Detection of Secondary Metabolites as Biomarkers for the Early Diagnosis and Prevention of Type 2 Diabetes

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Background: Type 2 diabetes, or T2D, is a metabolic disease that results in insulin resistance. In the present study, we hypothesize that metabolomic analysis in blood samples of T2D patients sharing the same ethnic background can recover new metabolic biomarkers and pathways that elucidate early diagnosis and predict the incidence of T2D.

Methods: The study included 34 T2D patients and 33 healthy volunteers recruited between the years 2012 and 2013; the secondary metabolites were extracted from blood samples and analyzed using HPLC.

Results: Principal coordinate analysis and hierarchical clustering patterns for the uncharacterized negatively and positively charged metabolites indicated that samples from healthy individuals and T2D patients were largely separated with only a few exceptions. The inspection of the top 10% secondary metabolites indicated an increase in fucose, tryptophan and choline levels in the T2D patients, while there was a reduction in carnitine, homoserine, allothreonine, serine and betaine as compared to healthy individuals. These metabolites participate mainly in three cross-talking pathways, namely “glucagon signaling”, “glycine, serine and threonine” and “bile secretion”. Reduced level of carnitine in T2D patients is known to participate in the impaired insulin-stimulated glucose utilization, while reduced betaine level in T2D patients is known as a common feature of this metabolic syndrome and can result in the reduced glycine production and the occurrence of insulin resistance. However, reduced levels of serine, homoserine and allothreonine, substrates for glycine production, indicate the depletion of glycine, thus possibly impair insulin sensitivity in T2D patients of the present study.

Conclusion: We introduce serine, homoserine and allothreonine as new potential biomarkers of T2D.

Keywords: glucagon signaling, glycine production, bile secretion, insulin sensitivity/resistance.

Introduction

Type 2 diabetes or T2D is an epidemic metabolic disease-causing millions of cases worldwide.1 Incidence of the disease can be identified through measuring common clinical risk factors.2–4 Individuals with fasting glucose (IFG) of 5.6–7.0 mmol/L have an annual risk of 4.7% to be type 2 diabetic. Increased risk of T2D is associated with the impaired glucose tolerance (IGT) and the elevated HbA1c (39–46 mmol/mol [5.7–6.4%]).5–7 Ethnicity seems to influence the incidence rate of diabetes, and the conversion speed from normoglycaemia to dysglycaemia.8
Many plasma metabolites were reported as biomarkers associated with the development of T2D. Alterations in these metabolites can be traced via their biological pathways, examples of which are the amino acid catabolism and hexose metabolism. Previous studies indicated that plasma glycine level is a negative biomarker in patients with diabetes mellitus. Glycine metabolism pathway indicates the participation of many other important metabolites that mostly serve in the biosynthesis of glycine. However, many of these metabolites have not been proven to be associated with insulin resistance or diabetes. In T2D patients, low plasma glycine level was found to be negatively correlated with insulin sensitivity. Insulin sensitizer therapy with pioglitazone + metformin for patients with T2D resulted in increased plasma glycine concentration as compared to placebo, while reduced plasma glycine level was found to be a predictor for impaired glucose tolerance and T2D incidence. The agency of the European Prospective Investigation into Cancer and Nutrition claimed that higher serum glycine is associated with decreased risk of incident T2D as glycine receptors existing in human pancreatic β-cells might stimulate or provoke secretion of insulin. In addition, the reduced plasma betaine level was also found to act as a positive biomarker of diabetes mellitus and proved to be associated with incident T2D.

Bile acid (BA) metabolism was reported to be involved in cholesterol as well as glucose metabolism homeostasis. The latter authors claimed that insulin resistance is positively associated with hyperbileacidemia regardless of diabetes status, which indicates the important role of insulin resistance in the regulation of “bile secretion” pathway in human. BAs also exert a variety of metabolic reactions as they likely promote the secretion of incretin in the digestive tract, improve glucose tolerance, promote insulin secretion, thus positively increase insulin sensitivity. BAs were also found to be increased in diabetic population, while other authors claimed that total bile acids (TBAs) in serum were not changed in T2D patients as compared to control individuals.

In the present study, we hypothesize that metabolomic analysis in blood samples of T2D patients with similar ethnic background can result in the recovery of metabolic biomarkers and pathways that elucidate early diagnosis and predict the incidence of T2D towards the prevention of the disease and its subsequent complications.

Materials and Methods
Participant Recruitment
The study included 34 T2D patients and 33 healthy volunteers recruited from Endocrinology clinics at King Abdulaziz University Hospital (KAUH) in Jeddah, Saudi Arabia between the years 2012 and 2013. As an exclusion criteria, we selected subjects with no comorbid conditions such as ESLD, ESRD, cancer, new onset diabetes after organ transplant, or a recent cardiovascular event within the 3 months prior to blood sampling time. As an inclusion criteria, subjects are mostly from the same ethnic background (Saudi individuals). Blood samples were further obtained from subjects to make some primary metabolic and physical measurements. Ethical approval from King Abdulaziz University Hospital (KAUH), Unit of Biomedical Ethics Research Committee, Jeddah, Saudi Arabia, was issued and written informed consents were obtained from eligible participants (Tables S1 and S2).

This study was conducted in accordance with the Declaration of Helsinki.

Sample Collection and Preparation
Blood samples were collected in the morning (8:00–10:30 am) after a period of ≥10 hrs fasting to avoid variation due to circadian rhythm. The metabolites were extracted from each aliquot in homogenous mixture of −20°C methanol: methyl-tert-butyl-ether:water (1:3:1), then, incubated for 10 mins in an ice cooled ultrasonication bath. The, 650 µL of UPLC-grade methanol:water 1:3, the homogenate was added, and mixture was vortexed and spun for 5 mins at 4°C in a table-top centrifuge. The lower aqueous phase contains the polar and semipolar metabolites. Then, annotation was done using in-house database in order to detect important negatively and positively charged secondary metabolites. R-software (http://cran.r-project.org/) was used for statistical analysis and visualization (PCoA & Hierarchical Clustering (HCA)).

Results
Principal coordinate analysis or PCoA (Figures 1 and 2) and hierarchical clustering or HCA (Figures 3 and 4) were performed for uncharacterized negatively charged (NeC) and positively (PoC) secondary metabolites, respectively, in order to detect the quality of the recovered data. The PCoA plots showed a similar tendency in the distances within each group. For the NeC metabolites, the diagram showed that most values of healthy group were localized in the positive direction of PC1.
and negative direction of PC2, while those of T2D group were localized in the negative direction of PC1 and positive direction of PC2 (Figure 1). The diagram of PCoA for PoC metabolites showed that most values of healthy group were localized in the positive directions of PCoA 1 (PC1) and PCoA 2 (PC2), while those of T2D group were localized in the negative direction of PC1 and almost distributed evenly at the PC2 direction (Figure 2). HCA pattern of the NeC metabolites indicated partial separation between healthy and T2D individuals (Figure 3). While HCA pattern of the PoC metabolites mostly indicated a complete separation between healthy and T2D individuals (Figure 4). Overall, PCoA and HCA patterns across types of metabolites indicated that samples from healthy individuals and T2D patients were largely separated from only few outliers. These results demonstrate that such unannotated NeC and PoC metabolites can help in the separation of individuals with and without T2D (Tables S3 and S4, respectively). We also concluded that the whole yield of secondary metabolites, either PoC or NeC, are responsible for the separation, which proves that the discrimination between the two groups is due to a pattern. However, secondary metabolite data were not in linkage to features such as age, gender, BMI, ethnic background or any other confounding parameters utilized in the present study (Tables S1 and S2).

The total numbers of NeC and PoC metabolites in the present study are 351 and 295, respectively. The inspection of the top 10% NeC and PoC secondary metabolites separating the two groups revealed numbers of eight and 24 metabolites, respectively (Table S5). These 32 metabolites were annotated. Of which, two NeC metabolites, namely fucose and tryptophan, separated the two groups of individuals as they were enriched in the T2D individuals as compared to healthy individuals (Figures 5 and 6, respectively). In addition, a number of six PoC metabolites, namely choline, L-carnitine, betaine, homoserine, allothreonine, and serine separated the two groups of individuals in which choline is the only metabolite that was enriched in the T2D individuals as compared to healthy individuals (Figures 7, while the rest were repressed (Figures 8–10, respectively). Levels of the other NeC and PoC metabolites in this study were arbitrary in control and T2D individuals. The overall results for the contribution of different regulated PoC or NeC metabolites in their corresponding cross-talking pathways are summarized in Figure 11.

Three main cross-talking pathways seem to respond to the disease. They are “glucagon signaling”, “glycine, serine and threonine” and “bile secretion” pathways. Choline is shown to participate in the three pathways, while serine participated in the second and third pathways. Tryptophan, betaine, homoserine and allothreonine also participated in “glycine, serine and threonine” pathway, while carnitine participates in the “bile secretion” pathway. Thus, the regulated metabolites cross-linked the three pathways in a way to respond either positively or negatively to the occurrence of T2D incidence.
Discussion

Type 2 diabetes (T2D) is mainly characterized by impaired insulin production. It arises from insulin resistance of peripheral tissues followed by dysfunction of β-cells in the pancreas due to a sort of metabolic stress. The disease seems to result in differential metabolic activities that cross-link several pathways as a direct response to the disease. We firstly aimed to analyze the untargeted metabolomes of blood samples from patients suffering from T2D and healthy controls and found out that, although uncharacterized, metabolites were able to separate the two groups effectively. The associations of metabolomes of healthy and T2D patients with age, gender, BMI and general health seemed to be negative. The results shown in Figures 1–4 demonstrated that the unannotated NeC and PoC metabolites allowed a nearly perfect separation of both groups with little intermingling. This separation was not due to single or a small number of metabolites, rather,
Figure 5 Analysis of the negatively charged secondary metabolite fucose for control female (red) and male (blue) individuals (as well as female (green) and male (orange) T2D patients). Streaked-colored columns refer to the mean values of the different groups. Original data are shown in Table S3. Subject IDs refer to those described in Tables S1 and S2.

Abbreviations: FC, female control; MC, male control; FP, female patient; MP, male patient.

Figure 6 Analysis of the negatively charged secondary metabolite L-tryptophan for control female (red) and male (blue) individuals (as well as female (green) and male (orange) T2D patients). Streaked-colored columns refer to the mean values of the different groups. Original data are shown in Table S3. Subject IDs refer to those described in Tables S1 and S2.

Abbreviations: FC, female control; MC, male control; FP, female patient; MP, male patient.

Figure 7 Analysis of the positively charged secondary metabolite choline for control female (red) and male (blue) individuals (as well as female (green) and male (orange) T2D patients). Streaked-colored columns refer to the mean values of the different groups. Original data are shown in Table S3. Subject IDs refer to those described in Tables S1 and S2.

Abbreviations: FC, female control; MC, male control; FP, female patient; MP, male patient.

it is due to a large number of compiled metabolites. Previous studies have successfully applied untargeted metabolomes, albeit individual metabolites associated with T2D were also identified.37 We speculate that this work is one of the first that proves that broad coverage unbiased metabolome can be used as an overall biomarker
or predictor for early diagnostics of T2D within the participating cohort of Saudi population. Further analysis can be done to strengthen this important conclusion when studying larger cohorts of Saudi population.

**Figure 8** Analysis of the positively charged secondary metabolite L-carnitine for control female (red) and male (blue) individuals (as well as female (green) and male (orange) T2D patients). Streaked-colored columns refer to the mean values of the different groups. Original data are shown in Table S3. Subject IDs refer to those described in Tables S1 and S2.

**Abbreviations:** FC, female control; MC, male control; FP, female patient; MP, male patient.

**Figure 9** Analysis of the positively charged secondary metabolite betaine for control female (red) and male (blue) individuals (as well as female (green) and male (orange) T2D patients). Streaked-colored columns refer to the mean values of the different groups. Original data are shown in Table S3. Subject IDs refer to those described in Tables S1 and S2.

**Abbreviations:** FC, female control; MC, male control; FP, female patient; MP, male patient.

**Figure 10** Analysis of the three positively charged secondary metabolite homoserine, allothreonine and serine for control female (red) and male (blue) individuals (as well as female (green) and male (orange) T2D patients). Streaked-colored columns refer to the mean values of the different groups. Original data are shown in Table S3. Subject IDs refer to those described in Tables S1 and S2.

**Abbreviations:** FC, female control; MC, male control; FP, female patient; MP, male patient.
Levels of L-fucose and L-tryptophan in the present study were increased in T2D patients as compared to their levels in healthy individuals (Figures 5 and 6, respectively). Serum L-fucose and tryptophan were previously reported to be significantly increased in diabetics when compared to the healthy controls. Shantaram et al\(^{18}\) claimed that high L-fucose level in T2D patients can be a prognostic indicator for type II diabetes mellitus. L-Fucose is a monosaccharide that is a common component of many N- and O-linked glycans and glycolipids synthesized by mammalian cells. Chen et al\(^{19}\) also indicated that tryptophan level in T2D patients positively correlate with insulin resistance and secretion. It was concluded that high levels of these two metabolites can be potential markers associated with the risk of diabetes.

Choline level was increased in the present study in T2D patients (Figure 7). This metabolite has previously proven to promote insulin resistance in a mouse model. Excess amount of this metabolite acts as a signal to activate “glucagon signaling” pathway towards the increased production of plasma glucagon,\(^{40}\) thus the occurrence of hepatic glycogenolysis/gluconeogenesis and the elevated plasma glucose (Figure 11). Glucagon, a 29-amino acid peptide hormone, is important in glucose homeostasis and diabetes pathophysiology.\(^{41-44}\) Glucagon is counter-regulatory to insulin, stimulating hepatic glucose production, thereby increasing plasma glucose levels. Glucagon hormone generally opposes the action of insulin as it stimulates hepatic glucose production that results in the elevation of plasma glucose levels in T2D patients.\(^{45}\) In other words, insulin is acting as a glucose-depositing and anabolic hormone, while glucagon acts as a glucose-mobilizing and catabolic hormone. Thus, the elevated level of choline in T2D patients of the present study can be an important biomarker of high glucagon level and T2D incidence. In addition, choline participates in the “glucose, serine and threonine” pathway and acts two steps upstream betaine towards promoting its level. As betaine showed opposite results to those of choline in terms of their levels in healthy individuals and T2D patients (Figure 9), then we speculate that at least one of the two steps upstream betaine biosynthesis should have been knocked down (Figure 11).

In the “bile secretion” pathway, choline is shown to require the action of two major facilitator superfamily (MFS) transporters, namely organic cation transporter 1 and 2 (OCT1 and OCT2) in order to exit plasma and be incorporated in hepatocytes. OCTs are responsible for the hepatic and renal transport of metformin (Glucophage XR), the treatment of T2D (Figure 11). The latter two transporters were previously reported to be decreased in T2D patients.\(^{46}\) We speculate that the accumulation of choline in plasma of T2D patients can be due to the decreased levels (knockdown) of OCT1 and OCT2. As indicated in the same pathway, carnitine requires the action of OAT2 gene for interring hepatocytes (Figure 11). The latter conclusion was previously supported by Nowicki et al.\(^{47}\) In the present study, carnitine was reduced in T2D patients. High level of this metabolite is effective in improving insulin-stimulated glucose utilization and in reversing abnormalities...
of fuel metabolism associated with T2D. The encoded OAT2 protein was reported to possess transmembrane transporter activity that promotes amino acids availability in skeletal muscle cells and mammalian/mechanistic target of rapamycin complex 1 (mTORC1) activation. The latter authors indicated that insulin level also positively correlates with mTORC1 activation.

Betaine level was also shown in the present study to be reduced in T2D patients (Figure 9). Betaine is an osmolyte and methyl donor, and commonly reduced due to diabetes mellitus. Betaine is essential for our life, and we can obtain it exogenously or endogenously by the mitochondrial oxidation of choline. Many reports claimed that betaine deficiency is a common feature of metabolic syndrome such as T2D. The deficiency can result in the knockdown of downstream glycine biosynthetic steps in the “glycine, serine and threonine” pathway, thus promotes insulin resistance. Unfortunately, the list of the inspected top 10% NeC and PoC secondary metabolites separating the two groups did not include glycine and glucagon, thus, levels of the two metabolites were not detected in the present study.

For the first time, levels of serine, homoserine and allothrionine in the present study were shown to be reduced in T2D patients (Figure 10). In the “glycine, serine and threonine” pathway, these three metabolites were shown to be effective substrates for glycine production. As levels of these three metabolites in the present study were reduced in T2D patients, this indicates that glycine can be depleted. As indicated earlier, plasma glycine level positively correlates with insulin sensitivity, while negatively correlated with insulin resistance.

Reduced plasma glycine level was reported to be a predictor for impaired glucose tolerance in T2D patients. We speculate that the level of the three metabolites might also be predictors for impaired glucose tolerance in T2D patients. As indicated earlier, these are opposite levels of tryptophan and serine as the latter can be depleted towards the production of plasma tryptophan by the action of trpA and/or trpB genes (Figure 11). In healthy individuals, it seems that this reaction is knocked down, while possibly highly expressed in T2D patients.

In conclusion, we introduce serine, homoserine and allothrionine as new potential biomarkers of T2D and urge the detection of their possible participation in insulin resistance. This study can also help in the understanding of the complex biology of metabolic disorders like diabetes and provide a new insight into the biology of insulin sensitivity/resistance.

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Disclosure
The authors report that they have no conflicts of interest and have no affiliation with any organization with a direct or indirect financial interest in the subject matter discussed in the manuscript.

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