Picrorhiza kurroa Enhances β-Cell Mass Proliferation and Insulin Secretion in Streptozotocin Evoked β-Cell Damage in Rats

Shiv Kumar1,2, Vikram Patial1,2, Sourabh Soni1,2, Supriya Sharma1,2, Kunal Pratap1, Dinesh Kumar2,3* and Yogendra Padwad1,2*

1Pharmacology and Toxicology Lab, Food and Nutraceuticals Division, CSIR-Institute of Himalayan Bioresource Technology, Palampur, India, 2Academy of Scientific and Innovative Research, New Delhi, India, 3Natural Product Chemistry and Process Development Division, CSIR-Institute of Himalayan Bioresource Technology, Palampur, India

Autoimmune destruction of insulin producing pancreatic β-cells leads to insulin insufficiency and hyperglycemia in type 1 diabetes mellitus. Regeneration of β-cells is one of the proposed treatment for type 1 diabetes and insulin insufficiency. Picrorhiza kurroa is a medicinal herb and is traditionally being used for the treatment of various diseases. Previous studies reported the hypoglycemic potential of P. kurroa. However, its potential role in β-cell induction in insulin secretion have not been fully investigated. Here, we characterized the hydro alcoholic extract of P. kurroa rhizome (PKRE) and further studied its β-cell regeneration and induction of insulin secretion potential in streptozotocin (STZ) induced diabetic rats as well as in insulin producing Rin5f cells. 1H-NMR revealed the presence of more than thirty metabolites including picroside I and II in PKRE. Further, we found that PKRE treatment (100 and 200 mg/kg dose for 30 days) significantly (p ≤ 0.05) protected the pancreatic β-cells against streptozotocin (STZ) evoked damage and inhibited the glucagon receptor expression (Gcgr) in hepatic and renal tissues. It significantly (p ≤ 0.05) enhanced the insulin expression and aids in proliferation of insulin producing Rin5f cells with elevated insulin secretion. Furthermore it significantly (p ≤ 0.05) increased insulin mediated glucose uptake in 3T3L1 and L6 cells. On the contrary, in diabetic rats, PKRE significantly (p ≤ 0.05) decreased high blood glucose and restored the normal levels of serum biochemicals. Altogether, our results showed that PKRE displayed β-cell regeneration with enhanced insulin production and antihyperglycemic effects. PKRE also improves hepatic and renal functions against oxidative damage.

Keywords: streptozotocin, β-cell regeneration, insulin expression, glucose uptake, P. kurroa

Abbreviations: ALP, alkaline phosphatase; ALT, alanine transaminase; AST, aspartate transaminase; ATCC, American type culture collection; BUN, blood urea nitrogen; CAT, catalase; CK, creatinine kinase; DMEM, Dulbecco’s modified eagles medium; FBS, fetal bovine serum; Gcgr, glucagon receptor expression; GLB, glibenclamide; MDA, malondialdehyde; NADH, nicotinamide adenine dinucleotide; NBT, nitroblue tetrazolium; OGTT, oral glucose tolerance test; PKRE, Picrorhiza kurroa rhizome extract; PMS, phenazine methosulphate; SOD, superoxide dismutase; SRB, sulforhodamine B; STZ, streptozotocin; T1DM, Type 1 diabetes mellitus, 2NBDG, 2-deoxy-2-[(7-nitro-2,1,3-benzoxadiazol-4-yl)amino]-d-glucose; TBA, thiobarbituric acid; TCA, trichloroacetic acid.
INTRODUCTION

 Destruction of insulin producing β-cells is one of the serious chronic autoimmune disorders, characterized by infiltrating autoreactive lymphocytes, macrophages and other immune cells leading to diminished insulin secretion and hyperglycemia (Soltani et al., 2011). It is one of the most prevalent pediatric endocrine illnesses in developing nations and is termed as type 1 diabetes mellitus (T1DM). T1DM incidences increase globally by 2–5% per year (Poudel et al., 2015) and in India approximately 0.97 million children (below the age of 15) are affected (Kumar, 2015). T1DM generally manifests during early childhood and adolescence, potentially triggered by autoimmune, genetic, as well as environmental factors (Kakleas et al., 2015). It has been reported that 85–90% of β-cells are lost during onset of clinical symptoms, attributed to autoimmunity (Klinke, 2008). A meta-analysis study revealed that, 78% of insulitis occurs in children below 14 years, whereas 29% insulitis was diagnosed between 15 to 40 years of age. However, the severity of β-cells death decreased with age (Battaglia et al., 2015). Currently available antidiabetic drugs potentially sustain the blood glucose homeostasis with several adverse complications (Meier et al., 2015). Hence, alternative medications derived from medicinal herbs with potent hypoglycemic activity and minimal undesired complications are gaining importance. Ethnobotanical knowledge revealed that extracts and isolated molecules derived from medicinal plants have been widely used to treat type I and type II diabetes (Sheikh et al., 2015; Coelho et al., 2016; Devi et al., 2016).

Traditionally, roots and rhizomes of the herb P. kurroa found in Indian Himalayan region were widely used to treat fever, hepatitis, respiratory tract diseases, allergies and other inflammatory conditions (Dwivedi et al., 1992; Luper, 1998). Owing to the various pharmacological attributes of this medicinally important endangered herb, researchers developed efficient protocols for its in vitro mass multiplication and alternative agronomic practices (Nautiyal et al., 2001; Mahajan et al., 2016; Tanuja and Preeti, 2016). It is well reported that the pharmacological efficacy of this herb is due to kutkin, an active principal which comprises of iridoid glycosides (Picroside I and II) and kutkoside mainly localized in the roots and rhizome (Shitiz et al., 2013). Recently it was reported that iridoid glycosides potentially inhibit the formation of antiglycation products, resists hyperglycemia and regulates immunity in vitro as well as in vivo (Chen et al., 2016). Studies have shown that picroliv, a derivative of P. kurroa ameliorates dextran-sulfate-sodium-induced colitis in mice by attenuating the activity of SOD and myeloperoxidase. Further, it also reduces the transcript levels of TNF-α and IL-1β by inhibiting NF-κB expression (Zhang et al., 2012). P. kurroa rhizome extract exhibited promising antioxidant activity against indomethacin induced gastric ulcers in rats. Treatment significantly improved the SOD and catalase activity along with reduction in lipid peroxides (Ray et al., 2002). Preliminary observations by Joy and Kuttan (1999) revealed hypoglycemic effect of P. kurroa in alloxan induced diabetic rats, however, a systematic and detailed investigation on the potential of P. kurroa in curtailing insulin insufficiency is still lacking. Therefore present study was aimed to characterize the PKRE using metabolomic approaches and its β-cell proliferation efficacy, insulin gene expression as well as its secretion in vitro and in vivo. Study also investigates PKRE's impact on oxidative damages in diabetic rats along with glucose uptake in vitro. Phytomolecular profiling results revealed the presence of more than thirty one major metabolites including picroside I and II in PKRE. Treatment effectively elevates the pancreatic β-cell mass in vivo with enhanced proliferation and insulin gene expression in Rin5f cells. Also it further enhances the glucose uptake in differentiated adipocytes and myotubes. Study also reports that PKRE improves the hepatic and renal complications associated with hyperglycemia and restores the normal functioning.

MATERIALS AND METHODS

Chemicals and Reagents

Dexamethasone, insulin, protease inhibitor cocktail, rosiglitazone and 2-deoxy-2-(7-nitro-2,1,3-benzoxadiazol-4-yl)amino)-glucose (2-NBDG), TBA, SRB and trysin were purchased from Sigma Aldrich, United States. Phenazine methosulphate (PMS), nitroblue tetrazolium (NBT), and nicotinamide adenine dinucleotide (NADH) were procured from Himedia labs, India. Hydrogen peroxide was purchased from MP biomedicals, United States. Kits for ALT, AST and ALP, albumin, BUN, CK analysis were purchased from ERBA (Germany). Anti-mouse insulin monoclonal antibody (Thermo Fischer Scientific, United States) and anti-rabbit glucagon receptor N-term polyclonal antibody (Abcam, United Kingdom) were used for the study. Analytical grade chemicals and solvents were used in the current study.

Plant Material

Plant material of P. kurroa was collected from Holi-Nala village, district Chamba (Altitude 2800-3200 m) located in the Dhauladhar range of Himachal Pradesh, India. The plant material was authenticated and deposited in CSIR-IHBT, Palampur herbarium (PLP 11694).

Extraction and Analysis of Samples

One kilogram P. kurroa (rhizomes) was extracted with 80% hydro alcohol and dried using rota evaporator at 45 ± 5°C to yield 21%. PKRE at 30 mg/mL concentration and standards of Mannitol, Picroside I and II at 5 mg/mL were prepared with NMR grade D2O and methanol-d4 (80:20) for NMR spectroscopic analysis. PKRE at 10 mg/mL and picroside I and II at 1mg/mL concentrations were prepared with HPLC grade methanol for LC-MS analysis respectively. The same extract was used for in vitro and in vivo studies.

NMR Spectroscopy

PKRE was dissolved in 1 mL of D2O and methanol-d4 (80:20), 600 µL was transferred to 5 mm NMR tube. NMR spectra
was recorded at 25°C on a 600 MHz Bruker AVANCE III-600 spectrometer (Bruker, Germany) operating at a proton NMR frequency of 600.30 MHz, while carbon at 150.94 MHz. Each 1H-NMR spectrum comprised of 64 scans, in 6 min and 50 s acquisition time with the following parameters: 0.16 Hz/point and relaxation delay (RD) = 5 s. A presaturation sequence was used to suppress the residual H2O/D2O/methanol-d4 signal with low power selective irradiation at the H2O/methanol frequency during the recycle delay. FIDs were fourier transformed with LB H1 (COSY) and heteronuclear spectroscopy (HMBC) based spectra manually phased and baseline was corrected and calibrated at (point calibration curves by plotting the peak areas versus amount scan at wavelength range of 200–490 nm for preparing seven-coupled to a 1998 photodiode array (PDA) detector and used to

H-NMR spectrum comprised of 64 scans, in 6 min 0 Hz. The resulting spectra were manually phased and baseline was corrected and calibrated at 0.0 ppm, using Topspin (version 3.2, Bruker). Homonuclear (COSY) and heteronuclear spectroscopy (HMBC) based spectra were recorded on a 600.30 MHz (1H) and 150.94 MHz (13C) Bruker AVANCE III- spectrometer. The COSY spectra were recorded with 1.0 s relaxation delay, NS = 16, 8012.82 Hz spectral width in both dimensions while the window function was sinebell (SSB = 0). Whereas, HMQC spectra were recorded with 1 s relaxation delay, NS = 64, 6009.61 Hz spectral width in F2 and 36,056.78 Hz in F1 while window function of the HMQC was Q sine (SSB = 2.0). The HMBC spectra were recorded with the 1 s relaxation delay, NS = 64, 9615.38 Hz spectral width in F2 and 36230.55 Hz in F1. Same parameters as the HMQC spectra except for 36,056.78 Hz of spectral width in F2 were applied. The detailed study was done as described earlier (Kumar et al., 2016).

HPTLC Fingerprint of PKRE
The HPTLC fingerprint study was carried out on both normal and reverse phase HPTLC plates according to Kumar et al. (2016) method. PKRE was dissolved in HPLC grade methanol at 10 mg/mL concentration. Four and eight microliters sample was used for HPTLC fingerprint on normal and reverse phase plates (Merck, Switzerland). The solvent system used as mobile phase was ethyl acetate:methanol:acetic acid (16:2:0.2; v:v:v) for normal phase and methanol:water:formic acid (64:36:0.4; v:v:v) for reverse phase (RP-18). The separated bands on the HPTLC plates were viewed using CAMAG Reprostar 3 and recorded at 366 nm after derivatization of NP-HPTLC plate with anisaldehyde sulphuric acid reagent followed by heating at 118°C for 5 min whereas reverse phase HPTLC (RP-18) was viewed and recorded at 254 nm for good separation of picrosides. CAMAG winCats™ 1.4.1 was used for analysis.

UPLC Based Quantification of Picrosides (Picroside I and II) in PKRE
PKRE was dissolved in UPLC grade methanol to yield a 10 mg/mL. The standard stock solutions (Picroside I and II; 1.0 mg/mL) were diluted to a series of suitable concentrations. An aliquot (0.5 μL) of the diluted solutions was injected into a Waters ACQUITY UPLC system (Waters, United Kingdom) with a Waters BEH C18 column, 1.7 μm (2.1 mm × 100 mm) coupled to a 1998 photodiode array (PDA) detector and used to scan at wavelength range of 200–490 nm for preparing seven-point calibration curves by plotting the peak areas versus amount (μg) of each analyte. The mobile phase consists of 0.1% HCOOH in H2O (solvent A) and C2H3N (solvent B) at constant flow-rate of 0.3 mL/min. The linear gradient profile with the following combinations (v/v) of solvent B was used: [t(min), % B]: (0, 5), (0.5, 5), (9, 30), (12, 30), (13, 5), (15, 5) over the 15 min time period. The peaks were recognized by comparing with the reference standards retention time and UV spectra. The 270 nm wavelength was selected for the quantification (Kumar et al., 2016).

In Vivo Experiments
Animals
Young, male Wistar rats (n = 30), 6–8 weeks old, weighing 100–120 g were obtained from experimental animal facility of CSIR-IHBT, Palampur. Animals were acclimatized at standard laboratory conditions (12 h light/dark cycle; 50–60% humidity and 22 ± 2°C temperature) prior to the experiment. Animals were caged in polypropylene cages (n = 6) and fed with pelleted chow and water ad libitum throughout the experiment. The study was approved by Institutional Animal Ethics Committee, CSIR-IHBT, Palampur (Approval number; IAEC/IHBTP3-2013).

Oral Glucose Tolerance Test
In order to evaluate the effect of PKRE on peripheral blood glucose levels, OGTT was performed. Overnight fasted rats were randomly categorized into four groups. Group I; administered with distilled water, Group II; administered with GLB (10 mg/kg body weight dose; Husain et al., 2009; Rehman et al., 2014), while Group III and IV were administered with PKRE at 200 and 400 mg/kg dose, respectively. After 30 min of PKRE or GLB treatment, animals were administered with glucose (2 g/kg dose) orally. Blood glucose levels were measured at varying time points using glucometer (Contour TS, Blood glucose monitoring kit, Bayer Healthcare).

Induction of T1DM and Experimental Design
Rats maintained at fasting condition overnight were injected with a single dose of freshly prepared STZ (Calbiochem, United States; 50 mg/kg dose) in 0.1 M citrate buffer (pH 4.5) intraperitoneally while control animals were injected with citrate buffer alone (Rao et al., 2013; Tripathi and Kohli, 2014; Altinoz et al., 2015). After 1 week, fasting blood glucose was measured using glucometer and animals with hyperglycemic condition at blood glucose levels above 300 mg/dL were taken further for the study. Animals were categorized into five different groups (n = 6) as per experimental regime. Group I; normal control rats, group II; STZ intoxicated rats, group III; diabetic rats administered with standard GLB at the dose of 5 mg/kg daily for 30 days (Cheng et al., 2013; Tripathi et al., 2015; Panjeshahin et al., 2016), group IV; diabetic rats administered with 200 mg/kg dose of PKRE for 30 days and group V; diabetic rats administered with 400 mg/kg dose of PKRE for 30 days. The PKRE doses were finalized on the basis of previous studies by Joy and Kuttan (1999), Husain et al. (2009), Husain et al. (2014). Doses were administered orally using intragastric tube by trained expert in experimental animal facility. Parameters such as body weight and blood glucose were measured at 0, 15th and 30th day of the study. At the end of experiment, animals were
euthanized by CO₂ asphyxiation and dissected to collect blood and vital organs for further investigations.

**Serum Biochemical Estimations**

Blood from each animal was collected prior to dissection via retro-orbital plexus puncture and allowed to clot at room temperature followed by centrifugation at 5000 rpm for 15 min. Serum thus collected was used for biochemical estimations. The levels of ALT, AST, ALP, albumin, CK and BUN were quantitated using commercially available Erba diagnostic kits as per manufacturer’s protocols in Erba XL 200 analyzer (Erba Mannheim, Germany).

**Superoxide Dismutase and Catalase Measurement**

Hepatic and renal tissues were homogenized in 0.1 M ice cold phosphate buffer saline (PBS; pH 7.3) using homogenizer (IKA T-10 basic, Germany), and resultant crude tissue homogenate was centrifuged at 10,000 rpm and 4°C for 20 min. The supernatants obtained were subjected to the estimation of SOD and CAT activities. Results were expressed as units per mg tissue for SOD and CAT as described by Sharma et al. (2016).

**Malondialdehyde Quantification**

Malondialdehyde levels were measured in hepatic and renal tissues using TBA reaction method as described by Patial et al. (2015). Homogenates were prepared from 200 mg tissue in ice cold 0.1M PBS (pH 7.4) and resultant crude homogenates were centrifuged at 10,000 rpm and 4°C for 10 min. The supernatants obtained were used for MDA estimation. For the assay, 1 mL supernatant was maintained at 37°C for 3 h. Reaction was halted by adding 1 mL of 10% TCA (w/v). The contents were allowed to precipitate at room temperature followed by centrifugation at 2000 rpm for 10 min. 1 mL of supernatant was mixed with same volume of 0.67% TBA (w/v), boiled in water bath for 10 min, allowed to cool down and the absorbance was measured at 535 nm, using microplate reader (Synergy; BioTek, United States).

**Histopathology**

Portions of pancreas were fixed in fixative (10% neutral buffered formalin) for histopathological evaluation. Briefly, tissue sections were processed for dehydration, clearing and were embedded in paraffin. Five micrometer sections were obtained using microtome (Finesse me; Thermo Fischer Scientific, United States) and were further deparaffinized, dehydrated, cleared and mounted. Observations for cellular apoptosis were made under Olympus BX53F bright field microscope (Japan).

**Immunohistochemistry**

The 5 µm paraffin embedded pancreas sections were mounted on poly-L-lysine overlayed slides, deparaffinized with xylene and hydrated. Antigen retrieval was achieved by treating sections with sodium citrate buffer. Quenching of endogenous peroxidases was accomplished using BLOXALL blocking solution (ImmPRESS excel staining kit, United States) and exposed sites were blocked by treating sections with horse serum (2.5%) for 20 min in a humidified chamber. Blocked sections were probed overnight with anti-mouse insulin monoclonal antibody (Thermo Fischer Scientific, United States) in dilution of 1:100 and anti-rabbit glucagon receptor N-term polyclonal antibody (Abcam, United Kingdom) in dilution of 1:200. Unbound primary antibodies were removed by PBS wash followed by incubation with HRP-conjugated anti-mouse and anti-rabbit secondary antibodies. The sections were PBS rinsed twice and incubated with diaminobenzidine (DAB) substrate. Developed sections were counterstained with Mayer’s hematoxylin, dehydrated, and then mounted. Examinations were performed under Olympus BX53F bright field microscope for protein expression studies.

**Cell Culture**

The rat insulinoma (Rin5f), rat myoblast (L6) and mouse preadipocyte (3T3L1) cell lines were procured from ATCC, United States. Rin5f cells were grown in RPMI-1640, whereas L6 and 3T3L1 cells were grown in Dulbecco’s Modified Eagle’s medium (DMEM). The culture medium was supplemented with 10% (v/v) heat inactivated FBS (Life Technologies, United States) and 1% (v/v) antibiotic-antimycotic (Life Technologies, United States). Cells were passaged once a week after trypsinization (0.25% trypsin containing 0.2% EDTA) and residual medium was replaced with fresh medium twice a week. Cultures were maintained in CO₂ incubator (Thermo Fischer scientific, United States) with 95% humidity, 5% CO₂ at 37°C. After second passage cells were used for experiments.

**Cell Viability Assay**

Effect of PKRE on the viability of Rin5f, L6 and 3T3L1 cells was determined by SRB assay (Rana et al., 2016). Briefly, confluent monolayer of cells was trypsinized, counted and seeded in a flat bottom 96 well plate at density of 2 × 10⁴ cells/well with culture medium. After 24 h, residual medium was replaced and cells were incubated with new culture medium containing indicated concentrations (viz, 6.25, 12.5, 25, 50, 100, 200, 300, 400, and 500 µg/mL) of PKRE for 48 h. After desired incubation, cells were fixed with 50% (w/v) cold TCA and then plates were kept at 4°C for 1 h. Following 1 h incubation, plates were washed with distilled water four times and allowed to dry at room temperature. Dried plates were stained with 0.4% (w/v) SRB in 1% (v/v) acetic acid for 30 min in dark. Excess and unbound SRB
dye was removed by repeated washing with 1% (v/v) acetic acid and plates were allowed to air dry. Absorbance was measured at 540 nm using microplate reader by solubilizing the bound dye with 10 mM (w/v) Tris HCl.

**In Vitro Insulin Secretion Assay**

Rin5f cells were seeded in 6 well plates at density of 2 × 10⁵ cells/well and allowed to grow for 4–5 days to attained 80–90% confluency. Cells were washed twice with Kreb’s-Ringer bicarbonate buffer (KRB: 115 mM NaCl; 4.7 mM KCl; 1.28 mM CaCl₂; 1.2 mM KH₂PO₄; 1.2 mM MgSO₄; 24 mM NaHCO₃; 10 mM Hepes-free acid, 1 g/L bovine serum albumin, 1.1 mM glucose; pH 7.4) and then pre-incubated in the same buffer for 60 min at 37°C. After incubation, cells were again incubated with fresh KRB buffer containing 25 mM glucose with or without PKRE for 90 min. Post incubation supernatants were collected and insulin secretion was measured using enzyme linked immunosorbent assay (ELISA) performed as per manufacturer’s protocol (Raybiotech, United States).

**Total RNA Isolation and qRT-PCR**

Rin5f cells were seeded in 60 mm cell culture petri dish and allowed to grow up to 80–90% confluency. Cells were incubated in KRB buffer containing 1 mM glucose for 60 min followed by incubation in KRB buffer containing 25 mM glucose with or without PKRE for 90 min. Cells were washed twice with PBS and total RNA was extracted using Qiagen RNeasy mini kit (Qiagen, Germany) following manufacturer’s protocol. RNA was quantified spectrophotometrically using nanodrop (Thermo Fisher Scientific, United States) and purity of RNA was assessed by 260/280 nm absorbance ratio. The integrity of RNA was confirmed by identifying 18 and 28 s RNA bands on 1% agarose gel. One step qRT-PCR was performed using QuantiFast on 1% agarose gel. One step qRT-PCR was performed using QuantiFast SYBR® Green RT-PCR kit (Qiagen, Germany) according to manufacturer’s instructions. Primers were designed using the Primer Express software (Applied Biosystems). For insulin; forward primer, 5′-GCCCAGGCTTTTGTCAAACA-3′ and reverse primer, 5′-AAAAACAGGTTCGCCACACA-3′ and for GAPDH; forward primer, 5′-GGTTGACCTCATGGGCTTACA-3′ and reverse primer, 5′-CAGCAACTGAGGGCCTCTCT-3′ were used. RNA template at concentration of 50 ng was added to the final reaction volume of 10 μL. The PCR reactions were carried out in 96 well plates using the Step one Plus™ Real Time PCR system (Applied Biosystems, United States). GAPDH was used as housekeeping gene to normalize the expression of target gene.

**Fluorescence analysis of 2-[N-(7-Nitrobenz-2-oxa-1,3-dioxol-4-yl)amino]-2-Deoxyglucose (2-NBDG) Uptake by Flow Cytometry in 3T3L1 and L6 Cells**

L6 cells were seeded in 6 well plate at 1 × 10⁵ cells/well and allowed to differentiate into myotubes for 5 days in DMEM supplemented with 2% horse serum (Himedia labs, India). The medium was replaced on alternate days. On the 5th day of seeding the residual medium was replaced with fresh medium with or without PKRE concentrations (6.25, 12.5, 25, 50, and 100 μg/mL) for 24 h. Rosiglitazone (10 μM) was used as positive control. After 24 h, cells were washed twice with KRB buffer (Sigma Aldrich, United States) and then pre-incubated in the same buffer for 30 min. Starved cells were treated with 100 μM 2-NBDG (Fluorescent analog of glucose) for 1 h. After incubation, cells were washed twice with ice cold KRB buffer, trypsinized, resuspended in same buffer and subjected to flow cytometric analysis using AMNIS ImageStream®X MK II Imaging flow cytometer (Merck, United States). Data acquired as 5000 single circular events.

3T3L1 cells were plated at a density of 1 × 10⁶ cells/well in 6 well plate and allow them to grow. At 2 days post confluence stage, cells were induced to differentiate in differentiation medium supplemented with 0.5 mM 3-isobutyl-1-methylxanthine (IBMX), 5 μg/mL insulin, 250 nM Dexamethasone in DMEM and 10% FBS for 48 h. After 2 days, the residual medium was replaced with maintenance medium containing DMEM with 10% FBS and 5 μg/mL insulin. This medium was replaced alternatively up to 9 days. At the end of 9th day, cells were treated with different concentrations of PKRE (6.25, 12.5, 25, 50, and 100 μg/mL) and rosiglitazone at 10 μM concentration for 24 h. On next day, treatment medium was removed and cells were incubated in serum free low glucose medium with or without PKRE for 3 h and further stimulated by adding insulin (5 μg/mL) for 10 min. Stimulated cells were treated with 100 μM 2-NBDG for 1 h, 2-NBDG labeled cells were trypsinized, centrifuged at 125 g for 7 min and the supernatant were decanted whereas cell pellet were suspended in ice cold KRB buffer for data acquisition. Flow cytometry was performed using AMNIS ImageStream®X MK II Imaging flow cytometer (Merck, United States) and data were analyzed using INSPIRE ImageStream system software. Data acquired as 5000 single circular events.

**Statistical Analysis**

Results are expressed as mean ± standard error mean (SEM). Differences were analyzed using GraphPad software by one way ANOVA followed by Dunnett’s multiple comparison range test. Values with p ≤ 0.05 were considered as statistically significant.

**RESULTS**

**Phytochemical Profiling of PKRE**

Thorough phytomolecular characterization of PKRE was carried out by NMR based metabolomic approach and further validated by HPTLC and UPLC. The obtained fingerprint deciphered the identification of major metabolites and revealed the structural and functional diversity of *P. kurroa*. ¹H-NMR spectra of PKRE is divided into three major regions such as amino acids/organic acids/cucurbitacins, sugars/polyols and phenolics/iridoids/flavonoids (Figure 1A). Moreover, the intensity of signals remained very low and overlapped for few regions. Hence, they were difficult to interpret. Therefore, 2D NMR analysis techniques such as ¹H–¹H COSY (homonuclear
correlation), HMQC, and HMBC were employed along with 1D NMR to overcome these problems. The NMR based chemical profiling of PKRE was shown in Figure 1A (Supplementary Figure S1a and Table S1). 1D and 2D NMR based chemical profiling highlighted the presence of 31 metabolites belonging to different classes. The HPTLC fingerprint confirmed the presence of picroside I and II in majority, (Figure 1B).

The UPLC-DAD based analysis also revealed the presence of picrosides. The PDA chromatogram of PKRE and standards was depicted in Figure 1C while the calibration curves of picroside I and II were used for the quantification (Supplementary Figures S1b,c). The total picrosides (picroside I and II) were found 13.87% while the individual picroside I and II were 9.57 and 4.30%, respectively.

PKRE Ameliorate Oral Glucose Tolerance
Amelioration of glucose tolerance obtained from integrated glucose to insulin response that maintains the normal glycemic conditions (Anhê et al., 2015). OGTT was performed to check the hypoglycemic potential of bioactive molecules. Results suggest that, initial blood glucose was 61.4 mg/dL and 55.6 mg/dL in PKRE assigned groups in comparison to normal control (75.4 mg/dL) and GLB assigned group (60 mg/dL). After 30 min, blood glucose levels reached at peak with 89.2 mg/dL at 200 mg/kg and 94.6 mg/dL at 400 mg/kg dose group in PKRE treated groups compared to normal control group (91.4 mg/dL) and GLB treated group (90.8 mg/dL). Further, improvement was observed in the glucose tolerance and after 90 min significant response was recorded. PKRE significantly lowered the blood glucose to 70 mg/dL (p < 0.05) at 200 mg/kg and 59.6 mg/dL (p < 0.001) at 400 mg/kg dose when compared with normal control group (91.2 mg/dL) after 90 min. However, GLB treatment also effectively lowered the blood glucose to 67.2 mg/dL, (p < 0.01) after 90 min of glucose loading (Figure 2A).

PKRE Improves Hyperglycemia and Weight Loss toward Normalcy
Impaired insulin secretion results from destruction of pancreatic β-cells that disturbs glucose homeostasis (Cerf, 2013). Increased blood glucose leads to serious metabolic complications and oxidative stress leading to cellular damages (Asmat et al., 2016). PKRE treatment for 15 days, significantly lowered the blood glucose levels to 190.8 mg/dL and 246 mg/dL (p < 0.01) in 200 and 400 mg/kg dose group respectively when compared to STZ intoxicated group (281.8 mg/dL, p < 0.001). After 30 days, PKRE treatment significantly (p < 0.0001) lowered the blood glucose levels to 136 mg/dL and 166 mg/dL in 200 and 400 mg/kg dose treated group in contrast to STZ intoxicated group (239.4, p < 0.01). Similar results were observed for GLB treated group (127.2 mg/dL) after 30 days. Blood glucose levels of STZ intoxicated group remained higher (239.4 mg/dL) till the end of experiment (Figure 2B).

STZ, GLB, and PKRE treatment doesn’t exert much effect on the body weight of control and diabetic rats throughout the experiment. At 0 day, 132 g body weight was recorded for both PKRE treatment groups, whereas 131.4, 123.5, and 126 g were recorded for normal control, STZ intoxicated and GLB treatment groups respectively. Further, increase in the body weight of animals was observed after treatments. After 15 days, it was recorded to 142.4 g in 200 mg/kg and 132.8 g in 400 mg/kg PKRE treated groups as compared to control group (163.8 g). The increase was recorded to 132.2 g for STZ intoxicated group and 129.4 for GLB treated group after 15 days. Further it was found that after 30 days, 147.4 and 152 g weight was recorded for 200 and 400 mg/kg PKRE treated groups respectively. Whereas, 152.25 g (p < 0.01) was recorded for STZ intoxicated group and 141.2 g for GLB treated group in contrast to control group (170 g) (Figure 2C).
PKRE Treatment Maintained the Normal Levels of ALT, AST and ALP

An elevated serum enzymes ALT, AST and ALP are the markers of hepatic damage and indicates the hepatotoxic effect of STZ (Srinivasan et al., 2014). ALT catalyzes the alanine cycle and its enhanced level suggests the existence of hepatic disorders. In the present study, increased level of ALT (350 IU/L, \( p < 0.001 \)) was measured in STZ intoxicated group compared to its control group (147.3 IU/L). On the other hand PKRE treatment effectively lowered the ALT levels to 60.0 IU/L (\( p < 0.05 \)) and 110.1 IU/L in 200 and 400 mg/kg dose groups respectively. However, GLB treatment also efficiently lowered the ALT level to (67.8 IU/L, \( p < 0.05 \)) (Figure 3A). AST is one of the important enzyme in amino acid metabolism and also a marker of hepatic health. In STZ intoxicated group, the serum level of AST decreased significantly (6.6 IU/L, \( p < 0.01 \)) compared to normal control group (28.9 IU/L), but PKRE administration restored the AST levels to 38.3 IU/L in 200 mg/kg and 57.1 IU/L (\( p < 0.001 \)) in 400 mg/kg treated groups. Further, GLB treatment also increased the AST to 38.6 IU/L (Figure 3B). On the other hand, serum ALP levels tend to increase due to its increased cellular synthesis during biliary stress. Leakage of ALP in bloodstream increases in hepatic abnormalities (Ashraf et al., 2013). The levels of ALP increased to 212.7 IU/L in STZ intoxicated group in comparison to control group 103.0 IU/L, whereas PKRE treatment lowered the serum ALP levels to 155.8 IU/L in 200 mg/kg and 175.8 IU/L in 400 mg/kg dose treated groups. GLB also effectively reduced the ALP level to 113.8 IU/L (Figure 3C).

PKRE Treatment Improves Serum Albumin, BUN and CK Levels

Serum albumin, BUN and CK levels are significantly altered in hyperglycemic situation as a result of renal damage, protein catabolism, extracellular dehydration and lack of glucose metabolism (Go et al., 2015). Serum albumin is the most abundant plasma protein imparting several physiological attributes including antioxidant property. Hyperglycemia induces the formation of advanced glycation end products (AGEs) that binds with albumin protein and significantly reduces its level (Raghav and Ahmad, 2014). The STZ intoxicated group showed reduced levels of serum albumin (1.8 g/dL) compared to normal control group (1.8 g/dL). PKRE treatment effectively enhanced the serum albumin level to 2.2 g/dL (\( p < 0.05 \)), 2.5 g/dL at 200 and 400 mg/kg dose treated groups whereas GLB also increased albumin level to 2.1 g/dL (Figure 3D). BUN and CK are the nitrogenous wastes and their increased serum levels are the markers of renal damage. Hyperglycemia mediated reactive oxygen species (ROS) affects glomerular filtration rate, which contributes in the accumulation of nitrogenous wastes in the blood stream (Altinoz et al., 2015). BUN was found increased in STZ intoxicated group (52.4 g/dL) as compared to control group (22.5 g/dL). PKRE treatment significantly (\( p < 0.0001 \)) reduced the levels to 25 g/dL in 200 mg/kg and 24.4 g/dL in 400 mg/kg dose treated groups, GLB also brought down the BUN level to 27.9 g/dL (Figure 3E). CK levels were also found elevated in diabetic rats (2.5 U/L) compared to control group (1.5 U/L). PKRE significantly decreased the CK level to 0.66 U/L.
PKRE改善肝肾功能并正常化抗氧化参数

高血糖诱导的氧化应激结果在降低抗氧化水平并增加自由基（Florence et al., 2014）。这些自由基损害细胞成分，并触发特定的信号通路，导致分泌炎性细胞因子（Naowaboot et al., 2009）。在本研究中，肝和肾组织中测量了SOD、CAT和MDA。SOD通过将超氧阴离子转化为水和氧（Ren et al., 2015）来减少细胞内的超氧阴离子浓度。STZ处理组测量的SOD水平（53.5 U, p < 0.05）与对照组（67.7 U）相比显著降低。PKRE治疗组显著提高了SOD水平（87.4 U和73 U）。

MDA，一种高度反应性物种，和细胞氧化压力的标记，来自不饱和脂肪酸（PUFA）的脂质过氧化反应（Adefegha et al., 2014）。STZ处理组的MDA水平（1.9 nM）显著升高，而对照组（1.6 nM）水平较低。

PKRE保护胰岛β细胞免受STZ诱导的细胞损伤并增强胰岛素表达

胰岛素缺乏是STZ诱导的高血糖的后果，是由于感染的自身免疫性淋巴细胞的胰岛β细胞的破坏（Zhao et al., 2015）。这导致血糖升高，进而导致代谢性变化。据报道，β细胞的再生能够逆转STZ诱导的糖尿病（Soltani et al., 2011）。STZ诱导的β细胞的损坏和脂质过氧化反应的增加模仿了T1DM的病理生理学。组织病理学分析表明，STZ诱导的β细胞的损坏和脂质过氧化反应的增加模仿了T1DM的病理生理学。组织病理学分析表明，STZ诱导的β细胞的损坏和脂质过氧化反应的增加模仿了T1DM的病理生理学。
**FIGURE 4** | PKRE treatment ameliorated the activity of SOD (A) catalase (B) and simultaneously inhibited the MDA production (C) in hepatic and renal tissue homogenates of different experimental groups. Data are expressed as mean ± SEM (n = 6). ∗p < 0.05, **p < 0.01 indicate significant difference in group compared to normal control group.

**FIGURE 5** | PKRE treatment significantly protects β-cells from STZ evoked cellular damage and enhanced the insulin production in diabetic rats. (A) Representative histopathological photographs showing STZ evoked β-cells alterations in different experimental groups detected by H and E staining. (a) Normal group, (b) Streptozotocin group, (c) Glib control, (d) PKRE 200 mg/kg and (e) PKRE 400 mg/kg. (B) Microscopic images of TUNNEL assay in various experimental groups for apoptosis assessment revealed the protective effect of PKRE in diabetic rats. (a) Normal group, (b) Streptozotocin group, (c) Glib control, (d) PKRE 200 mg/kg and (e) PKRE 400 mg/kg. (C) Representative Immunohistochemistry images of insulin expression in different experimental groups. PKRE treatment increased the insulin expression in pancreatic β-cells was evaluated through IHC. The normal control group was found to have highest expression of insulin upto 40%, however, it was found drastically less in STZ intoxicated group (6.1%, p < 0.0001). In contrast, Glib (10.6%, p < 0.001) and PKRE treated groups showed the enhanced...
insulin expression in β-cells as compared to STZ intoxicated group. PKRE significantly enhances the insulin expression up to 13.9% ($p < 0.001$) and 16.4% ($p < 0.01$) at 200 and 400 mg/kg doses respectively (Figure 5C).

**PKRE Treatment Inhibits the Expression of Hepatic and Renal Gcgr**

Evidence demonstrates that inhibition of in vivo glucagon signaling leads to hypoglycemia. Glucagon binds to its receptors, resulting in glucose release with enhanced gluconeogenesis and glycogenolysis. Gcgr are widely distributed in hepatic, renal and other tissues (Jiang and Zhang, 2003). IHC results revealed that hepatic Gcgr expression increases significantly ($p < 0.001$) to 26.5% in STZ intoxicated group when compared with normal control (9.05%). PKRE on the other hand effectively brought down the Gcgr expression to 16.3 and 9.3% at 200 and 400 mg/kg doses respectively. GLB treatment also brought down the Gcgr expression to 16.4% ($p < 0.001$) at 200 and 400 mg/kg dose treatment group. Further, GLB treatment also inhibited the Gcgr expression to 12.6% ($p < 0.001$) at 200 and 400 mg/kg dose treatment group. Further, GLB treatment also inhibited the Gcgr expression to 12.4% ($p < 0.01$) (Figure 6B).

**PKRE Proliferates Rin5f cells with Enhanced Insulin Gene Expression and Insulin Secretion**

Cell viability measures the cytocompatibility as well as protective efficacy of test molecules. It enables to find the safe dose of a test component with minimal toxicity. Cellular viability on Rin5f cells was determined by SRB assay and we found that PKRE treatment significantly supports their proliferation in concentration dependent manner after 48 h exposure (Figure 7A). Further, non-lethal concentrations of PKRE significantly enhanced the insulin gene expression. Reduction in insulin levels results from destruction of β-cells in T1DM, which is directly linked with hyperglycemia and associated complications (Subash-Babu et al., 2015). PKRE and GLB significantly increases the transcript levels post 90 min treatment. It was found that PKRE significantly increases the transcript levels to 3.1 fold ($p < 0.01$) at 6.25 µg/mL as compared to high glucose control (1.16 fold) and GLB control (2.3 fold, $p < 0.01$). Further, insulin gene expression was upregulated to 3.4 ($p < 0.01$), 4 ($p < 0.001$), and 5.2 fold ($p < 0.001$) at 12.5, 25, and 50 µg/mL of PKRE treatment. However, highest transcript levels were recorded up to 74.7 fold ($p < 0.0001$) at 100 µg/mL (Figure 7B).

In order to further validate our above findings, cells were treated with 6.25, 12.5, 25, 50, and 100 µg/mL of PKRE for 90 min and the insulin secreted by the Rin5f cells were measured by ELISA. The insulin secretion was 2.1 µIU/mL ($p < 0.01$) when cells were treated at 6.25 µg/mL compared with high glucose control up to 1.1 µIU/mL. This was further found to be elevated in concentration dependent manner and could recorded as high as up to 3.68 µIU/mL ($p < 0.05$) at 100 µg/mL compared to GLB treated cells which recorded up to 2.8 µIU/mL (Figure 7C).

**PKRE Protects Cellular Viability and Enhances the Glucose Uptake by L6 and 3T3L1 Cells**

Cytocompatibility of PKRE with L6 cells revealed that PKRE does not affect the cellular viability and in contrast supported the cellular proliferation up to 4.2% at 100 µg/mL concentration. However, we noticed decrease in cellular viability up to 2.3% with further increase in PKRE concentrations after 48 h exposure (Figure 8A). In 3T3L1 cells, we found that PKRE maintained the cellular integrity up to 95.5% ($p < 0.001$) at 100 µg/mL and further 79.9% ($p < 0.001$) viability was recorded at 500 µg/mL (Figure 8B). Afterward, sub-lethal concentration of PKRE were evaluated for their 2-NBDG uptake as a measure of glucose uptake potential in 3T3L1 and L6 cells. Insulin stimulated uptake in 3T3L1 and L6 cells significantly increases with PKRE treatment as compared to rosiglitazone. Rosiglitazone, a PPAR-γ agonist is an effective antidiabetic drug which modulates the secretory function of adipocytes and improves insulin sensitivity (Shyny et al., 2014). PKRE and rosiglitazone significantly enhances the glucose consumption in differentiated L6 cells. Mean
bioavailability of picroside I, II and apocynin, was detected in *P. kurroa* iridoid enriched fraction of Zahiruddin et al. (2016) reported the pharmacokinetics of in vitro validated at cellular and molecular level to ameliorate insulin insufficiency in vivo taken further systematically to determine the efficacy of PKRE doses in the glucose loaded rats. These observations were blood glucose level toward normalcy at both the treatment for OGTT, which revealed that PKRE effectively lowered the chemically characterized PKRE extract was further evaluated major iridoid glycosides using HPTLC and UPLC DAD. This approaches and then quantified and validated the presence of (\( p < 0.001 \)) compared to control. The mean fluorescence levels in cells treated with 6.25, 12.5, 25, 50, and 100 \( \mu \text{g/mL} \) of PKRE were found increasing with concentration and recorded as high as upto 27.7% (\( p < 0.001 \)) at 100 \( \mu \text{g/mL} \) (Figure 8D).

**DISCUSSION**

In the present study, phytomolecular profiling of PKRE revealed the presence of diverse metabolites belonging to different classes. We extensively performed the chemical characterization of the rhizome extract by 1D and 2D \( ^1\text{H}-\text{NMR} \) metabolomic approaches and then quantified and validated the presence of major iridoid glycosides using HPTLC and UPLC DAD. This chemically characterized PKRE extract was further evaluated for OGTT, which revealed that PKRE effectively lowered the blood glucose level toward normalcy at both the treatment doses in the glucose loaded rats. These observations were taken further systematically to determine the efficacy of PKRE to ameliorate insulin insufficiency *in vivo* which was further validated at cellular and molecular level *in vitro*. Recently, Zahiruddin et al. (2016) reported the pharmacokinetics of iridoid enriched fraction of *P. kurroa* rhizome. Significant bioavailability of picroside I, II and apocynin, was detected in plasma and negligible amount was reported in urine reflecting its proper systemic consumption and metabolism. This provides the molecular basis for antihyperglycemic attributes of *P. kurroa in vivo*.

Pathophysiology due to T-cell mediated destruction of insulin producing β-cells leads to insulin shortage in the vital organs resulting in life threatening macro and micro vascular complications. Autoreactive lymphocytes, macrophages and cytokines such as IFN-γ and TNF-α also play prominent roles in pancreatic β-cells destruction (Padgett et al., 2013). In the background of above perspectives, the present study investigated the effect of PKRE on β-cell mass regeneration in STZ intoxicated rats and its effect on proliferation and insulin gene expression in Rin5f cells. Our data suggests that PKRE lowered the fasting blood glucose levels at both the treatment doses compared to STZ intoxicated rats. Blood glucose clearance was due to PKRE stimulatory effects on pancreas, which could lead to enhanced blood insulin levels and glt 4 translocation in the peripheral tissues. Increased insulin secretion is either due to the regeneration or by stimulating the existing pancreatic β-cells (Ong et al., 2011; Soltani et al., 2011). Expression analysis by IHC showed significant (\( p \leq 0.05 \)) increase in the insulin secretion by regenerating β-cells. Remarkably, PKRE also reduced β-cells apoptosis and reversed the insulin insufficiency, whereas STZ intoxicated group was marked by the presence of apoptotic bodies. Interestingly, proliferation of insulin producing Rin5f cells was observed in concentration dependent manner when cytocompatibility of PKRE was evaluated. Transcript quantification and ELISA demonstrated that PKRE enhanced the insulin secretion by elevating the insulin transcript levels to levels as high as upto 74.7 fold.
It has been reported that Gcgr−/− mice significantly reverses T1DM complications and also suggests that Gcgr antagonists as well as glucagon suppressors can be used as possible therapeutic agents to maintain blood glucose homeostasis (Lee et al., 2011). Our IHC results revealed the increased expression of Gcgr in STZ intoxicated group compared to normal untreated group and also throws light on the fact that PKRE effectively inhibits the hepatic and renal Gcgr expression, which further supported its anti-hyperglycemic property.

Altered levels of ALT, AST and ALP in the blood stream are the markers of hepatic dysfunction or integrity (Adefegha et al., 2014; Al-Malki and El Rabey, 2014). Imbalance in their levels is reported to be associated with the complications due to hyperglycemia (Arunachalam and Parimelazhagan, 2013) which was evident in STZ intoxicated group. However, PKRE administration effectively brought down these markers to their normal levels when compared with normal control animals. On the other hand, hyperglycemia significantly altered the levels of serum albumin, BUN and CK in STZ intoxicated group. Alterations in their levels are directly related to renal damage, which might be due to decreased protein synthesis and enhanced tissue proteolysis in diabetic rats (Nabi et al., 2013; Tzeng et al., 2013). Further we found that, PKRE increases the serum albumin while decreasing the BUN and CK levels toward normalcy in diabetic rats, indicating lower tissue proteolysis and normalized glomuerl filtration. These results suggested that PKRE lowered the blood glucose and oxidative stress, which supported its protective effect against hyperglycemia mediated hepatic and renal damage.

Streptozotocin induced free radicals impaired the activity of antioxidant enzymes (SOD and CAT) with concurrently enhanced lipid peroxidation in STZ intoxicated rats compared to normal (Arunachalam and Parimelazhagan, 2013; Efiong et al., 2013). PKRE treatment activated SOD and CAT in both hepatic and renal tissues, respectively. Further, enhanced oxidative stress resulted in increased serum lipid peroxides due to oxidation of PUFA (Tangvarasittichai, 2015) as evident from STZ intoxication. PKRE also lowered the MDA levels to normal and restores the cellular oxidative status. Improvement in the oxidative stress may directly or indirectly protects β-cells from ROS mediated destruction, though the results are statistically not significant.

Skeletal muscles are considered as major site for the insulin stimulated glucose uptake as well as oxidation. Defective insulin sensitivity in skeletal muscle and adipose tissues leads to hyperglycemia (Ueda et al., 2012) Insulin stimulated glucose uptake is accompanied with glut 4 transporters, predominantly distributed in muscles and adipose tissues (Uldry and Thorens, 2004). In order to evaluate the effect of PKRE on insulin stimulated glucose uptake, we performed glucose uptake assays in L6 and 3T3L1 cells and compared their response with that of rosiglitazone. Our observations revealed that PKRE in varying concentrations significantly (p ≤ 0.05) enhanced the insulin

**FIGURE 8** | (A) Cytocompatibility ability of different PKRE concentrations (6.25, 12.5, 25, 50, 100, 200, 300, 400, and 500 µg/mL) on the L6 cells viability was achieved by SRB assay after 48 h incubation. (B) PKRE treatment significantly also protected the viability of 3T3L1 cells after 48 h treatment, measured by SRB assay. (C) 2-NBDG measurement in L6 cells by flow cytometry. PKRE treatment significantly enhanced 2-NBDG uptake in differentiated myotubes. Mean fluorescence intensities were measured using Image Stream X MKII, Merck flow cytometer by acquiring single circular events. (D) Insulin stimulated 2-NBDG uptake was enhanced by PKRE treatment in differentiated 3T3L1 cells. Mean fluorescence intensities were measured using Image Stream X MKII, Merck flow cytometer by acquiring single circular events. Experiments were done in triplicate on twice (n = 6). Data are expressed as mean ± SEM (n = 6; triplet of two separate experiments). ∗p < 0.05, ∗∗p < 0.01, ∗∗∗p < 0.001, and ∗∗∗∗p < 0.0001 indicate significant difference in groups compared to normal untreated control group.
stimulated glucose uptake in concentration dependent manner and supported the in vivo findings of enhanced insulin secretion from regenerated pancreatic β-cells.

Present study has provided insights for the presence of more than 31 major metabolites including iridoid glycosides in the PKRE. It exhibited β-cell regeneration potential as evidenced by IHC, which further complimented the enhanced insulin production. Additionally, results were validated in vitro on insulin producing Rin5f cells and it was found that PKRE proliferates β-cells at all treatment concentrations. It also results in elevated insulin gene expression and insulin secretion by Rin5f cells. Furthermore, PKRE was also found to enhance the insulin stimulated glucose uptake in 3T3L1 and L6 cells.

CONCLUSION

In summary, present study demonstrated that metabolomically characterized hydro alcoholic extract of P. kurroa rhizome exerted strong β-cell regeneration potential along with enhanced insulin expression, antihyperglycemic effects and improves the hepatic and renal functions. Furthermore, PKRE possesses increased insulin stimulated glucose uptake potential but the underlying mechanism still needs to be explored in order to elucidate the insulin signaling in skeletal muscles and adipose tissues.

REFERENCES

Adefegha, S. A., Oboh, G., Adefegha, O. M., Boligon, A. A., and Athayde, M. L. (2014). Anti-hyperglycemic, hypolipidemic, hepatoprotective and antioxidative effects of dietary clove (Syzygium aromaticum) bud powder in a high-fat diet/streptozotocin-induced diabetic rats model. J. Sci. Food Agric. 94, 2726–2737. doi: 10.1002/jsfa.6617

Al-Malki, A. L., and El Rabey, H. A. (2014). The antidiabetic effect of low doses of Moringa oleifera Lam. seeds on streptozotocin induced diabetes and diabetic nephropathy in male rats. Biomed. Res. Int. 2015:381040. doi: 10.1155/2015/381040

Altinoz, E., Oner, Z., Elbe, H., Cigremis, Y., and Turkoz, Y. (2015). Protective effects of saffron (its active constituent, crocin) on nephropathy in streptozotocin-induced diabetic rats. Hum. Exp. Toxicol. 34, 127–134. doi: 10.1177/0960327114538989

Anbé, F., Roy, D., Pilon, G., Dudonné, S., Matamoros, S., Varin, T. V., et al. (2015). A polyphenol-rich cranberry extract protects from diet-induced obesity, insulin resistance and intestinal inflammation in association with increased Akkermansia spp. population in the gut microbiota of mice. Gut 64, 872–883. doi: 10.1136/gutjnl-2014-307142

Arunachalam, K., and Parimelazhagan, T. (2013). Antidiabetic activity of Ficus amplissima Smith. bark extract in streptozotocin induced diabetic rats. J. Ethnopharmacol. 147, 302–310. doi: 10.1016/j.jep.2013.03.004

Ashraf, H., Heidari, R., Nejati, V., and Ilkhanipoor, M. (2013). Effects of aqueous extract of Berberis integerrima root on some physiological parameters in streptozotocin-induced diabetic rats. Iran J. Pharm. Res. 12, 423–434.

Asmat, U., Abad, K., and Ismail, K. (2016). Diabetes mellitus and oxidative stress - A concise review. Saudi Pharm. J. 24, 547–553. doi: 10.1016/j.sjps.2015.03.013

Battaglia, M., Mark, A., and Atkinson, M. A. (2015). The streetlight effect on type 1 diabetes. Diabetes Metab. Res. Rev. 64, 1081–1090. doi: 10.2337/db14-1208

Cerf, M. E. (2013). Beta cell dysfunction and insulin resistance. Front. Endocrinol. 4:37. doi: 10.3389/fendo.2013.00037

Chen, Y., Wu, Y., Gan, X., Liu, K., Lv, X., Shen, H., et al. (2016). Iridoid glycoside from Cornus officinalis ameliorated diabetes mellitus-induced testicular damage in male rats: involvement of suppression of the AGEs/RAGE/p38 MAPK signaling pathway. J. Ethnopharmacol. 194, 850–860. doi: 10.1016/j.jep.2016.10.079

Cheng, D., Liang, B., and Li, Y. (2013). Antihyperglycemic effect of Ginkgo biloba extract in streptozotocin-induced diabetic rats. BioMed Res. Int. 2013, 7. doi: 10.1155/2013/162724

Coelho, G. H., Martins, V. S., do Amaral, L. V., Novaes, R. D., Sarandy, M. M., and Gonçalves, R. V. (2016). Applicability of isolates and fractions of plant extracts in murine models in type II diabetes: a systematic review. Evid. Based Complement. Altern. Med. 2016, 25. doi: 10.1155/2016/3537163

Devi, S., Kumar, D., and Kumar, M. (2016). Ethnobotanical values of antidiabetic plants of M.P. region India. J. Med. Plants Stud. 4, 26–28.

Dwivedi, Y., Rastogi, R., Garg, N. K., and Dhawan, B. N. (1992). Picroliv and its components kutkoside and picroside I protect liver against galactosamine-induced damage in rats. Pharmacol. Toxicol. 71, 383–387. doi: 10.1111/j.1600-0773.1992.tb0566.x

Efiong, E. E., Igule, G. O., Mbeje, B. I. A., Out, E. A., and Ebong, P. E. (2013). Hepatoprotective and anti-diabetic effect of combined extracts of Moringa oleifera and Vernonia amygdalina in streptozotocin-induced diabetic albino Wistar rats. J. Diabetes Endocrinol. 4, 45–50. doi: 10.5897/JIDE2013.0069

Florence, N. T., Benoit, M. Z., Jonas, K., Alexandra, T., Désiré, D. D., Pierre, K., et al. (2014). Antidiabetic and antioxidant effects of Annona muricata (Annonaceae), aqueous extract on streptozotocin-induced diabetic rats. J. Ethnopharmacol. 151, 784–790. doi: 10.1016/j.jep.2013.09.021

Go, H. K., Rahman, M. M., Kim, G. B., Na, C. S., Song, C. H., Kim, J. S., et al. (2015). Antidiabetic effects of yam (Dioscorea batatas) and its active constituent, allantoin, in a rat model of streptozotocin-induced diabetes. Nutrients 7, 8532–8544. doi: 10.3390/nu7105411

Husain, G. M., Rai, R., Rai, G., Singh, H. B., Thakur, A. K., and Kumar, V. (2014). Potential mechanism of anti-diabetic activity of Picrorhiza kurroa. TANG 4, 27. doi: 10.5667/tang.2014.0013

Husain, G. M., Singh, P. N., and Kumar, V. (2009). Antidiabetic activity of standardized extract of Picrorhiza kurroa in rat model of NIDDM. Drug Discov. Ther. 3, 88–92.

AUTHOR CONTRIBUTIONS

SK performed majority of the experiments. VP, SoS, SuS, KP, and DK assisted in performing different experiments. SK, VP, and DK analyzed and interpreted the data as well as drafted the manuscript. YP conceptualized, supervised the study and edited the manuscript. All authors contributed in the finalization of manuscript.

ACKNOWLEDGMENTS

The authors are thankful to the Director, CSIR-IHBT for providing necessary facilities. The authors are also thankful and acknowledge Dr. Bikram Singh, PI, CSC0130. We are thankful to the CSIR network project “NaPaHa” (CSC0130) for financial support. We are also thankful to Ms. Ridhi Sood and Ms. Kiran Rawat for their help in experimentation and manuscript editing. SK is grateful to University Grants Commission (UGC, New Delhi) for providing Rajiv Gandhi National Fellowship.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fphar.2017.00537/full#supplementary-material
Jiang, G., and Zhang, B. B. (2003). Glucagon and regulation of glucose metabolism. *Am. J. Physiol. Endocrinol. Metab.* 284, E671–E678. doi: 10.1152/ajpendo.00492.2002

Joy, K. L., and Kuttan, R. (1999). Anti-diabetic activity of *Picrorhiza kurroa* extract. *J. Ethnopharmacol.* 67, 143–148. doi: 10.1016/S0378-7741(98)00243-8

Kakkle, K., Soldatou, A., Karachalou, F., and Karavanaki, K. (2015). Associated autoimmune diseases in children and adolescents with type 1 diabetes mellitus (T1DM). *Autoimmun. Rev.* 14, 781–797. doi: 10.1016/j.autrev.2015.05.002

Klinke, D. J. (2008). Extent of beta cell destruction is important but insufficient to predict the onset of type 1 diabetes mellitus. *PLoS ONE* 3:e1374. doi: 10.1371/journal.pone.0001374

Kumar, D., Kumar, R., Singh, B., and Ahuja, P. S. (2016). Comprehensive chemical profiling of *Picrorhiza kurroa* Royle Ex Benth. using NMR, HPTLC and LC-MS/MS techniques. *Comb. Chem. High Throughput Screen.* 19, 200–215. doi: 10.2174/18662073116610110492538

Kumar, K. M. (2015). Incidence trends for childhood type 1 diabetes in India. *Indian J. Endocrinol. Metab.* 19, S34–S35. doi: 10.4103/2230-8210.155378

Lee, Y., Wang, M. Y., Du, X. Q., Charron, M. J., and Unger, R. H. (2011). Glucagon receptor knockout prevents insulin-deficient type 1 diabetes in mice. *Diabetes Metab. Res. Rev.* 60, 391–397. doi: 10.2337/db10-0426

Luper, S. (1998). A review of plants used in the treatment of liver disease: part 1. *Altern. Med. Rev.* 3, 410–421. doi: 10.2337/db10-0426

Mahajan, R., Kapoor, N., and Singh, I. (2016). Somatic embryogenesis and callus proliferation in *Picrorhiza kurroa* Royle ex Benth. *J. Exp. Biol. Agric.* Sci. 4, 201–209.

Meier, C., Schwartz, A. V., Egget, A., and Lecka-Czernik, B. (2015). Effect of diabetes drugs on the skeleton. *Bone* 82, 93–100. doi: 10.1016/j.bone.2015.04.026

Nabi, S. A., Kasetti, R. B., Sirasanagadla, S., Tilak, T. K., Kumar, M. V., and Rao, C. A. (2013). Antidiabetic and antihyperlipidemic activity of Piper longum root aqueous extract in STZ induced diabetic rats. *BMC Complement. Altern. Med.* 13:37. doi: 10.1186/1472-6882-13-37

Naowaboot, J., Pannangpetch, P., Kukongviriyapan, V., Kongyingyoes, B., and Shitiz, K., Pandit, S., Chauhan, R. S., and Sood, H. (2015). Picroside contents in the rhizomes of *Picrorhiza kurroa* Royle ex Benth. traded for the herbal drugs in the market of North India. *Int. J. Med. Arom. Plants* 3, 226–233.

Shin, G. L., Kavitha, S., Indu, S., Arya, A. D., Anusree, S. S., Vneetha, V. P., et al. (2014). Chebulagic acid from *Terminalia chebula* enhances insulin mediated glucose uptake in ST31 adipocytes via PPARγ signaling pathway. *Biofactors* 40, 665–667. doi: 10.1002/biof.1193

Soltani, N., Qiu, H., Aleksic, M., Glinka, Y., Zhao, F., Liu, R., et al. (2011). GABA exerts protective and regenerative effects on islet beta cells and reverses diabetes. *Proc. Natl. Acad. Sci. U.S.A.* 108, 11692–11697. doi: 10.1073/pnas.110715108

Srinivasan, S., Satish, G., Jayanthi, M., Muthukumarana, J., and Ramachandran, V. (2014). Ameliorating effect of genonol on hyperglycemia by attenuating the key enzymes of glucose metabolism in streptozotocin-induced diabetic rats. *Mol. Cell Biochem.* 385, 159–168. doi: 10.1007/s10110-013-1824-2

Subash-Babu, P., Ignacimuthu, S., and Alishawi, A. A. (2015). Nymphaly becomes glucose-stimulated insulin secretion by Rin 5F cells and Glut 4 mediated insulin sensitization in type 2 diabetic rat liver. *Chem. Biol. Interact.* 226, 72–81. doi: 10.1016/j.cbi.2014.12.011

Tangvasisitchai, S. (2015). Oxidative stress, insulin resistance, dyslipidemia and type 2 diabetes mellitus. *World J. Diabetes* 6, 456–480. doi: 10.4239/wjd.v6.i3.456

Tanuja, T., and Preeti, C. (2016). Efficient in vitro establishment and multiplication of *Picrorhiza kurroa* Royle Ex Benth through in vitro raised seedlings. *Res. Crop.* 17, 800–807. doi: 10.5958/2348-7542.2016.00135.2

Tripathi, A. K., and Kohli, S. (2014). Pharmacognostical standardization and antidiabetic activity of *Syzygium cumini* (Linn.) barks (Myrtaceae) on streptozotocin-induced diabetic rats. *J. Complement. Integr. Med.* 11, 71–81. doi: 10.5137/cim.2014.0011

Tripathi, Y. C., Shukla, P., and Tewari, D. (2015). Phytochemical evaluation and antihyperglycemic effects of *Elaeocarpus ganitrus* root (Rudraksha) in streptozotocin-induced diabetes. *Int. J. Pharm. Pharm. Sci.* 7, 280–283.

Tzeng, T. F., Liu, S. S., Chang, C. J., and Liu, I. M. (2013). The ethanol extract of *Zingiber zerumbet* attenuates streptozotocin-induced diabetic nephropathy in rats. *Evid. Based Complement. Alternat. Med.* 2013:340645. doi: 10.1155/2013/340645

Ueda, M., Hayashibara, K., and Ashida, H. (2012). Pharmacokinetics and comparative metabolic profiling of iridoid glycosides from Picrorhiza kurroa rhizomes of *Picrorhiza kurroa* derivates, Picroliv, attenuates the development of dextran-sulphate-sodium-induced colitis in mice. *Mediators Inflamm.* 2012:751629. doi: 10.1155/2012/751629

Uldry, M., and Thorens, B. (2004). The SLC2 family of facilitated hexose and polyol transporters. *Pflugers Arch.* 447, 480–489. doi: 10.1007/s00424-003-1085-0

Zahruiddin, S., Khan, W., Nehra, R., Alam, M. J., Malik, M. N., Parveen, R., et al. (2016). Pharmacokinetics and comparative metabolic profiling of iridoid enriched fraction of *Picrorhiza kurroa*-An Ayurvedic Herb. *J. Ethnopharmacol.* 179, 157–164. doi: 10.1016/j.jep.2016.07.072

Zhang, D. K., Yu, J. J., Li, Y. M., Wei, L. N., Yu, Y., Feng, Y. H., et al. (2012). A *Picrorhiza kurroa* derivative, Picroliv, attenuates the development of dextran-sulfate-sodium-induced colitis in mice. *Mediators Inflamm.* 2012:751629. doi: 10.1155/2012/751629

Zhao, Y., Scott, N. A., Fynch, S., Elkerbout, L., Wong, W. W., Mason, K. D., et al. (2015). Autoimmune T cells induce necrosis and not BCL-2-regulated or death
β-Cell Regeneration Potential of PKRE receptor-mediated apoptosis or RIPK3-dependent necroptosis of transplanted islets in a mouse model of type 1 diabetes. *Diabetologia* 58, 140–148. doi: 10.1007/s00125-014-3407-5

**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.