPC), based on several gene markers as described previously.13–15 To assess the developmental trajectory of these four SPG subsets, we performed monolce pseudotime trajectory analysis, which aligns individual cells along a developmental trajectory. It was revealed that these four SPG subsets had the following developmental order: undiffed SPG → diffing SPG → diffed SPG → pre-L-SPC (figure 1G,H). Meanwhile, the existence of branched point from the linear path of the pseudotime trajectory was observed, demonstrating that one branched group of cells represented the differentiation from undiffed SPG to diffed SPG and pre-L-SPC; the other branched group of cells represented the transition state that bridged the undiffed SPG and diffing SPG stages, which were likely to be SSC pool enriched, simultaneously generating progenitor SPG that initiated spermatogenic differentiation (figure 1I).

In order to explore the mechanism of SPG differentiation retardation in MetS sheep model, Gene Ontology (GO) analysis was performed to demonstrate that distinct categories of differentially expressed genes were enriched in the ND versus MED group and ND versus HED group (figure 1J). ‘Reproductive process’ and ‘metabolic process’ were highly enriched in GO terms of SPG clusters in ND versus MED and ND versus HED, which suggested that metabolic process was consistent with reproductive process and was significantly altered during SPG differentiation.

The absorption of vitamin A was disrupted due to a drastic reduction in levels of bile acids in the MetS model

To address whether the arrest of SPG differentiation was closely connected with gut metabolism, we examined the untargeted metabolome profiles generated on intestinal digesta samples using ultra-high performance liquid chromatography–quadrupole time-of-flight mass spectrometry. Using partial least squares discriminant analysis, we observed distinct clustering of metabolites among the ND, MED and HED groups (figure 2A and online supplemental figure 5A). The excessive energy diet was...
Figure 1  High-caloric load induced by excessive energy intake disrupted spermatogenesis in a metabolic syndrome (MetS) model. (A) The representative testicular sections are stained with H&E. Scale bar=100 μm. (B) The percentage of abnormal tubules in testicular samples from each group (n=5). (C) The representative epididymal sections are stained with H&E. (D) Statistical analysis of sperm counts (n=5). (E) UMAP plot of germ cell clusters in testes defined by scRNA-seq analysis. (F) The percentage of each germ cell cluster from three groups. (G) Single-cell trajectories of SPG subsets are shown with cells ordered in pseudotime. Branch point represents distinct germ cell transition states. (H) The SPG trajectory is colored by three cell states. (I) Heatmap represents significant DEGs between each cluster at differentiated SPG and SSC pool stages at branch point. (J) Gene Ontology terms of differentially expressed genes (DEGs) in the SPG cluster from (a) ND versus MED and (b) ND versus HED groups. HED, high-energy diet; MED, medium-energy diet; ND, normal diet; scRNA-seq, single-cell RNA-seq; SPG, spermatocytes; SPG, spermatogonia; SSC, spermatogonia stem cell; UMAP, uniform manifold approximation and projection.
Gut microbiota sufficient to trigger widespread changes in metabolites (online supplemental figure 5B, C). Interestingly, these two comparisons had 250 common differential metabolites altered, and they were mainly enriched in the vitamin digestion and absorption KEGG pathway (figure 2B). The differential metabolites enriched in the vitamin digestion and absorption pathway were comprised of three fat-soluble vitamins and five water-soluble vitamins (figure 2C). Owing to the bile acid associated pathway also being enriched, we hypothesised that the absorption of fat-soluble vitamins was affected by bile acid levels. There are three fat-soluble vitamins: retinol, retinene and alpha-Tocopherol (vitamin E). Retinol and retinene are the bioactive forms of vitamin A, and they can be transformed into each other in vivo and have the same function. Retinene may be further oxidised to retinic acid, but this is an irreversible process. Vitamin E is an antioxidant that promotes vitamin A absorption. We found that the relative abundance of vitamin E was decreased in the MED and HED groups compared with the ND group (figure 2Dd). Taken together, these untargeted metabolome profiles demonstrated that the absorption of vitamin A was dysregulated by reduced bile acid levels in the MetS model.

Gut microbiota dysbiosis was induced in the MetS model

The gut microbiota plays a vital role in its hosts metabolism by interacting with the host’s signalling pathways; therefore, we examined bacterial 16S rDNA sequencing of intestinal digesta. Rarefaction curve examination revealed that the data were reliable for further analysis (online supplemental figure 7A). A distinct clustering of microbiota composition for the three treatment groups was observed using Bray-Curtis based

Figure 2  Absorption of vitamin A was disturbed in a MetS model. (A) Orthogonal projections to latent structures-discriminant analysis (PLS-DA) score plot for discriminating the intestine digesta metabolome from the ND, MED and HED groups (n=5). (B) Disturbed metabolic pathways in the ND versus MED and ND versus HED groups. (C) Heatmaps of the differential metabolites including eight vitamins that were altered through MED and HED feeding compared with ND feeding. (D) Comparison of the relative abundance of (a) vitamin E, (b) retinol, (c) retinene, (d) cholic acid, (e) glycocholic acid, (f) lithocholic acid and (g) tauroursodeoxycholic acid in the indicated groups. HED, high-energy diet; MED, medium-energy diet.
Gut microbiota

Figure 3  Profiling of the gut microbiota was altered in a MetS model. (A) Bray-Curtis principal coordinate analysis (PCoA) plot of gut microbiota based on the operational taxonomic unit metrics of the samples in the ND, MED and HED groups (n=9). (B) Bacterial taxonomic profiling at the phylum level of intestinal bacteria from different treated groups. (C) (a) Shannon index and (b) Simpson index of gut microbiota in three treated groups. (D) Heatmap of differential abundance from the MED and HED groups compared with the ND group. The color of the spots in the left panel represents the relative abundance at the genus level in each group. HED, high-energy diet; MED, medium-energy diet; MetS, metabolic syndrome.

principal coordinates analysis (figure 3A). To assess the overall composition of the bacterial community in different groups, the degree of bacterial taxonomic similarity of gut microbiota was analysed at the phylum level (figure 3B). Moreover, Shannon curves, OTU rank curves, Shannon index and Simpson index of gut microbiota were significantly altered, indicating that the richness and diversity of the gut microbiota was reduced in the MED and HED groups (figure 3C and online supplemental figure 7A). Furthermore, the twenty-nine bacteria altered at the genus level in the MED and HED groups compared with the ND group (figure 3D). Genus-level analysis showed that most of these altered bacteria belonging to Firmicutes, Ruminococcusaceae and among these genus-level bacteria, the relative abundances of Ruminococcus_2 and Ruminococcusaceae_NK4A214_group were significantly decreased in the MED and HED groups compared with ND in an energy dose-dependent manner after the long-term high-calorie challenge (online supplemental figures 7B and 8B).

The physiological function of bile acids includes the promotion of absorption of fat-soluble vitamin A. It is revealed that CA, LCA and TUDCA were related with retinol, and they were also correlated with retinene by using pearson’s correlation analysis (figure 4 and online supplemental figure 8A). To investigate the functional relationship between gut microbial dysbiosis and bile acid levels, Spearman’s correlation analysis was performed. As shown in figure 4B, Spearman’s correlation analysis between CA, LCA TUDCA and differential genus-level bacteria revealed that the numbered bacteria 4, 9, 15, 17, 19, 21 and 27 were significantly correlated with CA, LCA and TUDCA. Combined with the relative abundance of these bacteria (online supplemental figures 7B and 8B), Ruminococcusaceae_NK4A214_group showed a strong correlation with CA, LCA and TUDCA (figure 4C). Cumulatively, Ruminococcusaceae_NK4A214_group was dramatically downregulated, which in turn had a profound influence on bile acid levels and further affected host vitamin A absorption in the MetS model.

Vitamin A metabolism dysfunction was transferable to host testes in the MetS model

In order to deeply investigate the mechanism of impaired spermatogenesis, we verified vitamin A metabolism in the testis. First, we performed targeted metabolomics profiling of the vitamin A level in testis samples from different treated groups using liquid chromatography–MS. Vitamin A levels were significantly decreased in the HED group compared with the ND group (figure 5A). Furthermore, we detected the vitamin A level in serum using an ELISA kit and found that the data pointed towards a similar trend as that in the testis (figure 5B). Testicular cells receive vitamin A from the blood, and blood serum vitamin A is bound by retinol binding protein 4 (RBP4). Vitamin
Gut microbiota

A can also be transported into testicular cells via RBP4 delivery to membranes. Therefore, we examined the expression levels of RBP4 using immunofluorescent (IF) staining and western blot. Results showed that the expression level of RBP4 was significantly decreased in the MED and HED groups compared with that in the ND group (figure 5C). These results corresponded to the vitamin A level tests in the LS-MS and ELISA experiments.

The 10× scRNA-seq results given previously indicated that differentiation from undiffed SPG to pre-L-SPC was partly blocked. PLZF is expressed in SPG, which is a marker gene of undifferentiated SPG for maintaining self-renewal of SSCs.18 As shown in online supplemental figure 9, the expression level of PLZF was upregulated in the testis by using IF staining and western blot in the MED and HED groups compared with the ND group. In addition, the expression of marker genes of meiosis in pre-L-SPC are detected, and we found that STRA8 and SYCP3 expression levels displayed a sharp decrease, as shown by IF staining (figure 5D) and western blot (figure 5E,F).

In the present study, the damage during spermatogenesis resulted from the downregulated expression of meiotic gene markers, and vitamin A metabolism played a vital role in meiotic gene activation. To further assess whether RDH10 levels were changed in the testis, we identified the expression level of RDH10 using IF staining and western blot and found that it was downregulated in the MED and HED groups compared with the ND group (figure 5C, E,F). Together, these results suggested that vitamin A metabolism was abnormal, thus the reduced levels of vitamin A and RDH10 contributed to the compromised impaired spermatogenesis in testes in the MetS model.

Perturbative vitamin A metabolism led to impaired spermatogenesis in a gut microbiota dependent manner

To investigate the effect of the gut microbiota on impaired spermatogenesis phenotypes in the MetS model, microbiota of the intestinal digesta taken from the ND and HED groups was transplanted into mice by oral gavage (figure 6A). After 8 weeks of colonisation, the body weight (online supplemental figure 10A), serum TC levels (online supplemental figure 10B) and serum TG levels (online supplemental figure 10C) did not show any significant difference between ND-FMT (mice transplanted with microbiota from the ND controls) and HED-FMT (mice received the HED microbiota). However, HED-FMT displayed a significant decrease in testicular vitamin A levels, compared with those mice in the ND-FMT (figure 6B). To further confirm that the differential testis vitamin A levels in the FMT groups were due to the alteration of the colonised gut microbes instead of the vitamin A and bile acids levels transmitted from the donors, we examined the transmitted material from the donors ND and HED groups. The transplanted material was prepared by strict dilute and homogenise filter procedure, and we found that the levels of retinol, retinene and bile acids had no significant difference in the two donor groups (online supplemental figure 11A-C). In addition, we detected the abundance of the

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Figure 4  The alteration of gut microbiota was closely associated with the bile acid levels. (A) Pearson’s correlations of retinol and retinene levels with the levels of (a) CA, (b) LCA and TUDCA. Red indicates a positive correlation, and blue indicates a negative correlation. (B) Spearman’s correlation analysis of differential genus microbiota levels with CA, LCA and TUDCA levels. (C) Spearman’s correlation analysis of Ruminococcaceae_NK4A214_group abundance with (a) CA, (b) LCA and (c) TUDCA levels. CA, cholic acid; LCA, lithocholic acid; TUDCA, tauroursodeoxycholic acid.
Ruminococcaceae_NK4A214_group in the transmitted material from the donors and found that the abundance of the Ruminococcaceae_NK4A214_group was dramatically decreased in the HED group compared with that in the ND group (online supplemental figure 12A). We also analyse the abundance of the Ruminococcaceae_NK4A214_group in the ND-FMT and HED-FMT by PCR and qPCR and found that the abundance of Ruminococcaceae_NK4A214_group was significantly decreased in the HED-FMT group compared with ND group (figure 6C and online supplemental figure 12B).

To confirm that microbiota transplantation modulated spermatogenesis, we conducted H&E staining to examine testicular morphology. This revealed that testicular tissue from mice transplanted with ND microbiota exhibited normal testicular development, whereas tissue from mice transplanted with microbiota from the HED group exhibited thinning of the seminiferous epithelium and detachment of testicular cells from the basement membrane of the seminiferous tubules; in addition, some spermatogenic cells were dead and shed into the luminae of the tubules (figure 6D). The percentage of abnormal tubules was observed between the ND-FMT and HED-FMT; ND-FMT had approximately 12% abnormal tubules, while HED-FMT had more than 47% (figure 6E). The expression levels of STRA8 and RDH10 were determined by IF staining (figure 6F) and western blot (figure 6G,H), and HED-FMT had dramatically decreased protein expression, which was similar to the results of the MetS model. Collectively, these results demonstrated that the recipient mice recapitulated normal testicular morphology and the phenotype of testicular cells in the seminiferous tubules; in addition, some spermatogenic cells were dead and shed into the luminae of the tubules. This finding indicated that, through its ability to restore the homeostasis of gut microbiota, vitamin A metabolism might be a potential candidate for the treatment of male infertility in individuals with MetS (figure 7).

Adequate vitamin A is essential for male and female reproduction as well as embryonic development. Vitamin A is particularly important for its role in the process of meiotic initiation in the female gonad during embryogenesis and in the male gonad postnatally. In the first step, retinol is reversibly converted to retinaldehyde (RA) mainly by the enzyme RDH10. In the second step, RAL can then be oxidised to form RA mediated by members of the aldehyde dehydrogenase 1A family. Several studies report that vitamin A deficient rodents suffer male infertility mainly owing to a deficiency of RA. Therefore, the gut–testis axis in the aetiology of male infertility in MetS is worthy of exploration. First, we conducted a detailed phenotypic assessment of impaired spermatogenesis from a MetS model induced by an excessive energy diet and revealed a notable elevation in the number of abnormal tubules and a sharp decline in SPG differentiation. Furthermore, disruption of vitamin A absorption in the intestine was observed in the MetS model. Notably, the absorption of fat-soluble vitamins was affected by reduced bile acid levels and was strongly correlated with the abundance of Ruminococcaceae_NK4A214_group in the MetS model. These findings suggested that disruption of the absorption of vitamin A was mainly due to the dramatically reduced abundance of these bile acids induced by an imbalance in the homeostasis of gut microbiota. Lastly, the pivotal role of vitamin A metabolism in the gut–testis axis was uncovered in our study. This finding indicated that, through its ability to restore the gut microbial ecosystem, vitamin A metabolism might be a potential candidate for the treatment of male infertility in individuals with MetS.
on the relationship between vitamin A status and MetS. Furthermore, some reports suggested that vitamin A deficiency is a possible independent risk factor for MetS. However, whether an association between MetS and infertility in the vitamin A deficiency has not been uncovered. Organisms work as a whole, so an interference in one organ is likely to impact the body’s homeostatic balance and lead to damage in other related organs. To the best of our knowledge, the current findings are the first to show that the abnormal vitamin A metabolic process was strongly mediated by bile acid dysregulation and gut microbiota dysbiosis, resulting in the impairment of male fertility via circulating blood in the MetS model. Additionally, the expression of RDH10, an enzyme that is essential in the conversion of retinol to RAL, exhibited a sharp reduction, which was mainly due to feedback following the decreased abundance of vitamin A in the testis.

The gut microbiota is regarded as a metabolic organ that regulates its hosts metabolism. Meanwhile, the abundance of bile acids as a signalling molecule in metabolically active tissues such as the gut has aroused great interest in metabolic disorders. Furthermore, it is known that the interaction between the gut microbiome and bile acids is not unidirectional. On the one hand, the gut microbiota can regulate the size of the bile acid pool and the ratio of bile acid composition. On the other hand,
bile acid is one of the main components of bile; a healthy concentration is necessary for the dissolution, digestion and absorption of lipids and fat-soluble vitamins from food. In the present study, using KEGG pathway analysis, we found that the common differential metabolites in intestinal digesta from the MED and HED groups, when compared with the ND group, were also mainly enriched in the bile acid related pathway, which indicated that bile acids have an influence on vitamin A absorption. Additionally, correlation analyses of bile acid levels and genus-level bacteria revealed that CA, LCA and TUDCA were correlated strongly with Ruminococcaceae_NK4A214_group. Our findings suggest that the modulation of gut microbiota altered bile acids levels and further affected vitamin A absorption in the host gut, and the abnormal vitamin A metabolism was transferable to the testis through the blood circulation, resulting in damaged spermatogenesis in the MetS model. We therefore suggest that this study may have promising therapeutic value for male infertility induced by MetS.

Figure 7  Graphical abstract. Diagram illustrating the proposed vitamin A metabolism via the gut–testis axis being indispensable for spermatogenesis. The disrupted vitamin A absorption was affected by reduced bile acid levels, which mainly due to the dramatically decreased abundance of Ruminococcaceae_NK4A214_group. Notably, the abnormal vitamin A metabolism were transferable to the testicular cells through the blood circulation, which led to abnormal spermatogenesis in the MetS model. MetS, metabolic syndrome.
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