Results Cana reduced the lipid (such as TG, TC, and LDL-C) accumulation in serum, thus decreased atherogenic index of plasma and arteriosclerosis index values. More importantly, Cana decreased the thickness of the vascular basement membrane, improved cardiac mitochondrial homeostasis, and relieved oxidative stress (e.g. regulation of ROS, SOD, GSH, and MDA levels). And, Cana reduced the circulating markers of inflammation (such as TNFα, MCP-1, and IL-6). Myocardial injury was alleviated after Cana treatment with decreasing levels of serum cTn I (from 95 pg/mL to 75 pg/mL) and sCD40L (from 120 pg/mL to 90 pg/mL). Thus, the cardiovascular abnormality (from 95 pg/mL to 75 pg/mL) and sCD40L (from 120 pg/mL to 90 pg/mL) were relieved by elevating the CD31 expression level and mL to 90 pg/mL). Thus, the cardiovascular abnormality from 95 pg/mL to 75 pg/mL and sCD40L from 120 pg/mL to 90 pg/mL was relieved by elevating the CD31 expression level and improving the vascular basement membrane, improved cardiac mitochondrial and gut microbiota homeostasis, and relieved oxidative stress. Moreover, Cana subtly altered microbiota composition in T2DM mice with CVD, which contributed to the improvement of CVD. Collectively, the improvements of myocardial mitochondrial and gut microbiota homeostasis, may represent an important mechanism underlying the cardiovascular benefits of Cana treatment.

Conclusions Cana treatment improved CVD by decreasing the risk of atherosclerosis and reducing the thickness of the vascular basement membrane. Importantly, Cana treatment significantly elevated myocardial mitochondria homeostasis, thus ameliorated the oxidative stress and inflammatory states. Moreover, Cana subtly altered microbiota composition in T2DM mice with CVD, which contributed to the improvement of CVD. Collectively, the improvements of myocardial mitochondrial and gut microbiota homeostasis, may represent an important mechanism underlying the cardiovascular benefits of Cana treatment.

IDDF2021-ABS-0199 REVEALING MOLECULAR AND CELLULAR DISCRIMINANTS OF TYPE 2 DIABETES BY INTEGRATIVE ANALYSIS OF PANCREATIC SINGLE-CELL RNA-SEQ DATA

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Background Type 2 diabetes (T2D) cause the dysfunction of insulin produced in the pancreas by beta cell, which leads to the non-correct response in fat, liver, and muscle cells. However, we are still far from completely understanding the molecular and cellular discriminants for T2D on cell level.

Methods On collected and integrated single-cell RNA-Sequencing (scRNA-seq) data from 3 publicly available pancreas datasets, we have trained an interpretable deep neural network model, which can provide an efficient T2D classifier to estimate the cell states from normal or T2D individuals and selected the key genes. Next, these key genes were used as anchors in down-stream analysis. Then, the cell type, marker gene, gene activity in cells, and cell-cell communications were all extracted and used in estimating the cell-type proportions and cell-type specific differential expressions in individual-matched bulk sequencing data.

Results We trained and tested T2D classifier to predict the states of cells from multiple single-cell datasets, and acquired significantly high accuracy (ACC=0.9034) in independent validation. Among key genes ranked by our classifier, there were 4, 12, 63 differentially expressed genes between normal and T2D states detected in acinar, alpha and beta cells, suggesting the stronger signal of dysfunction of T2D-relevant beta cells than other types of cells. These genes are significantly correlated with metabolic processes relevant to abnormal metabolism of T2D, and mainly show loss of expression, activity, and signaling changes among cell-cell communication in T2D state. Especially, the analysis results suggest that the beta-cell heterogeneity identified in single-cell dataset may be smaller in T2D relative to a normal state; by contrast, the alpha cell shows an opposite effect.

Conclusions Along with the development of high-throughput single-cell sequencing technologies, we are now capable of investigating the T2D pathogenesis on each cell in addition to each individual. There are actually necessary requirements on investigating cellular heterogeneity underlying individual heterogeneity of T2D, for personalized diagnosis and prognosis of T2D.