We have previously reported that the fractalkine (FKN)/CX3CR1 system represents a novel regulatory mechanism for insulin secretion and β cell function. Here, we demonstrate that chronic administration of a long-acting form of FKN, FKN-Fc, can exert durable effects to improve glucose tolerance with increased glucose-stimulated insulin secretion and decreased β cell apoptosis in obese rodent models. Unexpectedly, chronic FKN-Fc administration also led to decreased α cell glucagon secretion. In islet cells, FKN inhibited ATP-sensitive potassium channel conductance by an ERK-dependent mechanism, which triggered β cell action potential (AP) firing and decreased α cell AP amplitude. This results in increased glucose-stimulated insulin secretion and decreased glucagon secretion. Beyond its islet effects, FKN-Fc also exerted peripheral effects to enhance hepatic insulin sensitivity due to inhibition of glucagon action. In hepatocytes, FKN treatment reduced glucagon-stimulated cAMP production and CREB phosphorylation in a pertussis toxin-sensitive manner. Together, these results raise the possibility of use of FKN-based therapy to improve type 2 diabetes by increasing both insulin secretion and insulin sensitivity.
Chronic fractalkine administration improves glucose tolerance and pancreatic endocrine function

Matthew Riopel,1 Jong Bae Seo,1,2 Gautam K. Bandyopadhyay,1 Pingping Li,1,3 Joshua Wollam,1 Heekyung Chung,1 Seung-Ryoung Jung,2 Anne Murphy,4 Maria Wilson,5 Ron de Jong,5 Sanjay Patel,5 Deepika Balakrishna,5 James Bilakivos,5 Andrea Fanjul,1 Artur Plonowski,5 Duk-Su Koh,2 Christopher J. Larson,4,6 Jerrold M. Olefsky,1,7 and Yun Sok Lee1,7

1Department of Medicine, Division of Endocrinology and Metabolism, UCSD, La Jolla, California, USA. 2Department of Physiology and Biophysics, University of Washington, Seattle, Washington, USA. 3State Key Laboratory of Bioactive Substance and Function of Natural Medicines, Institute of Materia Medica, Peking Union Medical College and Chinese Academy of Medical Sciences, Beijing, China. 4Department of Pharmacology, UCSD, La Jolla, California, USA. 5Cardiovascular and Metabolic Diseases Drug Discovery Unit, Takeda Pharmaceuticals, San Diego, California, USA. 6Sanford Burnham Prebys Medical Discovery Institute, La Jolla, California, USA. 7Graduate School of Medical Science and Engineering, Korea Advanced Institute of Science and Technology, Daejeon, South Korea.

We have previously reported that the fractalkine (FKN)/CX3CR1 system represents a novel regulatory mechanism for insulin secretion and β cell function. Here, we demonstrate that chronic administration of a long-acting form of FKN, FKN-Fc, can exert durable effects to improve glucose tolerance with increased glucose-stimulated insulin secretion and decreased β cell apoptosis in obese rodent models. Unexpectedly, chronic FKN-Fc administration also led to decreased α cell glucagon secretion. In islet cells, FKN inhibited ATP-sensitive potassium channel conductance by an ERK-dependent mechanism, which triggered β cell action potential (AP) firing and decreased α cell AP amplitude. This resulted in increased glucose-stimulated insulin secretion and decreased glucagon secretion. Beyond its islet effects, FKN-Fc also exerted peripheral effects to enhance hepatic insulin sensitivity due to inhibition of glucagon action. In hepatocytes, FKN treatment reduced glucagon-stimulated cAMP production and CREB phosphorylation in a pertussis toxin–sensitive manner. Together, these results raise the possibility of use of FKN-based therapy to improve type 2 diabetes by increasing both insulin secretion and insulin sensitivity.

Introduction

In type 2 diabetes mellitus (T2DM), progression to the full hyperglycemic state is associated with 2 major pathophysiological defects: insulin resistance and β cell dysfunction (1–3). While the relative contribution and temporal appearance of these events vary across different patients and populations, these 2 abnormalities usually exist concomitantly in the majority of patients (4). Therefore, an ideal antidiabetic therapeutic would involve strategies to enhance both insulin sensitivity and β cell function.

One component of insulin’s action to suppress postprandial hyperglycemia is to counteract the effects of glucagon (5, 6). In T2DM, insulin resistance potentiates α cell glucagon secretion and hepatic glucagon action, which stimulates hepatic glucose production (7, 8). Recently, it has been shown that inhibition of glucagon action using glucagon receptor–antagonizing antibodies normalizes blood glucose levels in streptozotocin-treated insulin-deficient mice (9). Glucose-lowering effects of glucagon antagonism have also been demonstrated in nonhuman primates and humans (10–13). Moreover, hepatocyte-specific deletion of the glucagon receptor improves glucose tolerance with improved insulin sensitivity (14), suggesting that inhibition of glucagon secretion and/or action is beneficial for improving glucose tolerance and insulin sensitivity.

An impaired insulin response to glucose has long been documented in T2DM subjects (15–19), and evidence indicates that β cell dysfunction involves a decline in both β cell responsiveness to insulin secretagogues and overall β cell mass (4, 20). Ex vivo glucose-stimulated insulin secretion (GSIS) is diminished in primary islets from T2DM patients compared with normal subjects (20), indicating that there is an islet-autonomous defect in GSIS in T2DM.

Secretion of insulin and glucagon is tightly regulated by blood glucose concentration, and, at the molecular level, the response to glucose is largely mediated through a common mechanism involving ATP-sensitive potassium (KATP) channel regulation (21). The postprandial rise in blood glucose increases the ATP/ADP ratio, leading to inhibition of KATP channel activity and plasma membrane depolarization in α and β cells. In β cells, this leads to action potential (AP) firing and insulin secretion (21). On the other hand, in α cells, basal membrane potential is already depolarized with spontaneous AP firing at low glucose levels, and further depolarization of membrane potential by glucose-dependent inhibition of KATP channels results in decreased AP amplitude and decreased glucagon secretion. Recently, it has been suggested that a slight increase in basal KATP channel conductance or incomplete inhibition of KATP channel activity by glucose may be responsible for the decreased glucose response in α and β cells in T2DM (22, 23).

Circulating fractalkine (FKN; also called CX3CL1) is a peptide released into the bloodstream by proteolytic cleavage of the
CX3CL1 extracellular domain and is readily detected in mouse and human blood (24, 25). Cellular KFN effects are largely mediated through its cognate receptor CX3CR1 (26). Recently, we reported that CX3CR1-deficient mice exhibit glucose intolerance mainly due to β cell dysfunction, while FKN treatment improves glucose tolerance and increases insulin secretion in vivo (27). FKN also potentiates GSIS in vitro in both mouse and human islets. Additionally, it has been shown that FKN treatment of human islets decreases glucagon release and protects against TNFα-induced decreased GSIS (28). In this study, we used a long-acting FKN-Fc fusion protein to conduct chronic treatment studies in obese mice. The results show that FKN-Fc administration leads to improved glucose tolerance, increased insulin secretion, and enhanced hepatic insulin sensitivity. These beneficial effects are mediated by distinct mechanisms in β cells, α cells, and hepatocytes.

Results
An FKN-Fc fusion protein exerts durable effects to enhance insulin secretion and improve glucose tolerance. We previously reported that CX3CR1 (FKN receptor) deficiency causes β cell dysfunction and glucose intolerance. Furthermore, acute administration of soluble FKN increases insulin secretion and improves glucose tolerance in lean chow-fed WT mice (27). These results suggested that FKN may have translational importance for antidiabetes therapy, raising the question of whether chronic FKN administration can exert durable beneficial effects to improve insulin secretion and glucose tolerance in obese/diabetic mice. Native soluble FKN is not adequate to test this idea, since it is rapidly cleared with a half-life of approximately 30 minutes (data not shown). To overcome this problem, we first generated a long-acting form of soluble FKN by fusing the chemokine domain (76 amino acids) of native FKN to a mouse IgG Fc fragment. This chimeric FKN-Fc fusion protein displayed a significantly increased in vivo half-life (~6 hours) in both normal chow diet (NCD) and high-fat diet (HFD) mice (data not shown). As seen in Figure 1, A–C, acute administration of FKN-Fc improved glucose tolerance and increased plasma insulin levels in both NCD and HFD mice. In NCD mice, these beneficial effects of FKN-Fc persisted up to 2 days at the dose 10 mg/kg (Supplemental Figure 1; supplemental material available online with this article; https://doi.org/10.1172/JCI94330DSI), and, upon repeated treatment at day 5, there was continued improvement in glucose tolerance (Figure 1B), indicating no decreased FKN sensitivity with repeated dosing. In HFD mice, the effect of FKN persisted for 2 days at a dose of 30 mg/kg (Figure 1C), a dose at which the effect persisted for 5 days in NCD mice (data not shown). Furthermore, plasma glucagon levels were decreased by acute FKN-Fc administration in both NCD and HFD mice (Figure 1D). Together, these data suggest that acute administration of FKN-Fc increases plasma insulin levels, decreases plasma glucagon levels, and improves glucose tolerance in both NCD and HFD mice.

We next tested the effects of chronic administration of FKN-Fc in HFD obese/diabetic mice. WT B6 mice were fed HFD for 10 weeks and then treated with FKN-Fc (30 mg/kg) every other day for an additional 8 weeks. During the 8 weeks of treatment, the mice were maintained on HFD. As seen in Figure 1, E and F, chronic FKN-Fc administration did not change body weight or food intake. Chronic FKN-Fc administration significantly improved glucose tolerance in HFD WT mice (Figure 1G), but not in HFD CX3CR1-KO mice (Figure 1H), showing that these beneficial effects of FKN-Fc are CX3CR1-dependent. Interestingly, chronic FKN-Fc administration decreased fasting plasma insulin level (Figure 1I), similarly to what has been reported in chronic GLP-1 analog-treated animals (29).

Chronic FKN-Fc treatment enhances insulin secretion and decreases β cell apoptosis in obese mice. β Cell dysfunction in T2DM is characterized by reduced GSIS activity and decreased β cell mass due to apoptosis (4). Interestingly, in primary mouse islets, FKN-Fc stimulated GSIS and inhibited the effect of palmitate treatment to cause apoptosis (Figure 2, A and B). On the other hand, insulin secretion was not affected by FKN-Fc treatment in low-glucose conditions (Supplemental Figure 2A). To assess this concept in isolated islets, we first measured GSIS in the islets isolated from vehicle- and FKN-Fc–treated mice. As seen in Figure 2C and Supplemental Figure 2, B and C, chronic FKN-Fc administration increased GSIS activity in the islets of HFD mice. In addition, the HFD-induced decrease in the expression of genes involved in β cell differentiation and function, such as Pdx1, Neurod, Glut2, Mafa, and Cx36, was prevented by FKN-Fc administration (Figure 2D and Supplemental Figure 2D). Next, we tested changes in β cell apoptosis and found that caspase-3/7 activity was significantly diminished in islets by FKN-Fc administration to HFD mice (Figure 2E).

To test this idea in the in vivo setting, we quantified apoptotic cell number in FKN-Fc–treated mice. In HFD mice, the number of apoptotic β cells (active caspase-3– and insulin–double-positive cells) was marginal (only 1–2 cells per islet) (data not shown), making it unfeasible to detect an FKN-Fc–induced decrease in apoptosis using immunohistochemistry. To overcome this problem, we examined other models of obesity and found that 15-week-old ob/ob mice exhibited a higher number (~4%) of apoptotic islet β cells (Figure 2F). Seven weeks of chronic FKN-Fc administration significantly improved glucose tolerance (Supplemental Figure 2E) and markedly decreased the number of apoptotic β cells in the ob/ob mouse islets (Figure 2F). Consistent with this, caspase-3/7 activity (Supplemental Figure 2F) and the expression of genes involved in β cell apoptosis, such as Tnixip, p21, and Icer, were decreased by FKN-Fc administration (Figure 2G), while the expression of genes involved in β cell differentiation was increased (Supplemental Figure 2G). In addition, overall β cell mass was significantly increased (Figure 2H), without changes in islet number per pancreatic area in HFD WT mice (Figure 2I).

Chronic FKN-Fc administration inhibits hepatic glucose production and improves hepatic insulin sensitivity. Since fasting plasma insulin was markedly decreased without change in fasting glucose level by chronic FKN-Fc administration in HFD mice (Figure 1I), we questioned whether the improved glucose tolerance is associated with enhanced insulin sensitivity. To assess this, we performed insulin tolerance tests. As seen in Figure 3A, chronic FKN-Fc administration significantly improved insulin tolerance in HFD WT mice at 90 and 120 minutes. To quantitate systemic insulin sensitivity and assess tissue-specific effects, we performed hyperinsulinemic-euglycemic clamp studies in these mice. As seen in Figure 3, B–D, the glucose infusion rate was increased 56% by chronic FKN-Fc administration, and this systemic effect was due to increased hepatic insulin sensitivity (Figure 3C) with no change...
in basal or insulin-stimulated glucose disposal rates (Figure 3D). Moreover, liver mass, triglyceride, nonesterified fatty acid content, and histologic indications of steatosis were all significantly reduced by chronic FKN-Fc treatment in HFD WT mice and ob/ob mice (Supplemental Figure 3).

Mechanism of FKN effects on hepatocytes. To determine whether this in vivo effect on hepatic insulin resistance is hepatocyte-autonomous, or through an indirect mechanism, we incubated mouse primary hepatocytes isolated from normal mice with glucagon, insulin, and/or FKN and measured glucose production from these cells. As seen in Figure 3E, FKN markedly reduced glucagon-stimulated glucose production from 14C-labeled pyruvate. Concomitant with this, FKN treatment reduced glucagon-stimulated cAMP levels (Figure 3F) and cAMP responsive element-binding protein (CREB) phosphorylation (Figure 3G), comparable to the effect of insulin. Mechanistically, FKN-dependent suppression of hepatic glucose production was blocked by pertussis toxin treatment (Figure 3H), but not by inhibition of MEK (Supplemental Figure 3G), indicating that FKN can suppress hepatic glucagon action by a Gαi-dependent mechanism. Next, we tested insulin-resistant HFD mouse hepatocytes. In these cells, FKN treatment caused a trend toward decreased glucagon-stimulated glucose production (Figure 3I), although this did not reach statistical significance. Notably, FKN potentiated insulin-dependent suppression of glucagon-stimulated glucose production and cAMP levels (Figure 3, I and J). Taken together, these results suggest that FKN activates the Gαi pathway in hepatocytes and can enhance hepatic insulin action by antagonizing glucagon.

Mechanism of FKN effects on β cells. Our studies show that FKN stimulates insulin secretion in vivo in mice and in vitro in isolated...
treatment with the MEK inhibitor U0126 or PD98059 (Figure 4, A and B), suggesting that FKN inhibits KATP channel through a MEK/ERK-dependent mechanism independent of changing mitochondrial activity. Consistent with the Min6 cell data, FKN treatment significantly inhibited KATP channel activity in primary β cells from WT mice, but not CX3CR1 KO mice (Figure 4, E and F). Furthermore, FKN treatment led to membrane depolarization (from –74 mV to –35.8 mV) and AP firing (2.2 ± 0.48 Hz) in low-glucose (2 mM) conditions in Min6 cells, when measured in a perforated whole-cell configuration (Figure 4G), while cellular metabolic flux (e.g., ATP/ADP ratio) was unchanged. Importantly, when administered after a high-glucose challenge, FKN significantly increased AP firing frequency (1.04 ± 0.21 vs. 2.02 ± 0.38 Hz) (Figure 4H), suggesting that FKN can potentiate glucose-stimulated electrical activity in β cells, leading to increased intracellular Ca2+ levels and GSIS. On the other hand, FKN treatment was without direct effect on VDCC activity (Figure 4, I and J). These results indicate that FKN inhibits KATP channel activity, leading to increased intracellular Ca2+ levels and GSIS.

**Mechanism of FKN effects on α cells.** Recently, it was reported that FKN treatment can suppress glucagon release in primary human mouse and human islets through CX3CR1. β Cells are electrically excitable by insulin secretagogues, and modulation of cation channels provokes action potential (AP) firing, leading to activation of voltage-dependent Ca2+ channels (VDCCs), increased intracellular calcium ion (Ca2+) levels, and Ca2+-dependent insulin exocytosis. Since our earlier studies showed that FKN stimulates GSIS by potentiating glucose-dependent increases in intracellular β cell Ca2+ levels (27), we have now conducted a series of electrophysiology studies to examine the molecular mechanisms of this effect. First, we examined whether FKN modulates β cell cation channels critical for GSIS, including the KATP channel and/or the VDCC, using patch-clamp techniques. When Min6 cells were incubated in low-glucose (2 mM) medium for 16 hours, resting KATP current density was 25.3 ± 2.86 pA/pF in an external solution containing 40 mM KCl. Upon FKN treatment, the KATP channels were rapidly inhibited within 30 seconds in a dose-dependent manner (81.5% ± 5.07% and 99.4% ± 0.51% by 10 and 100 ng/ml FKN, respectively), and this effect was not reversed during 2 minutes of washout (Figure 4, A and B). Moreover, this inhibitory effect of FKN occurred without changing cellular ATP levels, the ATP/ADP ratio, or the oxygen consumption rate (Figure 4, C and D, and Supplemental Figure 4C). The effects of FKN-Fc were prevented by pre-
islets (28). Consistent with this, we observed that FKN-Fc administration decreases plasma glucagon levels in mice (Figure 1D). Moreover, in vitro incubation of primary mouse islets with FKN suppressed low-glucose-induced glucagon release, without interfering insulin effects (Figure 5A). Since insulin suppresses α cell glucagon release (30, 31), we hypothesized that FKN treatment could decrease glucagon release from α cells, contributing to the improved glucose tolerance (6). Surprisingly, FKN also directly suppressed glucagon release in αTC1 mouse α cells (Figure 5B), raising the possibility that FKN can act on α cells to inhibit glucagon secretion. Consistent with this idea, immunohistochemistry analysis of mouse islets revealed that there are glucagon– and CX3CR1– double-positive cells in mouse islets (Figure 5C). However, pancreatic endocrine cells expressed relatively low Cx3cr1 mRNA levels, compared with monocytes (data not shown), and using immunohistochemical methods with the CX3CR1 antibody to measure the ratio between CX3CR1+ and CX3CR1– islet cells is not the ideal quantitative approach. Therefore, to obtain more precise insight into CX3CR1 expression in primary α and β cells, we performed flow cytometry analyses in dispersed islet cells. As seen in Figure 5D, 97.7% of glucagon-positive and 97.1% of insulin-positive cells were CX3CR1+, showing that most of the primary α and β cells express CX3CR1.

High glucose levels suppress α cell glucagon release by the same cellular mechanisms whereby glucose stimulates GSIS: inhibition of KATP channel activity. Compared with β cells, α cell KATP channel conductance is much less (1%–10%) in the absence of glucose, and relatively less ATP is required to attenuate whole-cell KATP currents in α cells than in β cells (21). Therefore, unlike in β cells, the basal membrane potential in α cells is already depolarized (about −40 mV) even at relatively low levels of intracellular ATP, as seen in the CX3CR1 antibody to measure the ratio between CX3CR1+ and CX3CR1– islet cells is not the ideal quantitative approach. Therefore, to obtain more precise insight into CX3CR1 expression in primary α and β cells, we performed flow cytometry analyses in dispersed islet cells. As seen in Figure 5D, 97.7% of glucagon-positive and 97.1% of insulin-positive cells were CX3CR1+, showing that most of the primary α and β cells express CX3CR1.

High glucose levels suppress α cell glucagon release by the same cellular mechanisms whereby glucose stimulates GSIS: inhibition of KATP channel activity. Compared with β cells, α cell KATP channel conductance is much less (1%–10%) in the absence of glucose, and relatively less ATP is required to attenuate whole-cell KATP currents in α cells than in β cells (21). Therefore, unlike in β cells, the basal membrane potential in α cells is already depolarized (about −40 mV) even at relatively low levels of intracellular ATP, as seen in the CX3CR1 antibody to measure the ratio between CX3CR1+ and CX3CR1– islet cells is not the ideal quantitative approach. Therefore, to obtain more precise insight into CX3CR1 expression in primary α and β cells, we performed flow cytometry analyses in dispersed islet cells. As seen in Figure 5D, 97.7% of glucagon-positive and 97.1% of insulin-positive cells were CX3CR1+, showing that most of the primary α and β cells express CX3CR1.

High glucose levels suppress α cell glucagon release by the same cellular mechanisms whereby glucose stimulates GSIS: inhibition of KATP channel activity. Compared with β cells, α cell KATP channel conductance is much less (1%–10%) in the absence of glucose, and relatively less ATP is required to attenuate whole-cell KATP currents in α cells than in β cells (21). Therefore, unlike in β cells, the basal membrane potential in α cells is already depolarized (about −40 mV) even at relatively low levels of intracellular ATP, as seen in the CX3CR1 antibody to measure the ratio between CX3CR1+ and CX3CR1– islet cells is not the ideal quantitative approach. Therefore, to obtain more precise insight into CX3CR1 expression in primary α and β cells, we performed flow cytometry analyses in dispersed islet cells. As seen in Figure 5D, 97.7% of glucagon-positive and 97.1% of insulin-positive cells were CX3CR1+, showing that most of the primary α and β cells express CX3CR1.
low-glucose conditions. This is sufficient to facilitate AP firing and glucagon release (22). However, \( \alpha \) cell AP firing occurs only over a narrow range of membrane potentials, and further depolarization beyond a threshold level can inactivate voltage-gated sodium channels and attenuate AP amplitude (21). Therefore, upon high-glucose challenge, closure of most \( \alpha \) cell K\(_{ATP} \) channels blocks glucagon release. Therefore, we hypothesized that FKN might suppress glucagon release by inhibition of K\(_{ATP} \) activity in \( \alpha \) cells. As seen in

Figure 4. Electrophysiology studies in \( \beta \) cells. (A) K\(_{ATP} \) channel current in Min6 cells. U0216 (200 nM) was pretreated 1 hour before the measurements.

(B) Dose-dependent inhibition of K\(_{ATP} \) channel activity by FKN in the presence or absence of 200 nM U0126 or 10 nM PD98059. 

(C) ATP/ADP ratio in Min6 cells with or without FKN (100 ng/ml) treatment for 60 min. 

(D) Oxygen consumption rate in Min6 cells. Oligo, oligomycin (ATP synthase inhibitor); FCCP, trifluoromethoxy carbonylcyanide phenylhydrazone (a protonophoric uncoupler). 

(E) Plasma membrane potential (\( V_m \)) and AP firing in low glucose conditions in Min6 cells. A representative figure is presented (left). Bar graphs (right) represent average electrical activities 30 seconds after FKN application. 

(F) V\(_m \) and AP firing in high glucose conditions in Min6 cells. Min6 cells were incubated in 2 mM glucose for 16 h and challenged by high glucose. 5 minutes after, FKN was applied. A representative figure from 7 independent experiments (left). Bar graphs represent mean ± SEM. See also Supplemental Figure 4.
Figure 5. Electrophysiology studies in α cells. (A) Glucagon secretion in primary mouse islets. Islets were incubated in 1 mM glucose medium for 16 hours and washed. Islet glucagon (Gcg) release was measured after incubation with insulin (100 nM) or FKN (100 or 500 ng/ml) for 1 hour. Mean ± SEM; n = 5 for control and 6 for other groups. *P < 0.05 vs. lane 1; **P < 0.01 vs. lane 1; ***P < 0.05 vs. lane 2; 1-way ANOVA. (B) Glucagon secretion in αTC1 cells. αTC1 cells were incubated in 1 mM glucose medium for 16 hours and washed. Glucagon release was measured after incubation of the cells with FKN for 1 hour. Mean ± SEM; n = 3 (lane 4) or 7 (lane 1–3). *P < 0.05 vs. lane 1; #P = 0.051 vs. lane 1; 1-way ANOVA. (C) IHC analysis of CX3CR1 expression in primary WT mouse islets. Images were obtained at ×20 magnification. (D) Flow cytometry analysis of CX3CR1 expression in primary α and β cells in dispersed islet cells. Right lower box in the graph is for glucagon+/insulin– subset. Left upper box is for the glucagon–/insulin+ subset. (E) KATP currents in the presence or absence of FKN (100 ng/ml) in αTC1 cells. (F) KATP currents in the presence or absence of FKN (100 ng/ml) in primary α cells. (G) Membrane potential (V_m) and AP firing before and after FKN (100 ng/ml) treatment in αTC1 cells. Cells were incubated in 0.5 mM glucose medium, and depolarization of membrane potential (∆V_m), average AP peak, and AP frequency upon FKN treatment were analyzed. Ctl, control (before FKN treatment). Mean ± SEM; n = 7. **P < 0.01; 2-tailed unpaired t test. (H) Intracellular Ca²⁺ levels in the presence (n = 47) or absence (n = 31) of FKN. Lowering extracellular glucose concentration from 10 to 0.5 mM triggered Ca²⁺ rise in αTC1 cells. ***P < 0.01; 1-way ANOVA. See also Supplemental Figure 5.

Figure 5, E and F, FKN suppressed K_ATP currents in αTC1 and primary α cells, and this effect was sensitive to the U0126 MEK inhibitor (Supplemental Figure 5A). Moreover, inhibition of K_ATP activity by FKN was sufficient to depolarize membranes from ~40 mV to ~28 mV with a reduction of AP peak size (Figure 5G). Consistent with these results, FKN suppressed the low-glucose-induced increase in intracellular Ca²⁺ levels (Figure 5H). These effects of FKN occurred without changes in cAMP levels (Supplemental Figure 5B).

Discussion
In this study, we demonstrate that chronic administration of a long-acting form of FKN, FKN-Fc, exerts durable effects to
improve glucose tolerance without changing body weight. This is associated with a direct effect of FKN-Fc to improve GSIS and decrease β cell apoptosis. In addition, we found that FKN-Fc treatment led to a tissue-specific improvement in hepatic insulin sensitivity. From a mechanistic point of view, we traced this latter effect to direct actions of FKN-Fc to decrease α cell glucagon secretion and inhibit hepatocyte glucagon action both in vitro and in vivo. As such, these findings demonstrate that chronic long-acting FKN treatment leads to improved glucose tolerance due to enhanced insulin secretion, decreased glucagon secretion, and improved hepatic insulin sensitivity. Taken together, these studies suggest the possibility that the FKN/FKN receptor axis (CX3CL1/CX3CR1) could be a new area for antidiabetic drug discovery, which can improve both insulin secretion and insulin sensitivity.

We previously reported that β cells express CX3CR1 and FKN increases insulin secretion by an ERK-dependent increase of the intracellular Ca²⁺ level (27). However, the molecular mechanisms of how FKN stimulates intracellular Ca²⁺ level were unknown. Since oxygen consumption rate in islet cells was not changed by FKN treatment, or deletion of CX3CR1, and mitochondrial content and morphology was normal in CX3CR1−KO mouse islets (27), it is possible that the FKN/CX3CR1 system enhances GSIS and intracellular Ca²⁺ levels by modulating the electrical activity of β cells. Consistent with this, we found that FKN treatment did change cellular ATP levels, the ATP/ADP ratio, and oxygen consumption rates in both low- and high-glucose conditions. To further address this question, we used electrophysiological studies to measure K_{ATP} channel and VDCC activity directly in β cells in the presence and absence of FKN. These experiments revealed that FKN inhibits K_{ATP} channel activity through an ERK-dependent and ATP-independent mechanism, without effects on VDCC activity. It is known that glucose triggers the insulin secretory pathway by increasing the β cell ATP/ADP ratio and inhibiting K_{ATP} channel activity (21, 32). This leads to membrane depolarization with opening of VDCCs, leading to a rise in intracellular Ca²⁺ levels and exocytosis of insulin secretory granules (21). Notably, in low-glucose conditions, FKN treatment-triggered AP firing did not lead to a substantial increase in insulin secretion. This is consistent with the literature that shows that increased intracellular Ca²⁺ levels induced by voltage-gated L-type Ca²⁺ channel (VDCC) activator treatment, K⁺ overload, K_{ATP} channel inhibitor treatment, or arginine treatment are not sufficient to stimulate insulin secretion, if not accompanied by increased β cell metabolism or other events normally induced by glucose (33–36). In T2DM, it has been suggested that decreased β cell glucose responsiveness is associated with lower mitochondrial ATP production capacity, leading to incomplete inhibition of K_{ATP} channel activity (21, 23, 32, 37, 38). Therefore, it is reasonable to suggest that the action of FKN to inhibit basal K_{ATP} channel conductance may compensate for lower mitochondrial ATP production in type 2 diabetic β cells and, thus, enhance GSIS.

Similarly to β cells, α cells also have an intrinsic glucose-sensing mechanism through K_{ATP} channel regulation. It has been suggested that a progressive and time-dependent increase in α cell K⁺ channel activity may be associated with the lack of glucagon suppression by glucose in T2DM (21, 22). Indeed, inhibition of K_{ATP} channel activity leads to partial restoration of glucose-dependent suppression of α cell glucagon release in islets from T2DM subjects (22). Interestingly, flow cytometry analyses revealed that CX3CR1 is detected in more than 90% of primary mouse α cells, and incubation of αTC1 cells with FKN inhibited low-glucose-induced glucagon release. Together, these results suggest that FKN can suppress glucagon release in a cell-autonomous mechanism. This is consistent with our findings that FKN-Fc treatment reduces circulating plasma glucagon levels. However, it is possible that the FKN effect to inhibit glucagon secretion is indirect. For example, FKN can increase insulin secretion, and insulin can suppress α cell glucagon release (27, 39, 40). Indeed, while we found that over 97% of α cells expressed CX3CR1, some years ago, Thorel et al. reported that 98% α cell depletion does not affect glucose tolerance, suggesting that only 2% of α cells may be sufficient to maintain adequate glucagon levels (41). While the effects of FKN were ERK-dependent, we do not know whether the role of ERK is direct or indirect, or whether this involves a specific K_{ATP} channel phosphorylation event.

An important component of β cell dysfunction in T2DM is decreased overall β cell mass, which advances in the later stage of the disease (42). While increased β cell apoptosis may be responsible for the majority of decreased β cell mass in advanced T2DM patients (42, 43), recently it has been suggested that an additional mechanism may exist. Thus, accumulation of dedifferentiated/imature β cells, which express markers of both α and β cells, has been reported in mouse models of diabetes as well as in human T2DM islets (4, 32, 43–45). While the origin of these dedifferentiated/imature β cells is still controversial in humans (32, 43), it is plausible that strategies to suppress β cell apoptosis and/or increase β cell differentiation/maturation would be effective at improving β cell function. Interestingly, chronic FKN-Fc administration enhanced GSIS and decreased β cell apoptosis. This was accompanied by increased β cell mass and the expression of the genes involved in β cell differentiation and function. It should be noted that chronic FKN-Fc administration began at 10 weeks of HFD, at which point β cell dysfunction and dedifferentiation already exist (46, 47).

Our findings that long-term FKN treatment leads to decreased insulin resistance due to a tissue-specific effect to improve hepatic insulin sensitivity represent an additional component of the antidiabetic effects of FKN. Glucagon is a key counterregulatory hormone to insulin, and stimulates AMPK production with activation of the protein kinase A (PKA) pathway, ultimately leading to increased endogenous glucose production. Postprandial increases in blood glucose and insulin levels normally suppress α cell glucagon secretion and inhibit hepatic glucagon action. In T2DM, a lack of glucagon suppression leads to elevated levels of basal and postprandial glucagon, contributing to hyperglycemia (48–52). There are many studies showing the efficacy of inhibitory glucagon receptor antibodies to improve glucose tolerance in preclinical models and in some clinical studies (12, 53–55). Thus, the concept of targeting glucagon action to achieve hepatic insulin sensitivity and glucose lowering is well accepted (14), even though current methods of glucagon receptor inhibition may have unwanted side effects (56). Using hyperinsulinemic-euglycemic clamp studies, we determined that chronic FKN-Fc treatment specifically improves hepatic...
insulin sensitivity in HFD/obese mice. We also show that this effect was direct and cell-autonomous by finding that FKN treatment of primary mouse hepatocytes directly inhibits glucagon action with a corresponding decrease in cAMP levels and PKA activity. Furthermore, all of these effects were pertussis toxin-dependent, but not sensitive to MEK inhibitors, demonstrating that the FKN/CX3CR1 pathway in hepatocytes signals through a classical Gαi-coupled pathway to inhibit adenylate cyclase activity. It is interesting to note that FKN treatment decreased hepatic glucagon action without causing a compensatory increase in plasma glucagon levels or a cell hyperplasia, as occurs in glucagon receptor–KO mice or after chronic treatment with a glucagon receptor antagonist (55, 57). Taken together with previously reported effects of the FKN/CX3CR1 system to improve hepatic fibrosis (58, 59), our results support the idea that chronic FKN administration can improve glucose tolerance and hepatic function through direct hepatic actions.

Sindhu et al. have reported a 45% increase in circulating soluble FKN levels in type 2 diabetic patients (60). However, other reports have found no significant increase (61). It should be noted that, to the extent that circulating FKN levels might be elevated in type 2 diabetics, this might raise the possibility of FKN resistance. Of course, the issue of FKN sensitivity or resistance in human subjects is entirely unknown and will await clinical trials for an answer.

Here, we report that chronic FKN-Fc administration exerts durable effects to improve glucose tolerance by enhancing insulin secretion and suppressing glucagon secretion. In addition, FKN-Fc administration leads to improved hepatic insulin sensitivity through direct hepatocyte-selective effects to inhibit glucagon action. These results raise the possibility that the FKN/CX3CR1 system could be targeted for antidiabetic drug discovery efforts.

Methods

**Animals and treatments.** Seven-week-old male WT C57BL/6J, Cx3cr1<sup>−/−</sup> knockin mice and B6.Cg-Tg(Insl1-EGFP)1Hara/J mice were obtained from The Jackson Laboratory and fed normal chow diet (NCD) until they were subjected to 60% HFD (Research Diets Inc.). For acute and chronic FKN-Fc injection studies, mice were randomly separated into 2 groups and subjected to vehicle or FKN-Fc administration. When 10 mg/kg FKN-Fc was injected i.p. into normal mice, the FKN-Fc level peaked at 598.9 μM in 4 hours and decreased to 31.6 μM in 24 hours. Since the dose of 100 ng/ml native FKN (chemokine domain) used in vitro studies corresponds to 10.5 μM, 10 mg/kg dose was used in the every-other-day injection protocol in NCD mice. We used 30 mg/kg in HFD mice to assure maximal effects. Intraportal glucagon and insulin tolerance tests were performed as described previously (62, 63). Mouse clamp experiments were performed as described previously (63, 64). During the clamps, insulin was infused at a constant rate of 8.0 mU/kg/min.

For testing acute effects of FKN-Fc administration, mice were fasted for 6 hours, i.p. injected with glucose (1 g/kg) alone or glucose (1 g/kg) + FKN-Fc (10 mg/kg) and blood glucose and plasma insulin levels were measured. For chronic FKN-Fc treatment studies, WT or CX3CR1 KO mice were fed HFD for 10 weeks and then i.p. injected with 30 mg/kg FKN-Fc every other day for an additional 8 weeks. 6 hours before measuring glucose tolerance and serum insulin levels, mice were given a final FKN-Fc administration and the food was withdrawn.
mouse fractalkine (mFKN), 1 mM glucose KRB + 100 nM GLP-1, 25 mM glucose KRB, or 1 mM glucose KRB + 10 ng/ml (or 100 ng/ml) mFKN + 100 ng/ml pertussis toxin. After the incubation period, the Bridge-It Cyclic AMP Designer Assay was used to determine intracellular CAMP levels per the manufacturer’s instructions (catalog 122934; Mediomics LLC).

Primary mouse hepatocyte isolation and in vitro glucose production assay. Mice were infused through the inferior vena cava with a calcium-free HEPES-phosphate buffer (pH 7.4) for 10 minutes followed by a collagenase solution (Liberase TM; Roche) for 10 minutes. The digested livers were excised, and hepatocytes were collected and washed 5 times in buffer by centrifuging at 70 g for 5 minutes. Cells were further purified by centrifugation (2,400 g for 10 minutes) over a Percoll density gradient (1.06 g/ml). Primary mouse hepatocytes were allowed to attach for 6 hours on collagen-coated plates in Williams’ Medium E (Catalog 12551-032; Life Technologies) fortified with nonessential amino acids, GlutaMAX (catalog 35050-061; Life Technologies), antibiotics, 10% FBS, and dexamethasone (10 nM) and cultured overnight in the same medium without serum. Cultures were then washed in HEPES phosphate-salt-bicarbonate (HPSB) buffer (10 mM NaH2PO4, 125 mM NaCl, 0.85 mM KH2PO4, 1.25 mM Na2HPO4, 1 mM MgCl2, 1 mM CaCl2, and 15 mM NaHCO3) containing 0.2% free fatty acid–free BSA, and incubated in the same buffer containing FKN, insulin (10 nM), and/or glucagon (10 ng/ml) and substrates for 3 hours in a 5% CO2 incubator. 14C-pyruvate (2 mM, 0.5 μCi pyruvate per incubation) was used as substrate. Incubations were carried out in 0.5 ml buffer in 24-well plates containing 0.25 million cells per well. At the end of incubation, the buffer solutions were transferred to 1.7-ml microcentrifuge tubes and added with 0.25 ml 5% ZnSO4 and 0.25 ml 0.3 N Ba(OH)2 suspensions to each tube, followed by addition of 0.5 ml water. After centrifugation, supernatants were transferred to a fresh set of tubes and assayed for radiolabeled glucose released into the medium by separation of radiolabeled glucose by mixed-bed ion-exchange resins, AG-50X8 resins (Bio-Rad). Two hundred milligrams of resins were added to each tube, vortexed intermittently for 15 minutes, and centrifuged, and the supernatants were transferred to scintillation vials for counting radioactivity. Cells on the plates were dissolved in IN NaOH for protein estimation.

Tissue lipid content measurement. Levels of lipid contents in liver samples were measured as described previously (63).

Electrophysiology. For single-cell experiments, Min6 or αTC1 cells were cultured on poly-L-lysine–coated coverglass chips and used 2 days after plating. Cells were transferred to culture medium containing 0.5 or 2 mM glucose 16 hours before experiments. KATP channel activity was measured 2 days after plating. Cells were transferred to culture medium containing 10 mM CaCl2. For time course measurements of VDCC activity, the external solution for identifying islet cell types contained (in mM): 120 NaCl, 2.5 KCl, 1 MgCl2, 0.5 d-glucose, 10 HEPES, 10 tetraethylammonium chloride (TEA-Cl), pH 7.4 adjusted with NaOH. VDCC activity in Min6 cells was recorded using the whole-cell patch-clamp technique with an EPC 9 patch-clamp amplifier (HEKA Elektronik). Ca2+ channels were activated by voltage steps from ~80 mV to various membrane potentials (~70 to +20 mV) with 10 mV increments with an extracellular solution containing 10 mM CaCl2. For time course measurements of VDCC activity, the external solution contained (in mM): 130 NaCl, 2.5 KCl, 2 CaCl2, 1 MgCl2, 10 TEA-Cl, 10 HEPES, and 10 d-glucose, pH 7.2 adjusted with NaOH. The 2 mM CaCl2 was replaced by 10 mM CaCl2 to enhance Ca2+ channel currents. The internal pipette solution contained (in mM): 130 Cs-glutamate, 10 CsCl, 10 HEPES, 1 MgCl2, 5 EGTA, 5 MgATP, and 0.3 NaGTP, pH 7.2 adjusted with CsOH. Membrane potential in Min6 and αTC1 cells was recorded in the current clamp mode of perforated whole-cell configuration in an internal solution containing (in mM): 76 K+SO4, 10 KCl, 10 NaCl, 1 MgCl2, 5 HEPES, and 0.25 mg/ml amphotericin B, pH 7.35 adjusted with KOH. Extracellular solutions consisted of (in mM): (a) 2 mM glucose external solution for Min6 cells: 135 NaCl, 2.5 KCl, 2 CaCl2, 1 MgCl2, 10 EGTA, and 0.6 Na-ATP, pH 7.2 adjusted with KOH. The current was expressed as current density (I, pA/μF) to compensate for different cell sizes. Recordings were performed at room temperature (22°C-24°C).
The recordings were semi-automatically analyzed using a macro written in Igor (WaveMetrics). For \( \Delta V_m \) calculation in Min6 cells, average \( V_m \) for 30 seconds right before high glucose challenge was subtracted from the average \( V_m \) during 1 minute starting 1 minute after first AP spike or 30 seconds after FKN application. Averages of AP frequency and peak size were calculated for the same 1 minute period of time or 30 seconds after FKN application.

**Fluorescence measurements of intracellular \( \text{Ca}^{2+} \).** To measure \([\text{Ca}^{2+}]_i\), in single cells, \( \alpha \)-TCI cells were loaded with 4 \( \mu \)M \( \text{Ca}^{2+}\)-sensitive Fura-2 AM for 30 minutes in the presence of 0.01% pluronic F-127 and 100 \( \mu \)M sulfipyrazone. Fura-2 dye was excited at 340 and 380 nm using a Polychrome IV (TILL Photonics, FEI Life Sciences), and the emission fluorescence was recorded at greater than 470 nm every 2 seconds using an EMCCD camera (Photometrics Technology) in an inverted Nikon TE2000 microscope equipped with a 20x objective lens. A cell-free region was used for background fluorescence correction. The external solutions contained the following (in mM): 10 mM glucose external solution: 135 NaCl, 3.6 KCl, 1.5 CaCl\(_2\), 0.5 MgSO\(_4\), 0.5 NaH\(_2\)PO\(_4\), 5 NaHCO\(_3\), 10 d-glucose, and 10 HEPES, pH 7.4 adjusted with NaOH; for 0.5 mM glucose external solution, 140 mM NaCl was used, instead of 135 mM. The \( \text{Ca}^{2+} \) measurements were performed similarly to current recordings (66).

**Relative apoptotic activity assays in primary mouse islet cells.** Primary mouse islets were incubated with trypsin solution, and the dispersed islet cells were plated. After overnight incubation for adhesion, cells were washed and incubated in low-glucose (2.8 mM) or high-glucose (16.7 mM) DMEM supplemented with 0.2% free fatty acid–free BSA or 400 \( \mu \)M palmitate-conjugated BSA in the presence or absence of FKN-Fc treatment for 48 hours. Relative apoptotic activity was calculated by normalization of caspase-3/7 activity by viable cell activity in each well.

**Oxygen consumption.** Min6 cells were plated at 20,000 per well in XF96 plates for 48 hours. Culture medium was changed to lower-glucose (2 mM) medium the night prior to the experiment. Two hours before the assay, the medium was changed to 120 mM NaCl, 3.5 mM KCl, 2.3 mM CaCl\(_2\), 0.04 mM KH\(_2\)PO\(_4\), 1 mM MgCl\(_2\), 0.1% BSA, 5 mM HEPES, pH 7.4 at 37°C, and 0.25 mM glucose. Successive additions from the ports were 16.45 mM glucose, 2 \( \mu \)M oligomycin, 1 mM FCCP, and then 0.25 \( \mu \)M rotenone plus 1 \( \mu \)M antimycin A.

**Statistics.** Statistical methods were not used to predetermine necessary sample size, but sample sizes were chosen based on estimates from pilot experiments and previously published results such that appropriate statistical tests could yield significant results. Statistical analyses used in the data presented are justified and described in all legends. Parametric tests were used that assume normal distribution, which we showed to be the case when data were plotted as frequencies. Variances were tested by Levene’s test for homogeneity of variance, and variances in the data were not significantly different. Experiments were not performed in a blinded fashion. The results are shown as means ± SEM. All statistical analysis was performed by 2-tailed Student’s t test or ANOVA, unless indicated; \( P \) less than 0.05 was considered significant. A representative figure for each experiment is presented without combining of data from different batch experiments, unless indicated in the figure legend.

**Study approval.** All animal procedures were performed in accordance with an IACUC-approved protocol and the research guidelines for the use of laboratory animals of the University of California, San Diego.

**Author contributions.** MR performed and analyzed in vivo mouse studies including glucose and insulin tolerance tests, tissue immunohistochemistry experiments, ELISA analyses of plasma hormones, ex vivo islet experiments, and in vitro Min6 and \( \alpha \)-TCI cell experiments. JBS performed patch clamp experiments and intracellular \( \text{Ca}^{2+} \) measurements. GKB performed ex vivo primary hepatocyte experiments and tissue lipid content measurements. PL performed mouse clamp experiments. JW, HC, and AP supported in vivo mouse experiments. AM measured oxygen consumption. SP and JB produced and tested activity of the recombinant FKN-Fc protein. DB and AF performed in vivo pharmacokinetics studies. DSK supported analyses and interpretation of patch clamp experiments. SRJ performed and supported patch clamp experiments. MW, RDJ, and CJL supported design of the study and analysis of the data. JMO and YSL designed the study, supervised the project, analyzed and interpreted the data, and wrote the paper. All authors discussed the results and commented on the manuscript.

**Acknowledgments.** This study was funded by the US National Institute of Diabetes and Digestive and Kidney Diseases (DK074868, DK063491, and DK101395), a fellowship from the American Heart Association (to MR), the TJ Park Science fellowship of POSCO TJ Park Foundation, and a grant from Takeda California Inc.

Address correspondence to: Yun Sok Lee, Stein Clinical Research Building, Room 231, 9500 Gilman Drive, MC# 0673, La Jolla, California 92093, USA. Phone: 858.822.6641; Email: yunsoklee@ucsd.edu. Or to: Jerrold Olefsky, Stein Clinical Research Building, Room 227, 9500 Gilman Drive, La Jolla, California 92093, USA. Phone: 858.534.6651; Email: jolefsky@ucsd.edu.

---

10. Raskin P, Unger RH. Hyperglucagonemia and its...


