Type 2 diabetes (T2D) is a long-term metabolic disorder disease characterized by high blood sugar and relative lack of insulin. Previous studies have demonstrated that *Dendrobium* has potent glucose-lowering effects and may serve as add-ons or alternatives to classic medications for T2D prevention and treatment, but the underlying molecular mechanisms were still unclear. We performed biochemical and transcriptional profiling (RNA sequencing [RNA-seq] and microRNA sequencing [miRNA-seq]) analyses on the pancreas and liver of *Dendrobium fimbriatum* extract (DFE)-fed diabetic rats and control animals. Our sequencing and experimental data indicated that DFE significantly alleviated diabetes symptoms through inhibiting inflammation and preventing islet cell apoptosis in diabetic pancreas. Transcription factors in Stat/nuclear factor κB (NF-κB)/Irf families combined with miR-148a/375/9a served as key regulators in the inflammation and apoptosis pathways under DFE administration. Meanwhile, DFE improved the energy metabolism, lipid transport, and oxidoreductase activity in the liver, and thus decreased lipid accumulation and lipotoxicity-induced hepatocyte apoptosis. Our findings revealed that DFE may serve as a potential therapeutic agent to prevent T2D, and also showed the combination of transcriptome profiling and regulatory network analysis could act as an effective approach for investigating potential molecular mechanisms of traditional Chinese medicine on diseases.

INTRODUCTION

Diabetes mellitus (DM) is a worldwide metabolic disorder disease caused by pancreatic β cell dysfunction and insulin dysbiosis, and mainly characterized by hyperglycemia and dyslipidemia. Most hypoglycemic agents used for diabetes treatment, including synthetic insulin and western medicine, may bring about side effects, such as hypoglycemia and myocardial infarction. Traditional Chinese medicines (TCMs) as health food resources could serve as an effective supplement for diabetes therapy.

TCM has more than 2,000 years of history and has been widely used in clinical studies for diabetes therapy. *Dendrobium*, as a traditional edible and medicinal plant in the Chinese Pharmacopoeia (National Pharmacopoeia Committee, 2015), is widely distributed in the tropical and subtropical regions of Europe/Asia/Oceania. Previous studies have demonstrated that *Dendrobium* has potent glucose-lowering effects and may serve as add-ons or alternatives for the prevention and treatment of diabetes. Dendrobium contains multiple active components, such as polysaccharides, alkaloids, and glycosides, among others, in which, the *Dendrobium* polysaccharides are the main components. Pharmacology research has proved that *Dendrobium* and the polysaccharide extracts possessed hypoglycemic, hepatoprotective, and hypolipidemic effects. *Dendrobium* could effectively reduce the levels of blood glucose, triglyceride, and serum glycosylated protein in hyperglycemic mouse and diabetic rat models. Previous studies demonstrated that the *Dendrobium* polysaccharides could inhibit JNK phosphorylation and promote AKT ser phosphorylation in the islets tissue of diabetic rats. However, rare research had systematically investigated the potential molecular mechanisms underlying the anti-hyperglycemia effects of *D. fimbriatum* on diabetes.

Next generation sequencing (NGS)-based transcriptome profiling could offer more comprehensive views for potential mechanisms underlying molecular mechanisms of traditional Chinese medicine on diseases. This is an open access article under the CC BY license (http://creativecommons.org/licenses/by/4.0/).
involved in diabetes and its complications in diabetic models or human samples.\(^{12,13}\) Furthermore, transcription factor (TF) and microRNA (miRNA) as two major regulators of gene expression at transcriptional and post-transcriptional levels may form a feed-forward loop contributing to the development of diabetes.\(^{14}\) However, few studies were conducted to explore the co-regulation of TFs and miRNAs on diabetic models, and rare studies focused on the molecular mechanisms of how *Dendrobium* showing hypoglycemic effects on diabetes.

In this study, we proved that the extracts of *D. fimbriatum* (DFEs) could increase the level of insulin and alleviate hyperglycemia in diabetic rats. To investigate potential molecular mechanisms of how the DFE regulates blood glucose, we performed transcriptome profiling (RNA sequencing [RNA-seq] and microRNA sequencing [miRNA-seq]) analysis and experimental validation on the pancreas and liver from DFE administration, diabetes, and normal rats. Our data imply that the DFE prevents β cell apoptosis and decreases hepatic lipid accumulation, which may be useful for the prevention and treatment of diabetes and its complications.

### RESULTS

#### DFE Significantly Alleviates Hyperglycemia and Improves Glucose Tolerance in Diabetic Rats

A detailed experimental design was shown in the Figure 1A. Compared with the normal rats, high-fat diet (HFD) and dexamethasone (DEX) administration severely impaired the glucose tolerance capacity and elevated fasting blood glucose (FBG) in the diabetic rats (diabetes group; Figures 1B and 1D). The peak concentrations of blood glucose appeared at 60 mins after oral glucose intake and thereafter returned to basal values (Figure 1C). The area under the curve (AUC) of the blood glucose level (source data of Figure 1C, sampled from the time points of 0–120 mins) in the diabetic rats was significantly larger than others (p < 0.01; Figure 1D). Moreover, a decrease of insulin concentration and an increase of serum free fatty acids (FFAs) level were observed in the diabetic rats (Figures 1E–1G). These results indicated that the diabetic model was successfully constructed.

To evaluate the hypoglycemic effects of DFE on diabetes, two different doses of DFE (100 and 200 mg/kg) were orally administrated, and the metformin (200 mg/kg) was used for positive control (Figure S2). Oral administration of 100 and 200 mg/kg DFE showed a similar hypoglycemic effect on the diabetic rats (Figure S2), hence the dose of 100 mg/kg was selected for the further study, and the rats with oral DFE administration were classified to the diabetes-DFE group in this study. DFE administration significantly decreased FBG and the blood glucose level compared with the diabetic rats (Figures 1B and 1C), which implied that the DFE could improve the glucose tolerance. The AUC values of blood glucose level were markedly smaller with DFE administration compared with the diabetes group (Figure 1D). Meanwhile, the concentrations of serum and pancreatic insulin were both increased with DFE administration (Figures 1E and 1F), whereas the level of serum FFAs was decreased (Figure 1G). Combined with the results above, our data implied that DFE administration
could significantly increase the insulin concentration and lower the blood glucose level in the diabetic rats, which may thereby alleviate hyperglycemia.

Transcriptome Profiling of the Pancreas and Liver for Normal, Diabetes, and Diabetes-DFE Groups

To investigate potential molecular mechanisms underlying DFE alleviating hyperglycemia, we performed transcriptome sequencing (RNA-seq and miRNA-seq) for the pancreas and liver tissues in the three groups (normal, diabetes, and diabetes-DFE). The group description and basic statistics of NGS data were summarized in Tables S1 and S2. In total, we found 13,231 genes and 276 miRNAs expressed in the pancreas (Figures S3A and S3B), while 14,285 genes and 177 miRNAs were detected in the liver samples (Figures S3C and S3D). Meanwhile, several miRNAs, such as the let-7 family members and miR-148/miR-143, were highly expressed in the pancreas and accounted for 70% of the total expression of all miRNAs (Figures S3E–S3G). A set of 12 miRNAs, including miR-122, let-7a/b/c/f, and miR-148, among others, accounted for 90% of total miRNA expressions in the liver samples (Figures S3H and S3I).

Bioinformatics Analysis Demonstrated that DFE Influenced Inflammatory-Related Processes in the Diabetic Pancreas

We detected 1,645 and 1,704 differentially expressed genes (DEGs) in the pancreas for diabetes-VS-normal and diabetes-DFE-VS-diabetes group comparisons, respectively. Among them, 588 DEGs showed the opposite expression pattern in the two comparisons, which indicated that these DEGs may attribute to the glucose-lowering effects of DFE in the pancreas (Figure 2A; Figure S4A). Interestingly, most of the 588 DEGs (97%, 571) were significantly downregulated in the diabetes-DFE-VS-diabetes comparison and upregulated in the diabetes-VS-normal comparison. Moreover, 74% of the 571 DEGs were strongly associated to inflammatory and immune processes (Figure 2B), which implied the main effects of DFE on the diabetic pancreas may focus on the anti-inflammatory. Furthermore, we identified 70 and 59 differentially expressed miRNAs (DEMs) in the diabetes-VS-normal and
diabetes-DFE-VS-diabetes comparisons, respectively (Figures S4B and S4C).

Notably, 87% of DEGs significantly upregulated in the diabetes-VS-normal group comparison were enriched in inflammation and immune-related processes, including inflammatory and immune response, the cytokine and chemokine pathway, JAK-STAT/nuclear factor κB (NF-κB)/NOD-like/Toll-like receptor signaling pathways, and apoptosis pathways (Figure 2C). Interestingly, the significantly downregulated DEGs in the diabetes-DFE-VS-diabetes comparison were mainly involved in three biological categories, which were upregulated in the diabetes-VS-normal comparison: (1) cytokine receptor activity, (2) inflammatory and immune processes, and (3) apoptosis (Figure 2C). Pathway cross-talk analysis demonstrated that the DFE may prevent the apoptosis of islet cells through suppressing the interleukin- (IL-) and interferon-induced inflammation/immune processes, which could activate downstream cell death signaling, including JAK-STAT/NF-κB/TNF pathways and others (Figures 3A and 3B). Additionally, the upregulated DEGs in the diabetes-DFE-VS-diabetes comparison were mainly enriched in the metabolic and insulin signaling pathways (Figure 4B). These results suggested that the DFE may decrease severe inflammation and immune response in the diabetic pancreas, which may thereby inhibit the islet cell apoptosis and increase insulin secretion (Figures 1E and 1F). Meanwhile, 22 upregulated and 11 downregulated DEMs displayed an opposite tendency between the DFE-VS-model and model-VS-control group comparisons (Figure 3C). Furthermore, some of these miRNAs have reportedly played vital roles in preventing the apoptosis of islet cells and facilitating the recovery of the pancreas functions. For example, let-7d could reduce insulin secretion and impair glucose tolerance; here it was significantly upregulated in the diabetes-VS-normal comparison and downregulated with the DFE administration.

To investigate the transcriptional regulatory interactions underlying the DFE alleviating hyperglycemia in the pancreas, we constructed a miRNA-TF-gene regulatory network using both DEGs and DEMs with opposite expression trends between the two comparisons (diabetes-VS-normal and diabetes-DFE-VS-diabetes; Figure S5). This network contained 21 miRNAs, 22 TFs, and 124 genes, which consisted of 1,245 regulatory pairs. TFs of Stat/Relb/NF-κB/Irf families and miR-148a/375/9a regulating the inflammation and immune-related signaling pathways were represented as hub nodes, which may contribute to the hypoglycemic effect of DFE on DM (Figure 3D). For example, miR-375 could target Jak2 and act as an inhibitor to JAK/STAT signaling pathways, and miR-148a is a repressor of IKKβ/NF-κB signaling, which inhibits the expression of inflammation-related genes, whereas miR-9 works as an inflammation inhibitor through mediating the TLR/NF-κB/mir-9 feedback loop and directly targeting the IL-1β/IRF/NF-κB/JAK/STAT pathway to positively regulate glucose-induced insulin secretion of β cells. Combining the results of regulatory network analysis and functional enrichment of DEGs, we inferred that DFE alleviated hyperglycemia probably through inhibiting severe inflammation-induced islet cell apoptosis in the diabetic pancreas.

Transcriptome Profiling Revealed the DFE Effects on the Energy Metabolism and Lipid Accumulation in the Liver

The liver as a major target organ of insulin plays key roles in lipid metabolism, and fatty liver is a major risk factor of diabetes. To reveal the potential molecular mechanisms of how DFE effects the diabetic liver, we investigated the transcriptome profiling in the liver as well. In the diabetes-VS-normal comparison, the expression levels of 1,222 genes (716 upregulated and 536 downregulated; Figure S6A) and 39 miRNAs (10 upregulated and 29 downregulated; Figure S6B) were significantly changed. The 716 upregulated DEGs were mainly enriched in the gluconeogenesis and non-alcoholic fatty liver disease (NAFLD) biological processes (Figure 4A), whereas the 536 downregulated DEGs were mainly associated with lipid metabolism, glutathione metabolism, and oxidoreductase activity and liver development processes (Figure 4A). Meanwhile, 418 DEGs were significantly upregulated, and 526 DEGs were downregulated in the diabetes-DFE-VS-diabetes comparison (Figure S6C). Notably, the DFE administration significantly restored multiple processes, which were dysregulated in the diabetic rats, such as gluconeogenesis, lipid metabolism, NAFLD, and oxidoreductase activity-related processes (Figure 4A). These results implied that DFE may improve the oxidoreductase activity and energy metabolism, and could decrease the lipid accumulation in the diabetic liver. For example, the expression levels of genes relevant to cholesterol and lipoprotein transport, such as Apoa1-2/Apoc1-3/Lcn2/Rbp4, showed reversed phase between the diabetes and diabetes-DFE groups (Figure 4B). Simultaneously, expression profiles of some miRNAs were markedly changed with DFE administration, such as miR-375/miR-9a/miR-143/miR-127 and miR-486/miR-451 (Figure 4C).

To investigate how the DEGs and DEMs were involved in the decrease of lipid with DFE administration, we constructed a miRNA-TF-gene regulatory network (219 nodes and 1,240 edges) using the DEGs and DEMs with opposite expression profiles in group comparisons of diabetes-DFE-VS-diabetes and diabetes-VS-normal (Figure S7). This regulatory network was involved in the energy and lipid metabolism, which may play important roles in decreasing lipid accumulation in the DFE group. Regulatory interactions between miR-375/9a/143/127/192 and their targets appeared as core modules in our network, suggesting their key roles in the liver underlying the DFE alleviating diabetes (Figure 4D). For example, the miR-192-Srebfl axis plays important regulatory roles in hepatic steatosis, lipid accumulation, and the development of NAFLD. The upregulation of miR-192 in the DFE-administered rats may result in suppression to the expression of TF Srebfl, which could repress the expression of downstream genes related to the lipid biosynthesis, such as Cyp7a1 and Atp5g1. TFs Klf9/Klf13, which were associated with hepatic steatosis and lipid accumulation, were targeted by miR-192/127/143 and acted as key nodes in the network to regulate the expressions of Apoa1 and Scl35a2. The downregulation of
Figure 3. Functional and miRNA-TF-Targets Analysis of the Inflammation and Apoptosis-Related Genes in the Pancreas

(A) Crosstalk of genes involved in the inflammation and apoptosis processes. (B) Heatmap of the crosstalk genes that are involved in apoptosis. (C) Heatmap of differentially expressed miRNAs with opposite profiling in the comparisons of diabetes-DFE-VS-diabetes and diabetes-VS-normal. (D) Core miRNA-TF-genes regulatory network contributed to ameliorated DM under DFE treatment. Blue rectangles: TFs; red triangles: miRNAs; green circles: non-TF genes.
Apa1 could bring about the reduced intake of plasma high-density lipoprotein cholesterol,\(^ {22}\) which thereby resulted in the decrease of lipid biosynthesis.

**Experiments Validated that DFE Could Alleviate Hyperglycemia through Inhibiting Inflammation-Induced Islet Cell Apoptosis**

To validate the anti-inflammation and anti-apoptosis effects of DFE on the diabetic pancreas predicted by the transcriptome profiling analysis, we performed cell and molecular biology experiments (Figure 5). Compared with the diabetes group, the levels of inflammatory markers, such as cytokines IL-1β and tumor necrosis factor alpha (TNF-α), were significantly decreased both in the serum and pancreas with DFE administration (Figure 5A), suggesting that DFE could alleviate severe inflammation in the diabetic pancreas. We also examined the cellular architecture and apoptosis of islet cells from the three groups. Tissue sections of the pancreas islets from normal and DFE groups showed well-preserved circular profiles, whereas pathological changes (massive destruction of the islets) were observed in the diabetic pancreas (Figure 5B). Meanwhile, terminal deoxynucleotidyl transferase (TdT)-mediated dUTP nick end labeling (TUNEL) assay demonstrated that DFE administration significantly decreased the islet cell apoptosis ratio (normal: ~10%, diabetes: ~82%, diabetes-DFE: 39%; Figure 5C). The expression levels of marker genes relevant to inflammation-induced apoptosis, such as Casp3/Bcl2/Bax/Inf-γ/Tnf-α/II-1β/Stat1/II-17α, evidenced that DFE could relieve severe inflammation and apoptosis in the diabetic pancreas as well (Figure 5D). Furthermore, the expression levels of insulin genes were significantly upregulated in the diabetes-DFE group compared with the diabetes group (Figure 5E), suggesting that DFE could inhibit the islet cell apoptosis and thereby increase the insulin level, which contributed to the prevention and treatment of diabetes.

**Experiments Validated that DFE Alleviated Disordered Metabolism and Cell Apoptosis in the Diabetic Liver**

The results of transcriptome profiling analysis indicated that DFE could decrease the glucose homeostasis, increase the oxidoreductase activity and lipid transport, partially attenuate NAFLD, and promote the regeneration of hepatocytes (Figure 6). To verify the effects of DFE on the diabetic liver, we measured the levels of some indicators related to the energy and lipid metabolisms, such as the total cholesterol (TC)/triglyceride (TG)/antioxidant capacity (TAC)/oxidative capacity, glycogen storage, and hepatocyte apoptosis ratio in the liver (Figure 6).

Consistent with the previous study, HFD and DEX could alter the glucose and lipid metabolism in the liver,\(^ {25}\) which resulted in lipid accumulation and lipotoxicity-induced cell death (Figure 6). Our results demonstrated that DFE administration significantly decreased the total cholesterol/triglyceride/oxidative capacity, whereas it increased the total antioxidant capacity and glycogen storage compared with the diabetes group (Figures 6A–6E). Meanwhile, the morphology and structure of hepatic cells were markedly changed with severe lipidosis in the diabetes group, whereas the DFE administration dramatically decreased lipid accumulation (Figure 6F). Furthermore, TUNEL experiments demonstrated the apoptosis of hepatocytes was significantly inhibited in the diabetes-DFE group compared with the diabetes group (apoptosis ratio: ~72.9% in the diabetes group and ~28.3% in the diabetes-DFE group; Figures 6G and 6H).

**DISCUSSION**

Diabetes mellitus is a metabolic disorder disease usually originating from the dysfunction of pancreatic β cells,\(^ {24}\) which could impair insulin target organs (e.g., the liver).\(^ {25}\) In this study, we assessed the anti-diabetic effect of DFE and explored the underlying mechanisms in the pancreas and liver tissues. Our data demonstrated: (1) DFE could reduce the severe inflammation in the diabetic pancreas and prevent the islet cell apoptosis, which could contribute to the protection or recovery of β cell function and mass; and (2) DFE could improve energy metabolism and strengthen the oxidoreductase activity and lipid transport in the diabetic liver, which may thereby decrease lipid accumulation and lipotoxicity-induced hepatocyte apoptosis.

The β cells make up 65%–80% of pancreas islet cells and are injury prone under excessive inflammation.\(^ {26}\) The low insulin and high glucose levels in diabetic rats (Figures 1E and 1F) may be caused by the loss of β cell mass (Figure 5C), which was consistent with a previous study.\(^ {27}\) Our data demonstrated that DFE administration could alleviate hyperglycemia through preventing β cell apoptosis via the anti-inflammation effect in the diabetic pancreas (Figures 5 and 7A). DFE administration significantly downregulated the expression level of a set of key genes (such as Il-1β, Il-17α, Inf-γ, Tnf-α, Stat, and Nf-xB, etc.) compared with the diabetes group, suggesting that Tnf-α and Fas/FasL–dependent apoptotic pathways may be inhibited in the islets with DFE administration.\(^ {28,29}\) Furthermore, DFE administration markedly increased the expression levels of genes involved in insulin signaling and metabolic pathways, including branch chain amino acids (BCAAs) degradation, fatty acid degradation, and PPAR signaling pathways (Figure 5B). The activation of PPAR signaling and fatty acid degradation pathways could reduce β cells apoptosis via upregulating fatty acid oxidation,\(^ {30}\) whereas increasing the BCAAs level could serve as a biomarker for impaired insulin action.\(^ {31}\) Additionally, miRNAs and TFs as important transcription regulators have participated in the genesis and development of diabetes.\(^ {32,33}\) The miR-21/22/146 as pro-apoptosis agents\(^ {34}\) were
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significantly downregulated in the DFE-VS-model comparison. The miR-148a/375/9a combined with TFs Stat/Relb/NF-κB/Srfs acted as core modules regulating inflammation and immune-related signaling pathways, which may contribute to the alleviation of hyperglycemia in the diabetes-DFE group (Figure 3D). The miR-375 and miR-148 play important roles in maintaining pancreatic cell mass, insulin biosynthesis, and normal glucose homeostasis, and here they were highly expressed in the pancreas under DFE administration compared with the diabetes group (Figure 3C). Meanwhile, TFs Stat/Relb/NF-κB/Srfs could promote cell apoptosis, and here they were significantly downregulated with DFE administration (Figures 3C and 5D). Given the above findings together, we inferred that DFE played anti-inflammation and anti-apoptosis roles on the diabetic pancreas (Figure 7A), which could prevent the loss of β cell mass and recover the secretion of insulin (Figure 5).

The liver as a major target tissue of insulin maintains the homeostasis of metabolic processes. Lipid accumulation in hepatocytes is highly prevalent in diabetes and could increase the diabetes risk through disordered lipid metabolism and transport, which may link the dysfunction of insulin to a metabolic disorder of the liver. Excessive lipid accumulation may result in NAFLD (Figure 4A), and further triggers hepatocyte apoptosis and lowers cell regeneration. Our results demonstrated that DFE could: (1) strengthen the oxidoreductase activity, lipid transport, and organ regeneration in diabetic livers (Figures 4A and 6); and (2) significantly decrease the glycogen consumption, lipid accumulation, and apoptosis in the hepatocytes (Figure 6). Genes (Apo family, Lcn2, and Rbp4, etc.) involved in lipid transport showed opposite expression profiles between the diabetes-DFE-VS-diabetes and diabetes-VS-normal comparisons (Figure 4B), suggesting their critical roles in decreasing lipid infiltration under DFE administration. Moreover, core regulatory modules with hubs miR-9/375/127/143/192 and Srebfl/Klf5 (Figure 4D) involved in the lipid metabolism and cell death may play important roles underlying the effects of DFE on the diabetic liver. For example, overexpressed miR-9 could decrease intracellular lipid content in the liver, and up-regulating the expression of miR-375 can inhibit mitochondrial autophagy in HCC cells, which may contribute to the anti-apoptosis effect of DFE on the hepatocytes in our study (Figures 6G and 6H). Additionally, the miR-192-Srebfl-Cyp7a1-Atp5g1 regulatory axis plays pivotal roles in the lipid metabolism/homeostasis. The miR-192/127/143 targeting TFs Klf9/Klf13 regulate their downstream genes and here acted as key modules in our regulatory network of the liver (Figure 4D). These modules were associated with hepatic steatosis and lipid accumulation, and may inhibit the genesis and development of NAFLD. For example, Sk35c2 as an Sk35 family member involved in the cell death process could enhance reactive oxygen species, which was co-regulated by mir-143 and Klf13 in our network (Figure 4D), and its downregulation may contribute to the increase of antioxidant capacity (Figures 6C and 6D). Combining those findings, the most noteworthy effects of DFE on the diabetic liver may focus on maintaining the balance between the energy and lipid metabolism, and then ameliorating the side effects (NAFLD and hepatocyte apoptosis, etc.) induced by disordered gluconeogenesis and lipid accumulation (Figure 7B).

However, due to the RNA quality, the number of sequencing samples was not uniform in different groups or tissues. To avoid this limitation, we validated the key results of transcriptome profiling through physiological and molecular experiments. The high consistency and positive correlation between transcriptome profiling and experimental results indicated that our analyzed results from sequencing data reflected the real situation. Another limitation is that due to the complex compounds within TCM and possible interactions among those materials, the detailed biological active ingredients of TCM are often difficult to identify, and the contribution of each compound on diabetes had not been investigated.

In summary, our findings establish the roles of DFE on regulating the β cell apoptosis and energy metabolism, suggesting that DFE has potential therapeutic implications for hyperglycemia and diabetes. Besides, transcriptome profiling analysis provides an effective approach to elucidate the regulatory pathways and mechanisms influenced by TCM intervention. The combination of transcriptome profiling and regulatory network analyses could offer comprehensive views for changes of gene expression and underlying mechanisms, which could provide new insights to explore the regulatory modules underlying the effects of TCM on diseases as well.

**MATERIALS AND METHODS**

**Preparation of the DFE**

The *D. fimбриatum* were obtained from Jiangxi Xiushui Miraculous Tea Industry (Jiagxi, China). DFE was extracted from dried stems of *D. fimбриatum*. In brief, the dried stem was pulverized and screened with a 100-mesh sieve and then was decocted with 50 vol (v/w) of boiling distilled water for 2 h. The filtrate was collected, and the residue was treated with the same procedure as above again. Finally, all the filtrates were put together and concentrated under a vacuum. Molecular Therapy: Nucleic Acids Vol. 19 March 2020

**Animal Procedure and Sample Preparation**

Male Sprague-Dawley rats weighing a range of 200–230 g and purchased from the Hubei Province Center for Disease Control and Prevention (China) were used in this study. Animal experiments were

**Figure 5. Experimental Validation of the DFE Effects on Inflammation and Apoptosis Processes in the Pancreas**

(A) The level of IL-1β and TNF-α in the serum/pancreas across the three groups (original magnification ×200). (B) Terminal deoxynucleotidyl transferase-mediated dUTP nick end labeling (TUNEL) staining for the islets of the three groups (original magnification ×400). (D) Quantitative real-time PCR results for selected genes involved in the inflammation and apoptosis. (E) The relative expression levels of insulin genes among the three groups. All quantitative data are means ± SEM. *p < 0.05 and **p < 0.01 were determined by one-way ANOVA followed by Newman-Keuls post hoc tests.
approved and performed according to the guidelines of the Institutional Animal Care and Use Committee of HuaZhong University of Science and Technology (permission number: SCXK2015-0018). All reasonable efforts were made to minimize animal suffering.

Rats had free access to food (a standard chow diet) and water with controlled conditions (22 °C ± 2 °C, 12-h light-dark cycle). After an acclimation for 7 days, rats were randomly assigned to three groups: (1) non-diabetic group (n = 6, normal group): feed standard chow diet and saline throughout the whole experiment; (2) diabetes model group (n = 6, diabetes group): administrated with high-fat diet and saline throughout the whole experiment; (3) DFE-treated group (n = 6, diabetes-DFE group): the same treatment as the diabetes group but additional DFE oral administration (100 mg/kg) from 2 to 5 weeks and daily intraperitoneal injection with dexamethasone (DEX; 0.8 mg/kg) from the 4th to 5th week; and (3) DFE-treated group (n = 6, diabetes-DFE group): the same treatment as the diabetes group but additional DFE oral administration (100 mg/kg) throughout the experiment period (Figure 1A).

After experiments, rats were weighed and anesthetized with an intraperitoneal injection of sodium pentobarbital (1.5%, 0.2 mL/100 g). Blood samples (no anticoagulant) were centrifuged at 10,000 g for 10 min at 4 °C. Aliquots of the supernatant were stored at −80 °C for biochemical analysis. The liver and pancreas tissues were rapidly excised and treated in ice-cold PBS with liquid nitrogen. Total RNA with DNase treatment was isolated using the standard TRIzol protocol and quantified using a spectrophotometer (Nanodrop 2000c). Both the tissues and RNAs were stored at −80 °C.


table

**Library Preparation, Sequencing, and Data Processing**

Libraries of RNA-seq (Ribo-Zero) and miRNA-seq were prepared and sequenced at BGI Bioinformatics Institute (Shenzhen, China). Library construction was on the basis of the TruSeq protocol (Illumina, USA), and the Illumina HiSeq2500 platform was employed to sequence these libraries using 2 × 150 bp pair-end strategy. The base-calling procedure was performed using the Illumina CASAVA v1.8.2 pipeline. All raw NGS data have been deposited in the Beijing Institute of Genomics Database (BIGD) (https://bigd.big.ac.cn/databases) under accession numbers BIGD: CRA000812 and CRA000813.

The adaptor removal and data filter procedures were performed using trimmomatic v.0.32 with default parameters. RNA-seq reads with the length less than 35 bp after adaptor trimming or with poly(N) (>5 base) or low-quality bases (quality value ≤ 5, ratio of low quality base > 10%) were removed. For miRNA-seq data, reads with base N or unexpected length (>45 or <15 nt) or low quality (mean quality ≤ 20) were discarded. All of the downstream analyses were based on the clean data with high quality.

**Abundance Estimation and Differential Expression Analysis for Genes and miRNAs**

Clean reads from miRNA-seq were mapped to GenBank, Rfam, and Piwi to identify rRNA, tRNA, small nuclear RNA (snRNA), small nucleolar RNA ( snoRNA), and PIWI-interacting RNA (piRNA). The unmapped reads above were aligned to canonical pre-miRNA sequences of rat from miRBase V21 to identify known miRNAs and generate expression profiles. The normalized values (transcripts per million reads [TPMs]) and read counts of expressed miRNAs in different samples were merged into matrix, respectively. The NOISeq were used to assess differentially expressed miRNAs across the three groups (false discovery rate [FDR] < 0.05 and |fold changes| > 1.5).

Transcript reassembly and quantification were processed according to the HISAT2-StringTie-ballgown pipeline using Rattus norvegicus reference genome (version 6.0, known as “Rnor_6.0”) and Ensembl v.83 annotation. The resulting transcripts were pooled across samples using the merge function of StringTie, discarding all contained and redundant isoforms. Furthermore, for a specific gene, we removed all the isoforms with a Jaccard distance score > 0.98 calculated by BEDTools2. The transcripts obtained above were used to build the “local” annotation file for further analysis. Normalized abundance of genes (fragments per kilobase of transcript per million annotated transcript (FPKM)) were calculated by BEDTools2. The transcripts obtained above were used to build the “local” annotation file for further analysis. Normalized abundance of genes (fragments per kilobase of transcript per million annotated transcript (FPKM)) were calculated by BEDTools2.
fragments mapped [FPKM]) were estimated using StringTie with the local annotation. Genes with FPKM > 1 in any condition (average across replicates) were kept for further analysis. The DEG analysis was performed using ballgown and NOISeq with default parameters and the significance threshold (FDR < 0.05 and |fold change| > 4).

**Biochemical Analyses**

Blood samples from tail veins were used to determine serum glucose concentrations by glucometer (Roche, Switzerland). Overnight-fasted rats were given glucose by oral gavage (2.5 g/kg), and blood samples were collected at 0 (before glucose administration), 30, 60, and 120 min after glucose administration. Insulin concentration was measured by ELISA kit (DRG, Germany). The levels of IL-1β and TNF-α were determined by ELISA kit (Cloud Clone, Wuhan, China) according to the manufacturer’s recommendations. The total cholesterol (TC), triglyceride (TG), antioxidant capacity (TAG), and oxidant activity (TOC) were detected by kits (Jiancheng Bioengineering, China).

**Histopathological Examination and Terminal Deoxynucleotidyl Transferase-Mediated dUTP Nick End Labeling (TUNEL) Staining**

Tissues were dehydrated in graded series of alcohol, embedded in paraffin, and sectioned in 5-mm thickness by using a microtome (ARM-3600; Histo-Line Laboratories, Italy). The sections were dewaxed in three changes of xylene, hydrated in two changes of 100% ethanol, followed by 95% and 80% ethanol, rinsed with water, and then stained with H&E. Histopathological examinations were carried out under light microscope with attached photograph machine (Nikon H600L, Japan).

Apoptotic cells were visualized by TUNEL staining according to the manufacturer’s manual. Tissue sections were digested with Proteinase K and kept for 15 min at 37°C. Slides were placed in a decolorization bed and washed three times (5 min for each time), and then were incubated with terminal deoxynucleotidyl transferase (TdT) buffer followed by TdT reaction solution containing TdT and dUTP for 60 min at 37°C. Further, slides were washed with PBS for 5 min and incubated with anti-digoxigenin peroxidase for 30 min. Color was developed using 0.05% diaminobenzidine (DAB). Sections were then washed, dehydrated, and mounted. Apoptotic cells were identified by a brown stain over the nuclei.

**Quantitative Real-Time RT-PCR**

cDNA was synthesized using PrimeScript RT reagent Kit with gDNA Eraser (QIAGEN, Germany); then real-time PCR was performed using SYBR Premix Ex TaqTMII (TaKaRa Biotechnology, China) in a Bio-Rad C1000 detecting system (Bio-Rad, USA). β-Actin was used as the control, and the fold change for all samples was calculated by the $2^{-\Delta\Delta C_T}$ method.

**Statistical and Regulatory Network Analysis**

SPSS v.17.0 was used for the biochemical analysis. Data from replicate samples are shown as mean ± SEM. Datasets with more than two groups were assessed by one-way ANOVA followed by Newman-Keuls post hoc tests, and p < 0.05 was considered the statistically significant threshold. Kyoto Encyclopedia of Genes and Genomes (KEGG) and Gene Ontology (GO) enrichment analyses were performed on DAVID. The gene list of TFs was downloaded from AnimalTFDB. The method used for regulatory network analysis was described in our previous work.21 Graphs were created in Cytoscape and R studio.

**Ethics Statement**

Animal experiments were approved and performed according to the guidelines of Institutional Animal Care and Use Committee of Hua Zhong University of Science and Technology.

**SUPPLEMENTAL INFORMATION**

Supplemental Information can be found online at https://doi.org/10.1016/j.omtn.2019.10.033.

**AUTHOR CONTRIBUTIONS**

Y.Z., X.Y., and A.-Y.G. designed this work. Q.Z. performed the whole bioinformatics analysis and wrote the manuscript. J.L. performed the experiments and helped edit the manuscript. Q.Z., M.L., and G.-Y.X. performed the data visualization. W.Z. and Y.W. assisted the experiments. Q.Z., Y.Z., and A.-Y.G. revised the manuscript.

**CONFLICTS OF INTEREST**

The authors declare no competing interests.

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