Visfatin Reduces Gap Junction Mediated Cell-to-Cell Communication in Proximal Tubule-Derived Epithelial Cells

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Visfatin • Gap Junctions • Proximal Tubule • Transforming Growth Factor Beta1 • Diabetic Nephropathy

Abstract
Background/Aims: In the current study we examined if the adipocytokine, visfatin, alters connexin-mediated intercellular communication in proximal tubule-derived epithelial cells.

Methods: The effects of visfatin (10-200ng/mL) on cell viability and cytotoxicity in HK2-cells were assessed by MTT, crystal violet and lactate dehydrogenase assays. Western blot analysis was used to confirm expression of Cx26, Cx40 and Cx43. The effect of visfatin (10-200ng/mL) on TGF-β1 secretion was confirmed by ELISA, and the effects of both TGF-β1 (2-10ng/mL) and visfatin (10-200ng/mL) on connexin expression were assessed by western blot. Functional intercellular communication was determined using transfer of Lucifer Yellow and paired-whole cell patch clamp electrophysiology.

Results: In low glucose (5mM), visfatin (10-200ng/mL) did not affect membrane integrity, cytotoxicity or cell viability at 48hrs, but did evoke a concentration-dependent reduction in Cx26 and Cx43 expression. The expression of Cx40 was unaffected. At 48hrs, visfatin (10-200ng/mL) increased the secretion of TGF-β1 and the visfatin-evoked changes in connexin expression were mimicked by exogenous application of the pro-fibrotic cytokine (2-10ng/mL). Visfatin reduced dye transfer between coupled cells and decreased functional conductance, with levels falling by 63% as compared to control. Although input resistance was increased following visfatin treatment by 166%, the change was not significant as compared to control. The effects of visfatin on Cx-expression and cell-coupling were blocked in the presence of a TGF-β1 specific neutralizing antibody.

Conclusions: The adipocytokine visfatin selectively evoked a non-toxic reduction in connexin expression in HK2-cells. The loss in gap-junction associated proteins was mirrored by a loss in functional conductance between coupled cells. Visfatin increased TGF-β secretion and the pattern of change for connexins expression was mimicked by exogenous application of TGF-β1. The effect of visfatin on Cx-expression and dye transfer were negated in the presence of a TGF-β1 neutralising antibody. These data suggest that visfatin reduces connexin-mediated intercellular communication in proximal tubule-derived epithelial cells via a TGF-β dependent pathway.
Introduction

Visfatin is an adipocytokine produced and secreted mainly by visceral adipose tissue. Alternatively known as the pre-B cell colony-enhancing factor, studies suggest that visfatin acts as an insulin-mimetic, capable of binding to the insulin receptor and inducing insulin-like effects [1, 2]. Associated with obesity and regulated by cytokines known to promote insulin resistance, these observations indicate that visfatin may have a role in the pathogenesis of type II diabetes [3]. Current evidence for its role in the disease is mixed. Some studies suggest that visfatin is reduced in patients with type I diabetes, in individuals with type II diabetes who exercise regularly and in the 3rd trimester of pregnancy where gestational diabetes is exhibited [4-7]. However, levels of visfatin have been reported to be elevated in the plasma of patients with type II diabetes [8], and a role for visfatin in the development of type II microvascular complications, for example diabetic nephropathy, cannot be ignored [8, 9].

Diabetic nephropathy is the single commonest cause of entry into the renal replacement therapy programme and with the incidence of the disease doubling in the past decade, diabetic nephropathy now accounts for approximately 50% of patients presenting with end-stage renal failure [10, 11]. Whilst our knowledge of a role for visfatin in mediating the underlying pathology of complications of type II diabetes is sparse, several reports confirm elevated visfatin in patient and animal models exhibiting diabetic nephropathy [3, 12], and increased levels of visfatin positively correlate to the degree of albuminuria in patients with type II diabetes, suggesting that endothelial dysfunction in early diabetic nephropathy may be linked to increased levels of this adipocytokine [8]. A number of studies have confirmed that the kidney is capable of synthesizing visfatin, a process exacerbated by high glucose [12]. Moreover, in mesangial cells and cells of the proximal tubule, visfatin promotes glucose influx and may be instrumental in the synthesis of pro-fibrotic molecules, including the beta-1 isoform of Transforming Growth Factor (TGF-β1), plasminogen activator inhibitor (PAI)-1 and type I collagen [3]. Through these local paracrine effectors, these data suggest that visfatin may, in part, contribute to the underlying pathology of diabetic nephropathy.

Characterised by an accumulation of extracellular matrix (ECM) in both the glomerular mesangium and tubular interstitium, the diabetic kidney exhibits multiple structural and functional changes, which culminate in excessive fibrosis and renal scarring [13, 14]. These fibrotic changes represent the crucial pathology underlying progressive chronic kidney disease (CKD) in diabetes. Changes in expression of several epithelial cell recognition and organizational proteins are central to the progression of fibrosis, and a pivotal step is the loss of cell-adhesion associated with reduced expression of epithelial protein E-cadherin (ECAD) [15, 16]. We have previously demonstrated that a loss in cell-cell adhesion is associated with a subsequent reduction in connexin expression [17]. Connexins (Cx) are membrane bound protein, which oligomerise into hexameric hemichannels (connexons) connecting the cytoplasm of adjoining cells and forming gap junctions. Gap-junctions (GJ) allow transfer of solutes, metabolic precursors and electrical currents [18], and are essential for synchronising activity to ensure appropriate function. To date our knowledge of a role for gap junctions in the kidney and in diabetic nephropathy is limited. Glucose decreases GJ-conductance and disrupts cellular homeostasis in a variety of cell systems [19, 20] and glucose-dependent down-regulation of Cx43 expression and GJ-communication has been reported in bovine retinal pericytes [21], endothelial [22], and epithelial cells [23]. Whilst the presence of GJs in the kidney has long been known, specific details regarding their function in the proximal tubule is sparse. Studies on renal vasculature have confirmed a role for various Cxs on renin secretion and the regulation of blood pressure [24], but minimal data exists on their role in tubular function where expression is also high. Using cells derived from the human proximal tubule, this study identifies a link between the adipocytokine visfatin, TGF-β1 and reduced cell-to-cell coupling. These changes will have profound effects on overall tissue integrity and function and may represent key events orchestrating loss of function in diabetic nephropathy in patients with type II diabetes.
Materials and Methods

Materials
Supplies for tissue culture were purchased from Invitrogen (Paisley, UK). Immobilon P membrane was from Millipore, Watford, UK and ECL from Amersham Biosciences, Buckinghamshire, UK. A Qproteome kit was obtained from Qiagen (Sussex, UK). Connexin antibodies were obtained from Santa Cruz (CA, USA). Anti-TGF-β1 neutralising antibody was obtained from R&D systems. Visfatin, TGF-β1, Crystal violet, MTT and Lucifer Yellow and all other general chemicals were all obtained from Sigma (Poole, UK). Anti-TGF-β1 ELISA was obtained from R&D systems. LDH assay was obtained from Abcam.

Model cell line
HK2 cells were obtained from the ATCC Bio-resource Centre (LGC Standards, Middlesex, UK). Cells (passages 18-30) were maintained in DMEM/Hams F12 (DMEM/F12) medium (17.5mM glucose), supplemented with 10% fetal calf serum (FCS), glutamine (2mM), and EGF (5ng/ml) and cultured at 37°C in a humidified atmosphere of 5% CO₂. Prior to treatment, cells were transferred to DMEM/F12 low glucose (5mM) for 48hr as described previously [25]. Cells were serum starved overnight before incubation with either visfatin (10-200ng/mL), TGF-β1 (2-10ng/mL) ± a TGF-β1 specific neutralising antibody (10μg/mL) for 48hrs.

MTT assay
The 3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay is widely used for cytotoxicity assessments of pharmacological and chemical agents. Viable cells transport MTT into their mitochondria, the compound is then reduced to formazin (purple in color), and the latter is quantified colorometrically. The amount of color formed corresponds to the number of viable cells. HK2 cells were cultured in 96-well plates (5×10³ cells/well) in 5mM glucose containing media for 48hrs prior to an overnight period of serum starvation. Cells were stimulated for 48hrs with visfatin (10-200ng/mL) and proliferation analyzed using the MTT colorimetric assay (Roche) according to manufacturers instructions. The values were presented as a percentage of the MTT uptake that was observed in visfatin treated cells as compared to control cells.

Lactate Dehydrogenase Assay
Cell death or cytotoxicity is classically evaluated by the quantification of plasma membrane damage. Lactate dehydrogenase (LDH) is a stable enzyme, present in all cell types, and rapidly released into the cell culture medium upon damage of the plasma membrane. Therefore, LDH is a common marker used to determine cytotoxicity. HK2 cells were cultured in 96-well plates (5×10³ cells/well) in 5mM glucose containing media for 48hrs prior to an overnight period of serum starvation. Cells were stimulated for 48hrs with visfatin (10-200ng/mL) and Lactate dehydrogenase levels assayed using the LDH-cytotoxicity assay kit II (Abcam) according to manufacturers instructions. The values were presented as a percentage of the LDH release that was observed in visfatin treated cells as compared to control cells.

Crystal Violet Assay
This is a simple assay useful for obtaining quantitative information about the relative density of cells adhering to multi-well cluster dishes. Crystal Violet stains DNA and upon solubilization, the amount of dye taken up by the monolayer can be quantitated in a plate reader. HK2 cells were cultured in 96-well plates (5×10³ cells/well) in 5mM glucose containing media for 48hrs prior to an overnight period of serum starvation. Cells were stimulated for 48hrs with visfatin (10-200ng/mL) and cell density determined with crystal violet staining. Briefly, media was removed and cells were fixed for 10mins with PFA. Following a brief wash with PBS, cells were incubated for 10mins at room temperature in a 1% Crystal Violet solution. After this time interval, all traces of dye were removed with distilled water and the stain solubilized with 1% SDS. The values were presented as a percentage of cells staining in visfatin treated cells as compared to control cells.

Quantification of TGF-β1
HK2 cells were cultured in 5mM glucose containing media for 48hrs prior to an overnight period of serum starvation. Cells were stimulated for 48hrs with visfatin (10-200ng/mL) under serum-free
conditions and total TGF-β1 was measured by specific enzyme-linked immunosorbent assay (ELISA) of cell culture supernatant collected from growth-arrested HK2 cells. Active TGF-β1 is measured directly and latent TGF-β1 can be measured indirectly following acid activation of samples. This assay has < 1 % cross-reactivity for TGF-β2 and TGF-β3. TGF-β1 concentration was normalized to mg/ml of protein. Data were obtained as picograms of TGF-β1 per milliliter per mg of protein.

**Immunoblotting**

Cytosolic proteins were prepared and separated by gel electrophoresis and electro-blotting onto Immobilon P membranes as described previously [26]. For determination of protein localization, proteins were harvested using the Qproteome cell compartment kit. Membranes were probed with specific polyclonal antibodies against anti-Cx26 (1:200), Cx43 (1:400) and Cx40 (1:500) (all Santa Cruz).

**Dye transfer**

Lucifer yellow was dissolved in 250 µl of fresh LiCl (150 mmol/l)/HEPES (10 mmol/l; pH 7.2). Individual cells within a cell cluster were injected using an Injectman/Femtojet 5247 delivery system (Eppendorf, Hamburg, Germany). The duration of injection was set at 1 sec with an injection pressure of 2 psi and a compensation pressure of 0.7 psi. Dye transfer between coupled cells was recorded over 4 min using Metamorph software (Molecular Devices) and a Cool Snap HQ CCD camera (Roper Scientific).

**Electrophysiology**

Cell covered 16 mm glass cover-slips were transferred to the recording bath and perfused (2 ml/min) with physiological saline solution composed of (mM): 127 NaCl, 1.9 KCl, 1.2 KH₂PO₄, 26 NaHCO₃, 1 MgCl₂, 2 CaCl₂ and 10 D-Glucose bubbled with 95% O₂/5% CO₂ and maintained at 32°C. Cells were visualised (x600 magnification) using IR-DIC optics (Olympus BX51W1 microscope and Hitachi CCD camera). Paired whole cell patch-clamp recordings were made from neighbouring cells using thick-walled borosilicate (Harvard) glass pipettes (3-8 MΩ) containing (mM): 135 potassium gluconate, 7 NaCl, 10 HEPES, 0.5 EGTA, 10 phosphocreatine, 2 MgATP, 0.3 NaGTP (~300 mOSM, pH 7.2). Recordings were made using an Axon Multiclamp 700B amplifier (Molecular Devices) and digitized at 20 kHz (Axon Digidata 1440a). Data acquisition and analysis was performed using pClamp (v10, Axon, Molecular Devices).

Following the establishment of paired whole cell recordings, the series resistance (10-16 MΩ) was continually monitored and compensated appropriately. If the series resistance markedly changed during the recording, then the data was excluded from analysis. Coupling was detected by the observation of a membrane potential change in one cell following injection of hyperpolarising and depolarising current steps in the other cell. At the end of recordings, the electrodes were removed from the cells and identical current steps were injected to ensure that there was no cross talk between electrodes.

**Estimating Coupling Strength**

Input resistance and coupling coefficients (ratio of membrane potential changes between the two cells) were determined from membrane potential responses following the injection of a series of hyperpolarising and depolarising current steps into each cell. The junctional conductance (Gj) was then calculated using the following equation [27, 28]

\[ G_{j12} = \frac{R_{1}k_{12}}{(R_{1}R_{2} - (R_{1}k_{12})^2)\} \]

Where R₁ and R₂ are the input resistances of the 2 cells and k₁₂ is the coupling coefficient. Estimates were calculated in both directions between pairs of cells.

**Analysis**

Autoradiographs were quantified by densitometry using TotalLab 2003 (NonLinear Dynamics, Durham, NC USA). Where data was quantified, the non-stimulated, low glucose control condition was normalized to 100% and data from all other experimental conditions compared to this. Statistical analysis of data was performed using a one-way ANOVA test with a Tukey’s multiple comparison post-test. Paired Patch data was analysed via t-test. Data are expressed as mean ± SEM, and ‘n’ denotes the number of experiments. Probability (P) * < 0.05 was taken to signify statistical significance.
Results

The effect of Visfatin on cell viability and cytotoxicity

Cells were cultured in 5mM glucose for 48hrs prior to being serum starved overnight. Cells were either non-stimulated (control) or stimulated for 48hrs with visfatin (10-200 ng/mL) under serum-free conditions and were assessed for cell viability by MTT uptake (uptake directly correlates with the number of viable cells). Incubation with visfatin at 10, 100 and 200ng/mL for 48hrs failed to significantly alter cell viability (Fig. 1A, P>0.05, n=3).

To confirm that the effects of visfatin, could not be attributable to toxicity at concentrations used in subsequent analysis, we used a crystal violet assay to measure the number of adherent cells after 48hr treatment with visfatin (10-200ng/mL), in serum-free conditions. Cells were fixed and stained with crystal violet (1% w/v). The extent of dye uptake in visfatin treated cells did not significantly differ to that from control cells at 48hrs (Fig. 1B, P>0.05, n=3). Data from all 3 complimentary strategies was used to inform on the concentration of visfatin to be used in subsequent analysis, i.e. 10-200ng/mL.

As a complimentary strategy to assess membrane integrity, we used the Lactate Dehydrogenase assay as a marker of LDH release from control versus visfatin-treated cells. Cells were stimulated for 48hrs with visfatin (10-200ng/mL) under serum-free conditions prior to measuring LDH release. Results, expressed as a % of LDH release as compared to control, indicate that there was no significant change in LDH release from cells treated with visfatin (Fig. 1C, P>0.05, n=3), suggesting that the adipocytokine (10-200ng/mL) does not damage the cell membrane.

Fig. 1. Impact of visfatin on HK2 cell viability, as assessed by MTT uptake, Crystal Violet staining and LDH release. HK2 cells were cultured in 5mM glucose containing media for 48hrs prior to an overnight serum starvation. Cells were stimulated for 48hrs with visfatin (10-200 ng/mL) under serum-free conditions and cell viability assessed by MTT uptake (panel A). Cell density was determined by the Crystal violet assay (panel B) at 48hrs (panel B). Finally cell membrane damage was assessed via LDH release at 48hrs (panels C). The values obtained are expressed as a % of control. Results are representative of 3 separate experiments.
**Fig. 2.** Visfatin evoked changes in connexin expression in HK2 cells. To assess the effect of visfatin on expression of gap junction proteins HK2 cells were cultured in 5mM glucose containing media for 48hrs prior to overnight serum starvation. Cells were stimulated for 48hrs with visfatin (10-200 ng/mL) under serum-free conditions and phenotypic changes of Cx26, Cx43 and Cx40 assessed. Whole cell expression was determined by western blotting and confirmed that Visfatin evoked a dose dependent decrease in expression of both Cx26 (panel A) and Cx43 (panel B). Compartmental localisation of Cx26 and Cx43 were determined for membrane (M), cytosol (C), nuclear (N) and cytoskeletal (CK) fractions +/-Visfatin (200ng/mL). Visfatin altered the cellular localization of both Cx26 and Cx43 as compared to control (panels Dii, and Div respectively). However, visfatin failed to exert a significant effect over expression and cell localisation of Cx40 (panel C and Dvi respectively). Upper panels show representative blots for each protein and re-probed for α-tubulin as a loading control. Lower panels show mean (±SEM) densitometry data, normalised against the non-stimulated low glucose control (100 %), from 3 or more separate experiments. Each lane in the representative blot corresponds to the associated bar in the graph. Key significances are shown, **P<0.01, ***P<0.001.

**Visfatin evokes a concentration-dependent down-regulation of Cx26 and Cx43 in HK2 cells**

HK2 cells were cultured in 5mM glucose prior to overnight serum starvation. Cells were then stimulated for 48hrs with visfatin (10-200ng/mL) under serum-free conditions and cell lysates analysed by immunoblotting to determine the effects of the adipocytokine on the expression of Cx26, Cx40 and Cx43. Visfatin decreased whole-cell expression of Cx26 in a concentration-dependent manner to 58±11%, 40± 3% and 21±2% as compared to control at 10,100 and 200ng/mL respectively (Fig. 2A n=3; P<0.01). Cell compartment analysis revealed that the loss in Cx26 expression was attributable to a decrease in expression from both the cell membrane and cytosol (Fig. 2Di and ii). Similarly, visfatin also decreased Cx43...
Visfatin increases secretion of TGF-β1

The pro-fibrotic cytokine TGF-β1 is a principal mediator of fibrotic changes in the kidney. TGF-β1 modulates the expression of several epithelial cell recognition and organizational proteins, whilst contributing to the reciprocal loss of tubular epithelial cells and accumulation of interstitial fibroblasts, changes associated with declining excretory function [14, 29, 30]. To determine if visfatin stimulated TGF-β1 secretion, cells were cultured in 5mM glucose prior to being starved of serum overnight. Cells were stimulated for 48hrs with visfatin (10-200ng/mL) under serum-free conditions and a TGF-β1 ELISA was used to measure total secretion from growth-arrested HK2 cells. ELISA of supernatant showed that exposure of HK2 cells to visfatin for 48hrs evoked a rise in total TGF-β1 secretion to 51±4 pg/mL, 58±6 pg/mL and 59±10pg/mL respectively at 10, 100 and 200ng/mL visfatin (Fig. 3), as compared to a 5mM glucose control; 32±7pg/mL (n=3; P<0.01). Differences in TGF-β1 were only detected following acidification of the samples, suggesting that TGF-β1 was produced in its latent form.

TGF-β1 down-regulates Cx26 and Cx43

Having confirmed that visfatin down-regulates Cx26 and Cx43 expression and increases secretion of TGF-β1 we wanted to determine the effects of TGF-β1 on Cx26, Cx43 and Cx40 expression to see if the effects of the pro-fibrotic cytokine matched those of visfatin. HK2 cells were cultured in 5mM glucose prior to overnight serum starvation. Cells were then stimulated for 48hrs with TGF-β1 (2-10ng/mL) under serum-free conditions. TGF-β1 decreased whole-cell expression of Cx26 in a concentration-dependent manner to 37±24%, 22±5% and 15±8% as compared to control at 2, 4 and 10ng/ml respectively (Fig. 4A n=3; P<0.001). Cell compartment analysis revealed that this loss in expression was attributable to removal of Cx26 from the cytosol (Fig. 4Di and ii). Similarly, TGF-β1 also reduced Cx43 expression with expression falling to 52±15%, 28± 9% and 9±18% as compared to control at 2, 4 and 10ng/ml respectively (Fig. 4B n=3; P<0.001). The loss of whole cell expression was primarily attributable to removal of Cx43 from the cell membrane (Fig. 4Diii and iv). In support of our previous data for visfatin, TGF-β1 failed to alter Cx40 expression (Fig 4C), which remained predominantly located at the membrane (Fig. 4Dv and vi).

Visfatin evoked changes in Cx26 and Cx43 expression are mediated by TGF-β1

To determine the role of TGF-β1 in mediating the visfatin-evoked loss in Cx26 and Cx43 expression we examined the effect of visfatin (200ng/mL) in conjunction with a TGF-β1 specific immunoneutralising antibody (Fig. 5 A and B respectively). HK2 cells were cultured in 5mM glucose prior to overnight serum starvation. Cells were then stimulated for 48hrs with visfatin (200ng/mL) +/- a TGF-β1 specific immunoneutralising antibody (10µg/mL)
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under serum-free conditions. Cell lysates were analysed by immunoblotting to determine the effects of the adipocytokine on the expression of Cx26 and Cx43. As previously observed, Visfatin (200ng/mL) decreased whole-cell expression of Cx26 to 32±15%, as compared to control (n=3, P<0.001) (Fig. 5A). However, co-incubation with TGF-β1-specific immuno-neutralising antibody negated the effect of visfatin, with Cx26 expression remaining at 90±6%; as compared to low glucose (5mM) control. Visfatin (200ng/mL) decreased Cx43 expression to 27±18%, as compared to control (n=3, P<0.001) (Fig. 5B). Co-incubation with TGF-β1-specific immuno-neutralising antibody negated the effect, with Cx43 expression remaining at 81±4%; as compared to low glucose (5mM) control. Incubation with TGF-β1-specific immuno-neutralising antibody had no effect on Cx26 or Cx43 expression.

Fig. 4. TGF-β1 evoked changes in connexin expression in HK2-cells. To assess the effect of TGF-β1 on expression of gap junction proteins, HK2 cells were cultured in 5mM glucose containing media for 48hrs prior to overnight serum starvation. Cells were stimulated for 48hrs with TGF-β1 (2-10ng/mL) under serum-free conditions and phenotypic changes of Cx26, Cx43 and Cx40 assessed. Whole cell expression was determined by western blotting and confirmed that visfatin evoked a dose dependent decrease in expression of both Cx26 (panel A) and Cx43 (panel B). Compartmental localisation of Cx26 and Cx43 were determined for membrane (M), cytosol (C), nuclear (N) and cytoskeletal (CK) fractions +/- TGF-β1 (2-10ng/mL). TGF-β1 altered the cellular localization of both Cx26 and Cx43 as compared to control (panels Dii, and Div respectively). However, TGF-β1 failed to exert a significant effect over expression and cell localisation of Cx40 (panel C and Dvi respectively). Upper panels show representative blots for each protein and reprobed for β-tubulin as a loading control. Lower panels show mean (±SEM) densitometry data, normalised against the non-stimulated low glucose control (100 %), from 3 or more separate experiments. Each lane in the representative blot corresponds to the associated bar in the graph. Key significances are shown, ***P< 0.001.
Visfatin-induced loss in Cx-expression is associated with impaired cell-coupling

To determine if visfatin-evoked loss in Cx-expression had functional implications for cell-to-cell communication we examined changes in dye-transfer and junctional conductance between HK2 cells using paired-patch recording. Cells were seeded onto coverslips and cultured in 5 mM glucose prior to being serum starved overnight. Cells were stimulated for 48 hrs with visfatin (10-200 ng/mL) under serum-free conditions and gap junction intercellular communication (GJIC) assessed (Fig. 6).

Lucifer yellow is a membrane impermeant dye, which moves between adjacent cells via gap junctions. The dye was injected into individual cells within a cell cluster (5-8 cells/cluster). Cells treated for 48 hrs with either TGF-β1 (2-10 ng/mL) or visfatin (10-200 ng/mL) failed to exhibit dye transfer (Fig. 6B and C respectively), whilst dye injected into non-stimulated control cells propagated away from the site of injection into neighbouring cells, suggesting gap junction-mediated intercellular communication (Fig. 6A). Incubation of cells with visfatin (200 ng/mL), in the presence of a TGF-β1 neutralising antibody (10 µg/mL), permitted dye transfer into neighbouring cells and away from the site of injection, suggesting that the loss of gap junction-mediated intercellular communication in response to visfatin is TGF-β1 dependent (Fig. 6D).

Paired whole cell patch clamp recordings were used to determine both the junctional conductance and the input resistance of HK2 cells ± visfatin (200 ng/mL). Cells were seeded onto coverslips and cultured in 5 mM glucose prior to starvation overnight without serum. Cells were stimulated for 48 hrs with visfatin (200 ng/mL). Representative recordings are illustrated for control (Fig. 6 Ei and F) and visfatin treated cells (Fig. 6 Eii and G). Input resistance was calculated from the slope of the current-voltage relationship around the resting potential of the cells. Under control conditions (Fig. 6 Ei) the resistance of cell 1 was 154 MΩ, and the resistance for cell 2 was 256 MΩ, the coupling coefficient was 0.48, giving a junctional conductance of 2.19 nS. In cells treated with visfatin (200 ng/mL) (Fig. 6 Eii) the resistance of cell 1 was 395 MΩ and the resistance of cell 2 was 163 MΩ, the coupling coefficient was 0.10, giving a junctional conductance of 0.61 nS.
Fig. 6. Visfatin-induced loss in Cx-expression is associated with impaired cell-coupling in HK2 cells. Panels A-D show Lucifer yellow dye transfer between HK2-cells. Monochrome plates illustrate phase images of HK2-cell clusters. Fluorescence (fluorescein) image of same cell clusters after single-cell injection with Lucifer yellow is shown at time 0. The same field of view is recorded 2 and 4 min after injection of dye in control cells (A), in the presence of TGF-β1 (10ng/mL; B), in cells treated with visfatin (200ng/mL; C) and lastly in cells treated with visfatin (200ng/mL) + TGF-β1 neutralising antibody (10µg/mL; D). The transfer of dye supports direct cell-to-cell communication between coupled cells and is only seen in control cells or following neutralization of TGF-β1 with visfatin. Panel E, show the positioning of electrodes (arrows) for paired whole cell patch clamp recordings from cells in a control culture (i) and in a culture treated with visfatin (ii). The electrophysiological data for these paired recordings is illustrated in (F, control) and (G, visfatin treated). F, Hyperpolarising and depolarising current steps (40 pA) injected into cell-1 produced voltage responses in cell-2 demonstrating that the cells are electrically coupled. The junctional conductance was calculated as 2.19 nS. F, Hyperpolarising and depolarising current steps (20 pA) injected into cell-1 produced voltage responses in cell-2 demonstrating that they are electrically coupled. The junctional conductance was calculated as 0.61 nS. Panel H, Graph plotting the mean junctional conductance for pairs of cells in control conditions versus cells treated with visfatin. The junctional conductance is significantly reduced in visfatin (2-tailed unpaired t-test, P<0.01). Panel I, shows mean input resistance for cells in control and treated with visfatin. Although there is a trend towards an increase in input resistance this is not significant (as there is a large variation in input resistance across cells).
The junctional conductance in HK2 cells in control conditions was 2.25±0.2 nS (n=6). Treatment with visfatin significantly reduced the junctional conductance (63% reduction to 0.83±0.19 nS; Fig. 6H; n=6; P<0.01). The fall in junctional conductance would be expected to increase the input resistance of the cells as less current will leak out of the membrane. (Fig. 6I).

Discussion

Produced and secreted mainly by visceral adipose tissue, adipocytokines have been studied for their association with insulin resistance and metabolic syndrome [1, 2]. Elevated in patients with type II diabetes and linked to endothelial dysfunction, a series of studies have addressed the potential role of visfatin in the pre-disposition of micro-vascular complications [3, 8, 9]. The current study suggests that visfatin can modulate gap junction mediated cell-to-cell communication in the human proximal tubule, a process that may have implications for renal function ahead of overt changes that are central to the progression of renal fibrosis. We have demonstrated that cells of the human proximal tubule express the gap junction proteins Cx26, Cx40 and Cx43 and that the expression of Cx26 and Cx43 are sensitive to increased levels of the adipocytokine. A loss in Cx-expression may be crucial in the development of complications of diabetes. In retinal capillaries of diabetic mice, Cx43 expression is reduced and apoptosis increased, resulting in a loss of cell-cell communication and a decline in the number of pericytes and acellular capillaries [19]. Similar findings in vascular endothelial cells, confirm that high glucose decreases Cx-expression/function and that this is an early trigger for apoptosis [20]. The data highlight the importance of GJ-mediated cell-to-cell coupling and suggests that a loss in cell-cell communication may contribute to some micro-vascular complications associated with diabetes.

More than a dozen fibrogenic factors affect renal function, however, it is widely recognised that the pro-fibrotic cytokine TGF-β1 and its downstream signalling partners represents the predominant pathway orchestrating renal fibrosis [15, 16]. In the current study we confirm that visfatin evokes an increase in TGF-β1 secretion in and that exogenous application of TGF-β1 to HK2 cells induces synonymous effects to visfatin in reducing both Cx26 and Cx43 expression. Furthermore, using a TGF-β1 specific neutralising antibody, we confirm that the effects of visfatin are, at least in part, mediated by TGF-β1. Abnormalities in TGF-β1 have been linked to a variety of disorders, including autoimmune diseases, malignancies, and chronic renal disease [11]. TGFβ1 is important in many tubulointerstitial diseases where disassembly of the adherens junction represents the initial overt change in epithelial organisation, well before any suggested cellular migration associated with epithelial-to-mesenchymal transition (EMT), a process in which cells convert from an epithelial to mesenchymal phenotype, and an event central to the progression of tubulointerstitial fibrosis. [15, 16]. In the current study, we have used dye transfer and paired patch clamp recordings to provide functional evidence that visfatin dramatically reduces connexin mediated intercellular communication in the proximal tubule, an effect that may be associated with the early renal damage in type II diabetes.

In podocytes and cells of the proximal tubule, that are able to synthesize visfatin, GLUT1-dependent glucose uptake is increased [3]. In 2010, Kang et al. confirmed that the concentration of visfatin in both the glomeruli and tubulointerstitium, was increased in a rodent model of type II diabetes mellitus [3]. Since then, elevated levels of circulating visfatin have been found in patients with the disease [31]. Plasma visfatin levels are significantly elevated in early stages of diabetic nephropathy and positively correlate with body weight, fasting plasma glucose and microalbuminuria [3, 9]. The current study demonstrates that when the concentration of this important adipocytokine increases, gap-junction mediated intercellular communication between cells of the proximal tubule decreases. Our data could help explain an early loss in renal function in patients with T2DM.
Conflict of Interest

The authors declare that there is no duality of interest associated with this manuscript.

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