Targeted CYP2E1 quantification and its correlation to currently acceptable clinical biochemical indices

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Abstract

Background: The Cytochrome P450 enzymes are commonly known for their major role in metabolism. Besides its metabolic role, CYP2E1 gene expression has been associated with the onset of diabetic nephropathy. CYP2E1 protein elevation has also been reported to be responsible for the production of reactive oxygen species. The aims of this study were (i) to optimize and validate a targeted proteomic approach for quantitating CYP2E1 and validating it as a suitable clinical test, (ii) to investigate the concurrency between ESI-LCMS-MS quantitated circulating CYP2E1 and gold standard indices in the context of outpatient point-of-care clinical settings involving various groups of diabetic patients and (iii) to investigate the concurrency profile of circulating CYP2E1 protein, CYP2E1 gene expression and reactive oxygen species (ROS). This is a cross sectional study involving three groups of subjects (n = 166): control, pre-diabetes, and diabetes. We optimized a targeted proteomic approach for absolute quantification of CYP2E1. “YPEIEEEK” and “GTVVVPTLYDNQEPDPDEK” were the representative peptides of CYP2E1 for our analytical method. Deuterated forms of “YPEIEEEK” and “GTVVVPTLYDNQEPDPDEK” were used as internal standards. Lymphocytes were isolated from whole blood, microsomes were prepared, followed by in-solution digestion for production of tryptic peptides. Amounts of “YPEIEEEK” and “GTVVVPTLYDNQEPDPDEK” from patients’ samples were calculated from a calibration curve.

Results: “YPEIEEEK” is a unique and reliable representative peptide for CYP2E1 quantification. “GTVVVPTLYDNQEPDPDEK” showed poor reproducibility and sensitivity. Incremental amounts of CYP2E1 protein in the peripheral circulation clearly showed concurrency with CYP2E1 gene expression and ROS levels in our study population. Elevations of CYP2E1 were observed even when gold standard clinical indicator for glycemic control (HbA1c) was within normal reference limits. Quantitated amounts of CYP2E1 protein in the pre-diabetes and diabetes groups showed significant difference relative to control group (p < 0.001). No significant differences were observed between the medians of pre-diabetes and diabetes groups (p = 0.870).

Conclusions: CYP2E1 protein in peripheral blood can be reliably quantitated by the validated targeted proteomic approach method. Quantifiable amounts of CYP2E1 preceded abnormal HbA1C levels which indicates quantitation of CYP2E1 could be useful as an additional tool for early indication of diabetic risks and its complications.

Keywords: Absolute quantitation, CYP2E1, Diabetes progression, Early risks predictor, LCMS/MS/MS, ROS
subtypes of tumour specific, while the CYP1A family has been identified predominantly in several types of organ cancers and CYP3A5 is the predominant subtype identified in lung tumour as well as in lung cancer cells [2]. Recent studies have also showed that CYP2E1 expression in peripheral lymphocytes [7, 8] is elevated in diabetes and kidney disorders [9]. The correlation between CYP2E1 level in peripheral blood lymphocytes and early pathogenesis of diabetic nephropathy has recently been demonstrated [10]. Notably, besides the important role that CYP450 superfamily plays in xenobiotics and endobiotics metabolism, individual CYP450 isoforms may pose as candidate markers of a specific disease or pathology [11]. If the proposed candidate markers are measurable earlier than conventional clinical indices, the former may be useful as early indicators of an onset of particular disease. Early indicators of diseases are essential for prevention of further progression and for early planning of effective interventions.

Rapid methods to quantitate these diseases’ specific CYP450 isoforms will allow early diagnosis, prognosis or monitoring a particular disease or may be used alongside conventional clinical tests as an alternative test for confirmation. Many methods have been described for qualitative and quantitative identification of CYP2E1. Earlier common qualitative methods for detection and identification of CYP450 subtypes were gel electrophoresis [12] and immunoblotting [13, 14] which lacked sensitivity and specificity. In the past decade, the quantitative proteomic approach gained popularity and many studies have attempted the application of trypic digest coupling with tandem LCMS analysis [15, 16] allowing absolute quantitation of individual CYP450 isoforms. These approaches demonstrated higher specificity and sensitivity but to date (and according to our knowledge) no studies have described a validated targeted proteomic method for analyzing CYP450s in human peripheral blood. Since a few isoforms of CYP450s have been associated with diseases, it will be worthwhile introducing a clinically applicable method for quantitating CYP450s.

Although various studies have convincingly dictated the association of CYP2E1 [6–10] and reactive oxygen species (ROS) [17] in diabetes and its complications, there is lack of evidence on the changing profile of ROS and CYP2E1 levels with varying glycemic levels. Also, there is lack of evidence on the concurrence of ROS and CYP2E1 with gold standard clinical indices for glycemic control (HbA1c). The knowledge of the latter will further reaffirm that CYP2E1 is a candidate early biomolecule in diabetes progression. Therefore, the aims of this study were firstly, to optimize a targeted proteomic approach for quantitating CYP2E1 in peripheral blood; secondly, to validate its use alongside with clinical indices (HbA1c and fasting blood sugar) currently used in the clinical laboratory, and thirdly, to investigate the correlation of circulating CYP2E1 protein amounts with CYP2E1 gene expression and ROS.

Results and discussion

The representative peptides for CYP2E1 used in this study were YPEIEEK [P1 (amino acid position 318–325)] and GTVVPTLYDNQFPDPEK [P2 (amino acid position: 386–408)]. P1 is suitable for universal quantification of CYP2E1 in human due to absence of polymorphic regions. P2 was included as a confirmatory peptide to the former because it has been employed in other studies [15]. P2 may not be suitable as a representative peptide for universal application in CYP2E1 quantification due to natural amino acid variation between individuals at position 389 [URL: http://www.uniprot.org]. An electrospray ionization source (ESI) was operated in the positive ionization mode for ion production. The most abundant and stable product ions from collision-induced dissociation of each analyte were selected as quantifier ions, while the second most abundant ions were selected as qualifier ions (Fig. 1a). An excellent separation of each analyte of interest was achieved with total run time for each injection of only 5 min (Fig. 1b). The latter shows that the analytical method is highly selective for the analytes of interest.

Partial validation of the optimized chromatographic method showed P1 is a reliable representative peptide of CYP2E1 (Table 1) for the intended purpose of this analytical method. P2 showed unsatisfactory reproducibility and sensitivity in the ESI-LCMS/MS analysis. The normal reference limits for CYP2E1 protein in peripheral circulation has not been documented to date. Therefore, to visually estimate the linear range, six independent calibration curves for each representative peptide were set up. Each calibration curve consisted of 20 individual concentrations ranging from 0 to 10,000 fg µL\(^{-1}\). The observed linear ranges for P1 and P2 were 5–100,000 fg µL\(^{-1}\) (Fig. 2a) and 100–1000 fg µL\(^{-1}\) (Fig. 2b), respectively.

CYP2E1 is a non-abundant protein in human peripheral blood. Current study showed that CYP2E1 was not quantifiable in individuals with normal blood sugar levels (euglycemia) but was quantifiable in some of the prediabetic and most of diabetic subjects at varying amounts (Fig. 3a). The mean values for Pre-D and D groups were significantly different from the control group (C) \((p < 0.001)\). No significant differences were seen between the means of Pre-D and D \((p = 0.870)\). CYP2E1 gene expression correspondingly showed up regulation when HbA1c was still within normal reference limits (Fig. 3b, c).

HbA1c is the gold standard point-of-care test for evaluating blood glucose levels and it is globally used by
Fig. 1 Representative chromatograms of analytes: YPEIEEK (P1), GTVVVPTLDSVLQNEFPDPEK (P2) and internal standards for YPEIEEK (P1-IS) and GTVVVPTLDSVLQNEFPDPEK (P2-IS). a 30 fg µL\(^{-1}\) mixture of P1, P2, P1-IS and P2-IS were prepared in pooled plasma from control subjects and injected into the Agilent 1100 UHPLC coupled to the Agilent 6410 ESI-LCMS/MS system for CYP2E1 analytical method development and optimization. Stable and reproducible transition ions (m/z) were selected for MRM analysis: P1 = (precursor: 454.3; quantifier: 647.3; qualifier: 518.3); P1-IS = (precursor: 476.7; quantifier: 518.3; qualifier: 647.3); P2 = (precursor: 855.0; quantifier: 456.3; qualifier: 585.3); P2-IS = (precursor: 807.1; quantifier: 501.3; qualifier: 585.3). MRM multiple reaction monitoring; P1 peptide 1 (YPEIEEK); P1-IS internal standard for peptide 1; P2 peptide 2 (GTVVVPTLDSVLQNEFPDPEK); P2-IS internal standard for peptide 2. b Representative chromatogram showing good resolution of each analyte of interest from a mixture containing 30 fg µL\(^{-1}\) of each analyte spiked pooled plasma from control subjects. The specific retention time for each analyte was P1: 1.041 min; IS-P1: 1.811 min; P2: 2.808 min and P2-IS: 4.118 min. Total run time for each sample was 5 min.

Table 1 Summary of method validation data

| Parameters       | YPEIEEK (P1) | GTVVVPTLDSVLQNEFPDPEK (P2) | IS-P1 | IS-P2 |
|------------------|--------------|-----------------------------|-------|-------|
| Linear range     | 5–100,000 fg/µl | 100–12,500 fg/µl             | NA    | NA    |
| Linearity        | \(r^2 = 0.991 \pm 0.007\) | \(r^2 = 0.998 \pm 0.001\)     | NA    | NA    |
| LLOQ             | 5 fg/µl      | 100 fg/µl                   | NA    | NA    |
| Specificity      | CV = 4.0 ± 0.04% | CV = 15.9 ± 0.42%           | CV = 2.42 ± 0.05% | CV = 14.2 ± 0.57% |
| Quality control: | L | M | H | L | M | H | NA | NA |
| Intra-day Accuracy, % | 99.2 | 99.8 | 98.2 | 41.0 | 69.2 | 70.6 | NA | NA |
| Precision, CV%   | 3.64 | 4.79 | 10.93 | 55.14 | 5.32 | 17.13 | NA | NA |
| Inter-day Accuracy, % | 91.75 | 105.08 | 90.09 | 70.8 | 82.6 | 84.14 | NA | NA |
| Precision, CV%   | 9.76 | 4.56 | 0.77 | 42.54 | 12.62 | 20.13 | NA | NA |

The optimized analytical method for CYP2E1 quantification was validated by assessing the linear range, linearity, LLOQ, specificity, intra and inter-group accuracy and precision. IS-P1 internal standard for P1; IS-P2 internal standard for P2; NA not applicable. Coefficient of variation (CV, %) is presented as mean ± standard deviation. Coefficient of variation, means and standard deviations were calculated using the Sigma plot 11.2 software. Overall, the analytical method showed higher sensitivity and specificity for P1 (LLOQ = 5 fg µL\(^{-1}\); CV = 4.0 ± 0.004) in comparison to P2 (LLOQ = 100 fg µL\(^{-1}\); CV = 15.9 ± 0.42). Higher intra-day accuracy (L = 99.2 %; M = 99.8 %; H = 98.2 %) and precision (L: CV% = 3.64; M: CV% = 4.79; H: CV% = 10.93) were observed for quantification of P1 in comparison to P2 (L: CV% = 41.0 %; M: CV% = 69.2 %; H: CV% = 70.6 %) for all levels of QC quantitated while P2 quantification showed slightly lower accuracy (L = 70.8 %; M = 82.6 %; H = 84.14 %) for all levels of QC in comparison to P1. Inter-day precision for quantification of P1 at all 3 QC levels showed CV% < 10 showing that the quantification method was highly precise for P1. P2 showed poor precision (L: CV% = 42.54 %; M: CV% = 12.62 and H: CV% = 20.13 %) for L and H QC levels. Overall, P1 showed higher reproducibility and reliability as representative peptide for CYP2E1 in the optimized targeted proteomic approach.
clinicians for diagnosis of diabetes as well as for evaluating glycemic control alongside fasting blood glucose. When blood glucose exceeds homeostasis level it naturally binds to haemoglobin to form glycosylated hemoglobin (HbA1c). HbA1c values indicate average blood glucose levels for the past 2–3 months from the day HbA1c is tested on the patient [18]. The HbA1c test does not reflect patients glycemic status beyond 3 months from the time of current test, unless HbA1c was tested once every 3 months repetitively. HbA1c levels continue to rise when blood sugar increases and decreases when patients consume oral antihyperglicemic drugs or when they undergo lifestyle modification as often prescribed by physicians. With regard to the latter, HbA1c does not represent cellular glucose metabolism processes during the hyperglycemia milieu which precedes the increase in blood glucose levels. Therefore, HbA1c values do not predict risks of diabetic complications.

Several studies commonly explained that within the hyperglycemic milieu ROS is grossly generated and promotes elevation of CYP2E1 protein levels [8–10]. CYP2E1 itself has oxidative properties [19] and along with glucose auto-oxidation is capable of generating more ROS which leads on to a positive turnover of ROS and CYP2E1; hence, enhancing a destructive environment at cellular level.

Measurement of ROS per se may not be suitable as an assessment of hyperglycemic related damage risk or severity of hyperglycemia. This is because the increase in ROS at any time may be transient due to the interplay of environmental factors, diet and various oxidants generated during normal intracellular metabolism. All these factors contribute to large inter-individual variations within each study group (Fig. 3d). The normal reference limit for ROS in human is ≤ 310 FORT units (FORT = free oxygen radical test) [20]. Moreover, in the human body an efficient natural antioxidant defence mechanism exists which aims to maintain physiological levels of ROS [21]. An imbalance of ROS homeostasis takes place during the onset of diabetes bias towards ROS generation at cellular level [17]. The latter processes create a toxic-driven catastrophic environment which
hypothetically triggers the up-regulation of CYP2E1 to play its physiological role to detoxify the environment. Consistent with diabetes being a systemic disease it is possible that this initial up-regulation of CYP2E1 gene occurs in peripheral lymphocytes and the CYP2E1 protein becomes quantifiable in peripheral lymphocytes. CYP2E1 itself has oxidative properties [1] therefore, sustained increment in CYP2E1 will not be a favourable condition and together with auto-oxidation of glucose the cellular environment becomes destroyed and can contribute to multiple forms of pathogenesis.

Therefore, an undiagnosed diabetic patient who has abnormal HbA1c level at the first visit to a clinician may already be at risk for diabetic complications. According to clinical guidelines [22], HbA1C $\geq$ 6.5 % is equivalent to normal blood glucose level, but our data showed that CYP2E1 protein was quantifiable in some individuals even when HbA1C was < 6.5 %. On average, CYP2E1

Fig. 3  Correlation of CYP2E1 protein levels, CYP2E1 gene expression and ROS, with glycemic control. a A scatter plot generated using the SigmaPlot 11.2 software, presenting measurable amounts of CYP2E1 protein in individual subjects (n = 166). Data from this scatter plot shows measurable amounts of CYP2E1 were seen even when HbA1c was 4.2 %. b CYP2E1 gene expression presented as a heat map generated using the GeneSpring 12.6 software. The heat map shows averaged CYP2E1 gene expression of study subjects who were grouped according to their HbA1c percentage. The averaged CYP2E1 gene expression for the groups of subjects who had HbA1c [(< 5.0 %); n = 8] and [(5.1–6.0); n = 20] showed down regulation and groups for subjects with HbA1c percentage [(6.1–7.0); n = 19], [(7.1–8.0); n = 11], [(8.1–9.0); n = 20], [(9.1–10.0); n = 17], [(10.1–11.0); n = 9], [(11.1–12.0); n = 6], [(12.1–13.0); n = 6] showed up-regulation. The gradual change in hues of green from dark green for group (6.1–7.0) to the lightest green in group (12.1–13.0) shows the increments in up regulation intensities of CYP2E1 gene as the HbA1c percentage increases. c CYP2E1 gene expression presented as a heat map generated using the GeneSpring 12.6 software. The heat map shows CYP2E1 gene expression for individual subjects with various HbA1c percentages. The heat map shows that CYP2E1 gene is up regulated even when HbA1c was 5.4 %, which is well within the normal reference limit for HbA1c in clinical practice. d A bar chart representing average ROS levels in the groups of subjects who had HbA1c [(< 5.0 %); n = 8], [(5.1–6.0); n = 20], [(6.1–7.0); n = 19], [(7.1–8.0); n = 11], [(8.1–9.0); n = 20], [(9.1–10.0); n = 17], [(10.1–11.0); n = 9], [(11.1–12.0); n = 6] and [(12.1–13.0); n = 6]. Error bars represent standard deviations (SD) within each group. SD was calculated using the Sigma Plot 11.2 software. Overall, the increment in ROS levels showed increment trends with increasing HbA1c percentage similar to CYP2E1 gene expression and protein amounts. This observation strongly supports the notion that the increase in ROS production during hyperglycemia induces up-regulation of CYP2E1 gene expression in peripheral lymphocytes and subsequently increases in CYP2E1 protein.
protein (Fig. 3a) as well as CYP2E1 gene expression (Fig. 3b, c) were higher in diabetic individuals who had higher glycemic levels (HbA1c). Findings from this study clearly suggest that the appearance of CYP2E1 protein in peripheral lymphocytes precedes glycosylation of haemoglobin. Data from this study also suggest that the conventional clinical criteria [22] for diagnosis of diabetes based on HbA1c and fasting blood glucose alone may prevent early recognition of the risks of onset of diabetes and possibly its complications.

Quantification of CYP2E1 in diabetic patients gives a prediction of cumulative oxidative damage and toxicity at cellular level due to hyperglycemia at the time the patient is tested for this marker. Since CYP2E1 was not detected in control subjects, the lowest quantifiable amounts during hyperglycemia should be regarded as a risk factor for diabetic complications because CYP2E1 is catastrophic by nature and is capable of exacerbating cellular damage and can be potentiated by other extrinsic and genetic factors.

With complete validation of our optimized targeted proteomic approach method, quantification of CYP2E1 in peripheral blood will be useful as an additional tool alongside current clinical indices for predicting risk of developing diabetes in patients with impaired glucose tolerance or impaired fasting glucose as well as for predicting risks for diabetic complications.

Our findings complement and strengthen previous discussions [17] on the role of ROS in diabetes progression. Data from this study clearly provide evidence that the amounts of ROS correlates incrementally with the severity of hyperglycemia indicated by increasing HbA1c levels (Fig. 3d).

Conclusions
Consistent with the aims of our study, we evidently showed that the optimized targeted proteomic approach for quantifying CYP2E1 in peripheral blood is suitable for clinical application. “YPEIEEEK” is unique to CYP2E1 and does not carry any polymorphic regions. “YPEIEEEK” showed high inter-day and intra-day accuracy and precision. Therefore, “YPEIEEEK” is a suitable and reliable probe for quantification of CYP2E1 in the general population.

The increment of CYP2E1 protein was consistent with increment of HbA1c values (gold standard marker for glycemic control) in peripheral blood, but the former precedes abnormal HbA1c values. This observation evidently suggests that CYP2E1 protein levels in peripheral blood is a potential candidate biomarker for risk of developing diabetes and probably its complications in prolonged hyperglycemia.

Data from this study showed in a cohesive manner, the amounts of circulating CYP2E1 protein amounts and gene expression in peripheral blood increases in parallel with ROS. This finding compliments and strengthens the notion that ROS is increasingly being generated during hyperglycemia and subsequently influences the up-regulation of CYP2E1 gene and protein expression in peripheral lymphocytes.

Methods
Materials
Reagents were obtained from Sigma Chemicals Co. (St. Louis, MO) and Agilent Technologies (M) Pte. Ltd. unless otherwise stated. Recombinant human CYP2E1 expressed in Escherichia coli was from Euro Science (M) Pte. Ltd. Synthetic peptides “YPEIEEEK (P1)” and “GTVVPTLYDQNQFDPDPEK (P2)” were ordered from First BASE Laboratories (M) Pte. Ltd. LCMS analytical column was purchase from Agilent Technologies (M) Pte. Ltd.

Research design and study subjects
This study was a cross sectional study. Study population (n = 166) consisted of three groups: normal healthy individuals (Control; n = 45), pre-diabetes (Pre-D; n = 25), and diabetes (D; n = 96). Our inclusion criteria were adult males and females aged 30–60 years and non-smokers. Our exclusion criteria were pregnant women and individuals who just recovered from any form of illness or surgery. Study subjects were grouped based on fasting plasma glucose levels (FPG), HbA1c and oral glucose tolerance test (OGTT) with reference to the American Diabetes Association (ADA) criteria for diagnosis of diabetes mellitus [23]. Control subjects (C) were individuals with fasting plasma glucose FPG < 5.6 mmol L$^{-1}$ and/or HbA1c < 6.5 % on the recruitment day. The OGTT test reveals individuals with impaired fasting glucose (IFG) and impaired glucose tolerance (IGT). IFG is when plasma glucose level after an overnight fast (8 h) is ≥ 7.0 mmol L$^{-1}$. IGT is when an individual has plasma glucose level ≥ 11.0 mmol L$^{-1}$ 2 h after consuming 75 g glucose. Individuals with IFG or IGT were assigned to the pre-diabetes group (Pre-D). Individuals who had 2 h post prandial glucose ≥ 7.8 < 11.0 mmol L$^{-1}$ or FPG ≥ 6.7 < 7.0 mmol L$^{-1}$ and/or HbA1c ≥ 5.7 < 6.5 % were assigned to the Pre-D group [23]. The diabetes group (D) consisted of individuals who had fasting plasma glucose (FPG) ≥ 7.0 mmol L$^{-1}$ and/or HbA1c ≥ 6.5 %, individuals with both IFG and IGT and individuals who are already on antihypertensive medication. All subjects were instructed not to consume any form of alcoholic beverages or food for 7 days before phlebotomy. Subjects who
had any major health conditions within 5 months from the date of phlebotomy were excluded from participation. Characteristics of our study population are summarized in Table 2. Informed written consent was obtained from all subjects following human research ethics approval from the Ministry of Health Malaysia, Medical Research and Ethics Committee (NMRR: 12-501-11926). This study conforms to the Declaration of Helsinki.

**Sample collection**

All subjects were instructed to fast overnight and phlebotomy was performed the following morning on each subject at our clinical research center (Monash University Medical Precinct). Approximately 8 µL of venous blood was collected into tubes containing EDTA anticoagulant. All phlebotomy sessions were between 8 a.m. till 11 a.m.

**Selection of CYP2E1: specific tryptic peptides**

CYP2E1 unique peptides were selected in silico [24, 25] using the Agilent Spectrum Mill peptide selector (Revision: 3.3.084). Tryptic peptides which are portions of the membrane spanning region of CYP2E1 and those containing natural variant amino acid sequences were identified and excluded using the UniProt software http://www.uniprot.org/uniprot/PO5181. Subsequently, “YEPIEEK” and “GTV-VVPPTLDSVLYDNSQFDPPEK” were selected as suitable unique analytical target peptides to quantify CYP2E1.

**Isolation of lymphocytes from venous blood**

Peripheral blood lymphocytes were isolated using NycoPrep media (Axis-Shield, Oslo, Norway) according to an established method [26]. 8 ml whole blood from each subject was diluted with equal volume of 0.9 % NaCl in a 50 mL Falcon tube. 16 µL of diluted blood was then carefully layered on 8 ml NycoPrep media in another 50 µL Falcon tube. Falcon tubes were then capped to prevent the formation of aerosols. All tubes were then centrifuged at 800×g for 20 min at room temperature in a swing out rotor (Beckman, Allegro X-12, USA). After centrifugation, the lymphocytes form a distinct band at the interface between the sample layer and the NycoPrep media. A pasture pipette was used to transfer the lymphocytes suspension into 15 µL glass centrifuge tubes without removing the upper layer. Equal volume of phosphate buffered saline (PBS) was added into centrifuge tubes containing lymphocytes suspension, mixed by gently inverting the tubes until the lymphocytes were homogenously diluted. Next, all tubes were centrifuged at 400×g for 10 min at room temperature. Supernatants were discarded and lymphocytes were resuspended in PBS and centrifuged again at 400×g for 10 min at room temperature. Supernatants were discarded and 1 µL 0.1 M Tris–HCl pH 7.4 was added into each tube, mixed gently to dislodge the lymphocytes and bring them to a homogenous suspension. All lymphocyte suspensions were then transferred into pre-labeled 1.5 µL microfuge tubes and immediately stored at −80 °C until used for microsome preparation.

**Preparation of lymphocyte microsomes**

Microsomes were prepared according to established method [27] with slight modifications. Isolated lymphocytes stored at −80 °C were thawed gradually at room temperature, mixed gently by tapping at the bottom of the microfuge tubes and placed on ice. Lymphocytes were disrupted by sonication for 3 × 60 s on ice, followed by centrifugation at 10,000×g for 30 min at 4 °C (Beckman Coulter, Optima L-100 XP Ultracentrifuge). The supernatants were transferred into new microcentrifuge tubes, and centrifuged again at 100,000×g for 90 min at 4 °C. From this step onwards, all supernatants were discarded and the pelleted microsomes were resuspended in 100 mM sodium pyrophosphate buffer, pH 7.4, to inhibit the NADPH destructing pyrophosphatase activity of the microsome preparation. This was followed by another centrifugation at 100,000×g for 60 min at 4 °C. The washed microsomes were resuspended in 50 mM potassium phosphate buffer, pH 7.4, containing 0.1 mM EDTA, 0.1 mM dithiothreitol (DTT) and 20 % glycerol. Microsome suspensions were transferred into 1.5 µL microcentrifuge tubes and stored immediately at −80 °C until used for subsequent protocol.

**Isolation of soluble CYP2E1**

CYP2E1 is a membrane bound protein. To facilitate the analysis and quantification this protein is enriched and solubilized in order to maximize cleavage by trypsin during the in-solution digestion procedure. CYP2E1 tryptic peptides were extracted similar to the procedure described by Liao et al. [28] with slight modifications to gain maximum

| Variables     | Study groupings | C (n = 45) | Pre-D (n = 25) | D (n = 96) |
|---------------|----------------|-----------|---------------|-----------|
| Gender        |                | M         | F             | M         | F         |
|               |                | 12        | 12            | 47        |            |
| Age (years)   |                | 42 ± 13   | 50 ± 11       | 54 ± 9    |            |
| HbA1c (%)     |                | 4.4 ± 0.36| 6.0 ± 0.39    | 8.7 ± 2.0 |            |
| FBS (mmol L⁻¹)|                | 4.9 ± 0.48| 6.0 ± 1.16    | 10.0 ± 4.68| |
| CYP2E1 (ag µL⁻¹) |              | 0 ± 0     | 7.66 ± 16.39  | 100 ± 20.79| |

Data are presented as mean ± SD

C control, Pre-D pre-diabetes, D diabetes, m male, f female, FBS fasting blood sugar

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recovery. Briefly, lymphocyte microsomes were sonicated (Sonicator 3000, Misonix Inc., Farmingdale, NY) followed by centrifugation at 153,000×g for 30 min. Lymphocyte lysate protein concentration was determined using Pierce bicinchoninic acid (BCA) protein assay kit (Thermo Scientific, IL, USA). Membrane-bound proteins were solubilized by placing the resulting total membrane pellets in 0.2 % sodium cholate in 25 mM NH₄HCO₃ and vortexed continuously at 700 rpm for 60 min at room temperature. Subsequently, the dissolved membrane proteins were heated at 90 °C for 5 min to denature the solubilized membrane proteins to make them amenable to cleavage by trypsin. All samples were allowed to cool at room temperature, then treated with trypsin (Sigma-Aldrich, catalog number: T0303) for 15 h at 37 °C. The substrate to trypsin ratio used was 50/1 (w/w). After hydrolysis, 50 fg µL⁻¹ internal standard peptides (deuterated “GTVVVPTLD5VLVDNQEPDPEK” and “YPEIEEEK”) were added to each sample and centrifuged at 153,000×g for 30 min. Supernatants for each sample were transferred to clean tubes and equal volumes of acetonitrile were added. All samples were dried using a Vacufuge (Eppendorf AG, Hamburg, Germany).

**Analysis and quantitation by MRM using UHPLC-ESI- LC–MS/MS**

Analytical parameters for quantitation of target peptides were optimized on the Agilent 1100 UHPLC system (Agilent, Santa Clara, USA) which was connected to an ESI triple quadrupole mass spectrometer (Agilent 6410). The targeted unique peptides were separated on ZORBAX Eclipse Plus C8 Rapid Resolution column (Agilent, RRHD; 2.1 × 50 mm; 1.8 µm particles) and eluted using a non-linear mobile phase schedule. The mobile phase consisted of 0.1 % formic acid in water (solvent A) and 0.1 % formic acid in 90 % acetonitrile/water (solvent B).

**Analytical method validation**

**Linearity and range**

Synthetic peptides (YPEIEEEK, and GTVVVPTLD5VLVDNQEPDPEK) were serially diluted to obtain final concentrations of 1000, 800, 600, 400, 200, 100, 80, 60, 40, 20, 10, 5 and 0 fg µL⁻¹ in pooled plasma from normal volunteers containing 50 fg µL⁻¹ of each internal standard. Triplicates of each calibrator was injected into the LCMS/MS system to generate standard curves (n = 6).

**Specificity**

Synthetic peptides and internal standard peptides were prepared as a mixture containing 30 fg µL⁻¹ of each peptide in pooled plasma from normal volunteers. 1 µL of peptide mixture was injected into the LCMS/MS system. The injections were repeated 10 times in the same day. Specificity was evaluated by comparing the repeatability of retention times of each peptide [29, 30].

**Accuracy**

Three levels of quality control (QC) standards for each peptide (10, 30 and 60 fg µL⁻¹) containing 50 fg µL⁻¹ of each internal standard were prepared in pooled plasma from normal volunteers. Triplicates of each QC standards were injected and data were collected for 6 runs. The observed values for each QC were then predicted from a 7 points calibration curve (500, 250, 125, 50, 25, 5, 0 fg µL⁻¹). Percentage of error (bias) was calculated as:

\[
\text{Error} \% = \frac{\text{Expected value} - \text{Observed value}}{\text{Expected value}} \times 100.
\]

Percentage of accuracy was calculated by subtracting error (%) value from 100 %.

**Precision**

Precision was assessed by comparing the peak area ratios from 6 runs of QC standards (3 levels) for each peptide. Peak area ratio is calculated by dividing the area under the curve (AUC) for analyte peptide by AUC for internal standard peptide [29, 30]. Preparations of QC standards were similar to the QCs prepared for accuracy evaluation. Mean (n = 6) percentage coefficient of variation (CV) for each QC should be ≤15 % [29, 30].

**Intra-day precision**

The procedure for precision and accuracy assessment was repeated for 6 days and the percentage of CV for 6 days was calculated.

**Inter-day precision**

The procedure for precision and accuracy assessment was repeated for 6 runs in a single day and the percentage of CV was calculated for observation.

**Measurement of reactive oxygen species (ROS)**

ROS was measured using a free oxygen radical monitor (FORM-OX; Callegari, Italy).

**Measurement of HbA1c**

Glycosylated hemoglobin (HbA1c) was measured using Afinion AS100 analyzer (Axis-Shield, USA) according to manufacturer’s protocol. This instrument conforms to the National Glycohemoglobin Standardization Program (NGSP) and diabetes control and complications trial (DCCT) standards.

**CYP2E1 gene expression**

Total RNA was extracted from each sample using RNeasyPlus Micro Kit (Qiagen, Germany) according to manufacturer’s protocol. Purity (A₂₆₀/A₂₈₀), RNA integrity number (RIN) and concentration of extracted total RNA were assessed using Agilent 2100 Bioanalyzer with RNA
6000 Nano Reagents (Agilent Technologies, Santa Clara, USA). Only samples with RIN 7.0 or higher were selected for the gene expression experiment. 100 ng of total RNA of each sample was processed and hybridized to SurePrint G3 Human Gene Expression Microarray platform (G4858A; Agilent Technologies, Santa Clara, USA) according to manufacturer’s protocol. Microarray slides were scanned using Agilent G4900DA Sure Scan Microarray Scanner (Agilent Technologies, Santa Clara, USA). After scanning the image data files were imported into the GeneSpring GX gene expression software, Version 12.6 (Agilent Technologies, USA) for detailed analysis.

Statistical analysis
Significant difference in CYP2E1 protein levels between the studied groups (C, PD, D) were calculated using SigmaPlot 11.2 software. Since our study samples were not drawn from normally distributed populations with the same variances, we used the Mann–Whitney rank sum test. Statistical analysis was performed in a pair-wise manner (C versus PD; C versus D and PD versus D). Differences between study groups were considered significant at \( p < 0.05 \). Intra-group variations for variables in subjects and study parameters were described as standard deviations (SD). SDs were also calculated using the SigmaPlot statistical software.

Authors’ contributions
CGY: Corresponding author, project planning, principal investigator, compilation of data, analysed data, preparation of manuscript. AZ: Contributed in project planning, discussion and conclusion. IO: Contributed in project planning, proteomics analysis. All authors read and approved the final manuscript.

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Competing interests
The authors declare that they have no competing interests.

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