

Autoregulation of insulin receptor signaling through MFGE8 and the $\alpha v \beta 5$ integrin

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The role of integrins, in particular αv integrins, in regulating insulin resistance is incompletely understood. We have previously shown that the $\alpha v\beta 5$ integrin ligand milk fat globule epidermal growth factor like 8 (MFGE8) regulates cellular uptake of fatty acids. In this work, we evaluated the impact of MFGE8 on glucose homeostasis. We show that acute blockade of the MFGE8/β5 pathway enhances while acute augmentation dampens insulin-stimulated glucose uptake. Moreover, we find that insulin itself induces cellsurface enrichment of MFGE8 in skeletal muscle, which then promotes interaction between the $\alpha v \beta 5$ integrin and the insulin receptor leading to dampening of skeletal-muscle insulin receptor signaling. Blockade of the MFGE8/β5 pathway also enhances hepatic insulin sensitivity. Our work identifies an autoregulatory mechanism by which insulin-stimulated signaling through its cognate receptor is terminated through up-regulation of MFGE8 and its consequent interaction with the $\alpha v\beta 5$ integrin, thereby establishing a pathway that can potentially be targeted to improve insulin sensitivity.

integrins | MFGE8 | insulin sensitivity | insulin receptor | insulin signaling

A cute insulin resistance can be viewed as a protective response under specific physiological conditions that necessitate increased insulin secretion. Nevertheless, the increasing prevalence of chronic insulin resistance (1) in the current obesity epidemic hastens the development of type 2 diabetes (T2D) and induces compensatory hyperinsulinemia. Hyperinsulinemia can produce potentially maladaptive consequences at least in part, due to the mitogenic roles of insulin (2–4). As such, there remains a critical need for new therapies to improve insulin sensitivity in order to prevent T2D, avoid the need for insulin treatment in patients with T2D, or reduce the insulin dose required to normalize blood glucose in such individuals.

Insulin binding to the alpha subunit of the insulin receptor induces a conformational change that triggers activation of insulin receptor beta subunit (IR β) tyrosine kinase activity (5–7). The activated insulin receptor phosphorylates target molecules that mediate downstream signaling leading to glucose uptake and other metabolic effects (8, 9). Dephosphorylation of IR β and insulin receptor substrate-1 (IRS-1) aids in termination of insulin signaling pathways (10, 11) and is the basis of clinical trials targeting putative phosphatases to treat diabetes (12). Despite their potential therapeutic relevance, there is a relative paucity of knowledge regarding molecular mechanisms that lead to termination of insulin receptor signaling.

The integrin families of cell surface receptors mediate bidirectional signaling between the cell and its external environment. Previous work has identified interactions between integrin receptors and other growth factor receptor tyrosine kinases (13–16) that lead to modulation of downstream signaling (17–19). For example, the $\alpha\nu\beta3$ and $\alpha\beta\beta4$ integrins function as coreceptors for insulin-like

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growth factor-1 and 2 (IGF1 and 2) and potentiate IGF1 receptor (IGF1R)-mediated signaling (19–23). Immunoprecipitation studies have demonstrated a physical association between the αv integrins and IR β (24, 25). The impact of these associations on glucose homeostasis has not been evaluated. A role for β 1 integrins in the regulation of glucose homeostasis is well established. This class of integrins appears to be particularly important in regulating insulin-mediated glucose homeostasis in the obese state. The effect of β 1 integrins on glucose homeostasis appears to be primarily due to obesity-associated matrix remodeling (26–30) rather than a direct effect secondary to a physical association between β 1 integrins and the insulin receptor.

Milk fat globule epidermal growth factor like 8 (MFGE8) is a secreted integrin ligand which binds the $\alpha\nu\beta3$, $\alpha\nu\beta5$, and $\alpha8\beta1$ integrins (31, 32). Several recent observations suggest a role for MFGE8 in modulating insulin resistance. In humans, serum MFGE8 levels are increased in the context of diabetes and correlate positively with the extent of hemoglobin glycosylation (33, 34). Indeed, serum MFGE8 levels correlate with indices of insulin resistance in two independent cohorts of patients with T2D or gestational diabetes from China (35, 36). A missense variation in the gene encoding MFGE8, present in South Asian Punjabi Sikhs, is associated with increased circulating MFGE8 levels and increased risk of developing T2D (37). Increased circulating levels of MFGE8 in diabetic patients may impact T2D through effects

Significance

The increasing prevalence of insulin resistance in the current obesity epidemic creates a significant need for ongoing investigations into the mechanisms that modulate insulin resistance. Long-term treatment of diabetic patients with high doses of insulin, especially in those that are obese, may cause further weight gain and exacerbate the risk of developing cancers. The work presented here demonstrates a previously unappreciated regulatory pathway by which insulin initiates dampening and/ or termination of its own signaling pathway through an integrin-dependent pathway. These data identify a pathway that can potentially be targeted to reduce insulin requirements in patients with diabetes.

The authors declare no competing interest.

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on inflammation and cardiovascular disease. Humans with increased MFGE8 expression have a greater risk of developing coronary artery disease (38). In contrast, in murine models, MFGE8 deficiency exacerbates cardiac hypertrophy and atherosclerosis (39, 40). MFGE8 also improves wound healing responses in diabetic foot ulcers (41, 42) by triggering apoptotic cell clearance and promoting resolution of inflammation (43–45).

Despite the notable links between MFGE8, insulin resistance, and T2D pathology, the biology underlying these associations has not been investigated. We therefore evaluated the effect of acute antibody-mediated disruption of the MFGE8/ β 5 pathway on glucose homeostasis in wild-type (WT) mice. We report here that MFGE8 markedly attenuates the effect of insulin on skeletal muscle glucose uptake. Antibody-mediated blockade of MFGE8 or $\alpha\nu\beta5$ enhances while recombinant MFGE8 (rMFGE8) reduces insulin-stimulated glucose uptake in vitro and in vivo. Mechanistically, insulin acts to promotes cell-surface enrichment of skeletal muscle MFGE8, which then binds to cell surface $\alpha\nu\beta5$ and increases the interaction between the integrin and the insulin receptor. This interaction subsequently aids in terminating insulin receptor signaling.

Results

MFGE8 and the $\alpha v\beta 5$ Integrin Dampen Insulin-Mediated Glucose Uptake. To determine whether the MFGE8/ β 5 pathway regulates glucose homeostasis, we performed a glucose tolerance test (GTT) in 8 wk old male WT mice in the presence of β 5 blocking or isotype control antibody. 65 blockade significantly reduced serum glucose levels in WT mice as compared with isotype control antibody (Fig. 1A). Of note, antibody-mediated blockade of \$65 did not affect glucose clearance in Mfge8-/- mice, indicating that MFGE8 functions upstream of $\beta 5$ in regulating glucose uptake. Furthermore, *Mfge8*⁻ mice used in the studies did not have altered glucose clearance in the GTT or body mass composition as compared with WT controls (Fig. 1A and SI Appendix, Fig. S1). Insulin levels during the GTT in the setting of $\beta 5$ blockade correlated with blood glucose levels, suggesting that enhanced glucose clearance was not due to a direct effect of the blocking antibody on pancreatic insulin release (Fig. 1B). A GTT in WT mice conducted after intraperitoneal (IP) administration of an MFGE8 blocking antibody significantly enhanced (Fig. 1C) glucose clearance. By contrast, IP administration of rMFGE8 significantly blunted glucose clearance compared to treatment with control MFGE8 construct (RGE), carrying a single

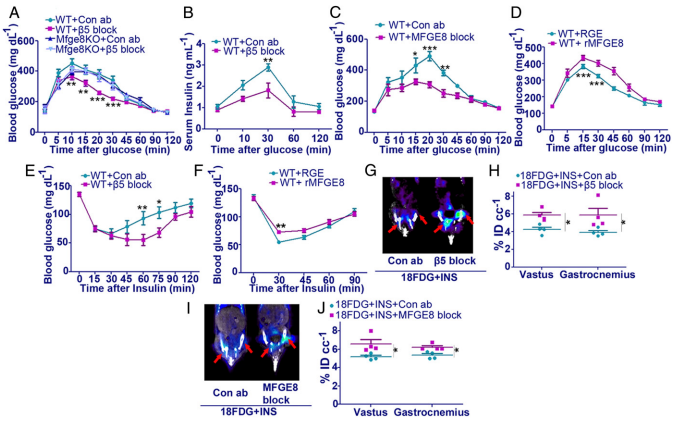


Fig. 1. MFGE8 regulates insulin-induced skeletal-muscle glucose uptake in vivo. (*A*) GTT in 7 to 8 wk old male WT and *Mfge8* ^{-/-} mice after IP injection of $\beta5$ blocking ($\beta5$ block) or isotype control antibody (Con ab). n = 3 to 4 for WT mice and n = 3 for *Mfge8* ^{-/-} mice per group per experiment and merged data from two to four independent experiments are presented. Statistical analysis compares the $\beta5$ blocking and control antibody groups in WT mice. (*B*) Serum insulin levels during GTT in WT mice. n = 3 to 4 mice per group per experiment, and merged data from two independent experiments are presented. (*C*) GTT in male WT mice after IP injection of MFGE8 blocking (MFGE8 block) or isotype control antibody (Con ab). n = 3 mice in each group per experiment, and merged data from two independent experiments are presented. (*D*) GTT in WT male mice after IP injection of recombinant MFGE8 (rMFGE8) or RGE control construct. n = 2 to 3 mice per group per experiment, and merged data from three independent experiments are presented. (*E*) ITT in WT male mice treated with $\beta5$ blocking or control antibody. n = 3 to 4 mice per group per experiment, and the merged data from four independent experiments are presented. (*F*) ITT after IP administration of rMFGE8 or RGE control in WT male mice. n = 2 to 3 mice per group per experiment, and merged data from three independent experiments, are presented. (*G* and *I*) PET/CT scan images showing radioactive 18FDG deposition (red arrowheads) in 7 to 8 wk old WT male mice treated with $\beta5$ blocking, (G) MFGE8 blocking, (I) or control antibody in presence of insulin (1 *U/kg*) and 18FDG. n = 4 independent experiments. (*H* and *J*) Quantification on 18FDG deposition in vastus and gastrocnemius skeletal-muscle compartments in the setting of $\beta5$ (*H*) or MFGE8 (*J*) blockade. Data are expressed as % injected dose of 18FDG per cubic centimeter (cc) tissue (%ID cc⁻¹). All data expressed as mean \pm SEM; **P* < 0.05, ***P* < 0.01, and ****P* < 0.001. Data

amino acid mutation that changes the RGD sequence to RGE (46) and precludes integrin binding (Fig. 1*D*).

The MFGE8/ β 5 Pathway Modulates Systemic and Skeletal Muscle Insulin Sensitivity. We next performed insulin tolerance tests and found that systemic β 5 blockade significantly improved insulin tolerance (Fig. 1*E*) while treatment with rMFGE8 had the opposite effect (Fig. 1*F*) in WT male mice. Moreover, systemic β 5 blockade significantly improved both glucose and insulin tolerance tests in female mice, indicating that the effects of targeting β 5 generalizes across sex. (*SI Appendix*, Fig. S2 *A* and *B*). Taken together, these data suggest that the MFGE8/ β 5 pathway modulates insulin sensitivity in vivo.

To evaluate the effect of the MFGE8/ β 5 pathway on skeletal muscle insulin sensitivity, we treated WT mice with 18-fluoro-deoxy glucose (18FDG) and insulin after administering β 5-blocking, MFGE8-blocking, or isotype control antibodies and then performed positron emission tomography/computed tomography (PET/CT) scanning. Systemic treatment with β 5- or MFGE8-blocking antibodies significantly enhanced insulin-stimulated 18FDG uptake in hind-leg skeletal muscles (vastus lateralis, gastrocnemius). These data indicate that disruption of the MFGE8/ β 5 pathway enhances skeletal muscle glucose uptake in vivo (Fig. 1 *G–J*).

We tested whether these effects are cell intrinsic by examining the insulin-responsive uptake of a nonhydrolyzable fluorescent glucose analog, 2NBDG, in primary WT and Mfge8^{-/-} skeletal muscle myotubes, human skeletal muscle myotubes (HskM), and differentiated C2C12 myotubes. In each system, rMFGE8 treatment dampened while 65 integrin blockade enhanced the ability of insulin to stimulate 2NBDG uptake. (Fig. 2 A-C). Moreover, $Mfge8^{-/-}$ myotubes had greater insulin-stimulated 2NBDG uptake than WT myotubes. This effect was reversed by treating the Mfge8^{-/-} myotubes with rMFGE8, further supporting the concept that MFGE8 is a direct inhibitor of insulin-stimulated glucose uptake. By contrast, neither fibronectin nor vitronectin, two unrelated avβ5 integrin ligands, affected insulin-stimulated 2NBDG uptake in C2C12 myotubes (SI Appendix, Fig. S3), indicating that the effect of avß5 integrin on insulin-stimulated glucose uptake is specific to MFGE8. To further characterize these findings, we evaluated the effect of $\beta 5$ blocking antibody on glucose uptake in response to a dose range of insulin in C2C12 myotubes. We found a similar and significant effect of $\beta5$ blockade on glucose uptake at insulin doses of 100 and 200 nM (Fig. 2D). We also performed a time course experiment in this system and found a significant effect of $\beta 5$ blockade on glucose uptake beginning 30 min after insulin administration that persisted at 45 and 60 min after insulin administration (Fig. 2E).

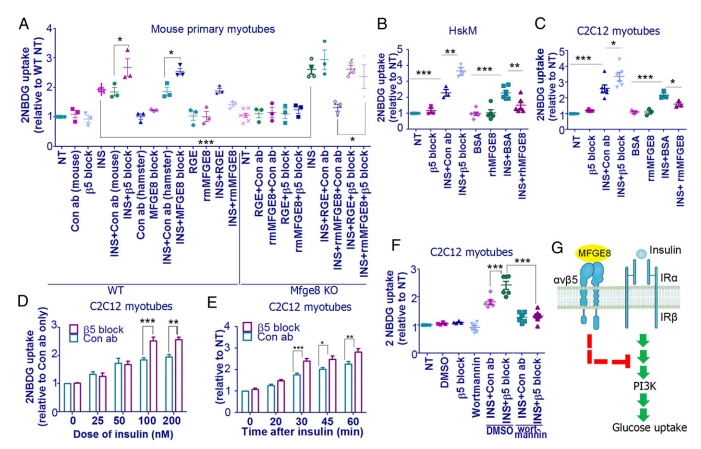


Fig. 2. MFGE8 regulates insulin-induced glucose uptake in vitro. (*A*–*C*) 2NBDG uptake assay in mouse primary *WT* and *Mfge8^{-/-}* myotubes (*A*, *n* = 3 independent experiments), primary HsKM (*B*, *n* = 3 to 6 independent experiments), and differentiated C2C12 myotubes (*C*, *n* = 3 to 6 independent experiments) in the presence or absence of rMFGE8 or RGE control protein, β 5 blocking, MFGE8 blocking, or isotype control antibody and insulin (100 nM). Data are expressed as relative fold changes compared to the untreated cells (NT). (*D* and *E*) Dose response (*D*, *n* = 4 to 6 independent experiments) and time course of insulin action (*F*, *n* = 6 independent experiments) on 2NBDG uptake in C2C12 myotubes in the presence of either β 5 blocking or control antibody. Data are expressed as relative fold changes compared to control antibody-treated cells in the absence of insulin. (*F*) 2NBDG uptake assay in β 5 blocking antibody-treated C2C12 cells in the presence and absence of insulin and wortmannin. Data are expressed as relative fold changes compared to classe (*N*), and ****P* < 0.001. Data analyzed by one-way ANOVA followed by Bonferroni's posttest (*A*, *D*, and ****P* < 0.001. Data analyzed by one-way ANOVA followed by Bonferroni's posttest (*A*, *D*, and dampens insulin-stimulated glucose uptake.

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Insulin stimulates glucose uptake in the myotubes via canonical activation of PI3 kinase pathway (47–50). To determine whether the enhanced insulin sensitivity by $\beta5$ blockade is sensitive to PI3K inhibition, we examined insulin-stimulated 2NBDG uptake in C2C12 myotubes treated with $\beta5$ blocking antibody in the presence of PI3K inhibitor wortmannin. Wortmannin significantly dampened insulin-stimulated 2NBDG uptake (Fig. 2*F*). $\beta5$ blockade failed to augment insulin-stimulated glucose uptake in the presence of wortmannin, indicating that $\beta5$ blockade potentiates the PI3K/AKT pathway to trigger glucose uptake in these cells (Fig. 2*F*). Taken together, these data indicate that MFGE8 and $\beta5$ integrin limit insulin-stimulated glucose uptake via regulation of canonical insulin signaling pathway in a cell-intrinsic manner (Fig. 2*G*).

To investigate whether augmented insulin signaling due to $\beta5$ blockade promotes tumor growth, we assessed cell proliferation of MCF-7 human breast-cancer cells treated with $\beta5$ blocking antibody in the presence or absence of insulin. Incubation of MCF-7 cells with $\beta5$ blocking antibody (with or without insulin administration) for 6 h induced marked cell death, as measured by propidium iodide staining followed by flow cytometry, mitigating concerns of $\beta5$ blockade promoting tumor growth (*SI Appendix*, Fig. S4).

MFGE8 and the $\alpha v\beta 5$ Integrin Regulate Insulin Receptor Signaling in Skeletal Muscle. We next investigated whether the MFGE8/65 integrin pathway regulates insulin receptor signaling. We measured the impact of antibody-mediated ß5 blockade on insulin-stimulated tyrosine phosphorylation of IR β and IRS-1 in C2C12 myotubes. We pretreated C2C12 myotubes with β 5 blocking or control antibody for 1 h, followed by insulin for 5 or 30 min. Subsequent immunoprecipitation of protein lysates using an anti-phospho-tyrosine antibody, followed by Western blotting for IR^β and IRS-1, demonstrated that $\beta5$ blockade enhanced insulin-stimulated tyrosine phosphorylation of IR β and IRS-1 at both time points (Fig. 3A). We repeated experiments at the 30 min time point with a dose range of insulin (25, 50, 100, and 200 nM) and observed enhanced tyrosine phosphorylation of IR β and IRS-1 at each of these doses after β 5 blockade (Fig. 3B and SI Appendix, Fig. S5 A-C). Additionally, treatment of C2C12 myotubes with rMFGE8 but not RGE control construct blunted insulin-stimulated tyrosine phosphorylation of IRß and IRS-1, indicating that the effect of rMFGE8 treatment is dependent on integrin binding (Fig. 3C).

To extend these findings in vivo, we administered β 5 blocking or control antibody to WT mice 1 h before treatment with IP insulin, harvested hind-leg skeletal muscles 15 and 60 min after insulin treatment, and performed immunoprecipitation of protein lysates using an anti–phospho-tyrosine antibody followed by Western blotting for IR β and IRS-1. β 5 blockade led to persistent tyrosine phosphorylation of both IR β and IRS-1 in skeletal muscle (Fig. 3*D*). Taken together, these data indicate that targeted disruption of the MFGE8/ β 5 pathway potentiates insulin receptor activation and signaling in vitro and in vivo.

ανβ5 Integrin Regulates Hepatic Insulin Signaling. To determine whether our findings extended beyond the skeletal muscle system, we assessed the effect of β5 blockade in the hepatic system. We performed a pyruvate tolerance test (PTT) in 6 wk old male WT mice in the presence of β5 blocking or isotype control antibody. β5 blockade significantly suppressed the rise in blood glucose levels in WT mice after administration of pyruvate (Fig. 4*A*). We next treated HepG2 liver cells and primary hepatocytes with insulin for 30 min in the presence of β5 blocking or control antibody and performed coimmunoprecipitation using a phosphotyrosine antibody followed by Western blotting for IRβ and IRS-1. We found that β5 blockade potentiated insulin receptor signaling in response to insulin in both HepG2 (Fig. 4 *B–D*) and primary hepatocytes (Fig. 4*E*). We performed similar coimmunoprecipitation experiment from total liver lysates of mice treated with

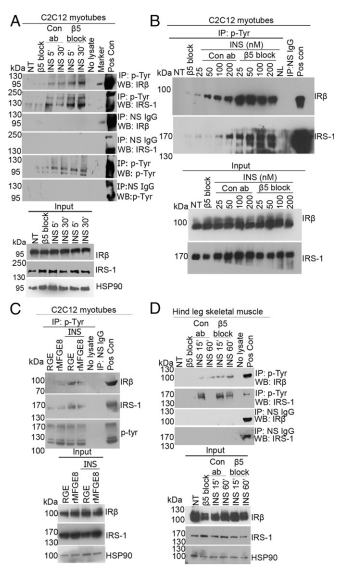


Fig. 3. Persistent activation of insulin receptor signaling with disruption of the MFGE8/_{β5} pathway in skeletal muscle. (A-C) Coimmunoprecipitation experiments demonstrating the effect of $\beta 5$ blockade and rMFGE8 treatment on IRB and IRS-1 tyrosine phosphorylation in presence or absence of insulin (INS). (A) C2C12 myotubes were treated with insulin (100 nM) for 5 min (INS 5') or 30 min (INS 30') in presence of either β5 (A) blocking or isotype control antibody or (C) rMFGE8 or RGE protein coadministered with insulin. Western blots are representative of three independent experiments. (B) Dose response of insulin action on IR β and IRS-1 tyrosine phosphorylation in presence of $\beta 5$ blocking or isotype control antibody. Western blots are representative of four independent experiments. (D) Coimmunoprecipitation studies showing tyrosine phosphorylation of IR β and IRS-1 in skeletalmuscle lysates from WT mice treated with IP insulin (1 U/kg) for 15 min (INS 15') or 60 min (INS 60') in presence of β 5 blocking or control antibody. Western blots are representative of three independent experiments. WT male mice were used for all in vivo experiments.

insulin for 15 and 60 min in the presence of β 5 blocking or control antibody. In this system, β 5 blockade significantly enhanced phosphorylation of IR β and IRS-1 at the 15 min time point but not at the 60 min time point (Fig. 4 *F*–*H*). We also evaluated whether prolonged treatment with β 5 blocking antibody (weekly administration for a total of 8 wk) impacted liver fat content and found no differences in liver triglyceride content or serum triglycerides (*SI Appendix*, Fig. S6 *A* and *B*). Taken together, these data indicate that

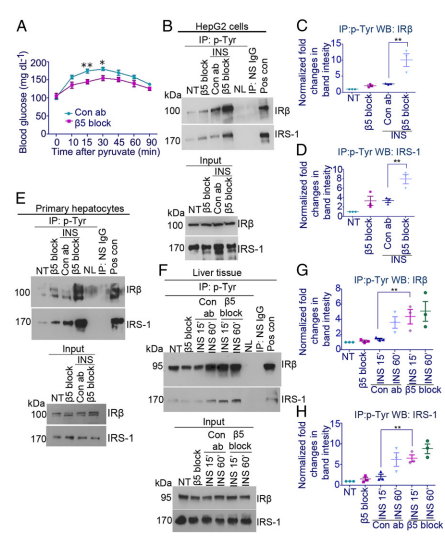


Fig. 4. Enhanced hepatic insulin sensitivity after β 5 blockade. (*A*) PTT in 6 wk old male WT mice after IP injection of β 5 blocking (β 5 block) or isotype control antibody (Con ab). *n* = 3 to 4 in each group per experiment; data merged from two independent experiments are presented. (*B*) Coimmunoprecipitation experiments demonstrating the effect of β 5 blockade on IR β and IRS-1 tyrosine phosphorylation in the presence or absence of insulin (INS, 100 nM for 30 min) in HepG2 cells after treatment with β 5 blocking or isotype control antibody. Western blots are representative of three independent experiments. (*C* and *D*) Densitometric analysis of the Western blots (including *B*) of IR β and IRS-1 tyrosine phosphorylation in HepG2 cells. (*E*) Coimmunoprecipitation experiments demonstrating the effect of β 5 blockade on IR β and IRS-1 tyrosine phosphorylation in HepG2 cells. (*E*) Coimmunoprecipitation experiments demonstrating the effect of β 5 blockade on IR β and IRS-1 tyrosine phosphorylation in HepG2 cells. (*E*) Coimmunoprecipitation experiments demonstrating the effect of β 5 blockade on IR β and IRS-1 tyrosine phosphorylation in HepG2 cells. (*E*) Coimmunoprecipitation experiments demonstrating the effect of β 5 blockade on IR β and IRS-1 tyrosine phosphorylation in the presence of insulin (INS, 100 nM for 30 min) in murine primary hepatocytes in the presence of β 5 blocking or isotype control antibody. Western blots are representative of three independent experiments. Both male and female mice were used for primary hepatocyte isolation. (*F*) Coimmunoprecipitation studies showing IR β and IRS-1 tyrosine phosphorylation from liver lysates of WT male mice treated with IP insulin (1 U/kg) for 15 min (INS 10°) no 60 min (INS 60°) in the presence of β 5 blocking or control antibody. Western blots are representative of three independent experiments. (*G* and *H*) Densitometric analysis of Western blots (including *F*) showing fold changes in IR β and IRS-1 tyrosine phosphorylation

the effects of the MFGE8/ β 5 integrin pathway on insulin signaling extend beyond the skeletal system to the liver.

Association between the $\alpha\nu\beta$ 5 Integrin and IR β . We next evaluated whether the $\alpha\nu\beta5$ integrin is part of a complex with the insulin receptor. First, we immunostained for $\beta5$ and IR β in untreated and insulin-treated C2C12 myotubes. IR β and $\beta5$ colocalized, and colocalization increased with insulin treatment (*SI Appendix*, Fig. S7). Next, we performed a proximity ligation assay (PLA) in differentiated C2C12 and HsKM myotubes observing a positive signal for proximity between $\beta5$ and IR β that was markedly increased 30 min after insulin treatment (Fig. 5*A*). $\beta5$ blockade reduced the PLA signal at baseline and after insulin treatment (Fig. 5*A*). We also performed coimmunoprecipitation studies of C2C12 cellular lysates using an antibody targeting the cytoplasmic domain of $\beta5$ and probing the pulldown product for IR β . Consistent with our

PLA results, we observed a baseline association between β 5 and IR β , which was enhanced with insulin stimulation. Furthermore, the baseline and insulin-stimulated association was reduced with β 5 blockade (Fig. 5 *B* and *C*) and enhanced by rMFGE8 (Fig. 5 *D* and *E*). These findings indicate that $\alpha\nu\beta$ 5 integrin associates with IR β and that both insulin and MFGE8 strengthen this association.

To validate these findings in vivo, we administered β 5 blocking or isotype control antibody in WT mice 1 h before IP insulin treatment and then performed the same immunoprecipitation studies from hind-leg skeletal muscle lysates after insulin treatment. We observed a baseline association between β 5 and IR β that increased markedly 60 min after IP insulin treatment (Fig. 5 *F* and *G*). Pretreatment with the β 5 blocking antibody significantly reduced this association 60 min after insulin stimulation (Fig. 5 *F* and *G*). We also immunostained for IR β and β 5 in hind-leg skeletal muscles of mice treated with insulin for 60 min in the presence of β 5 blocking

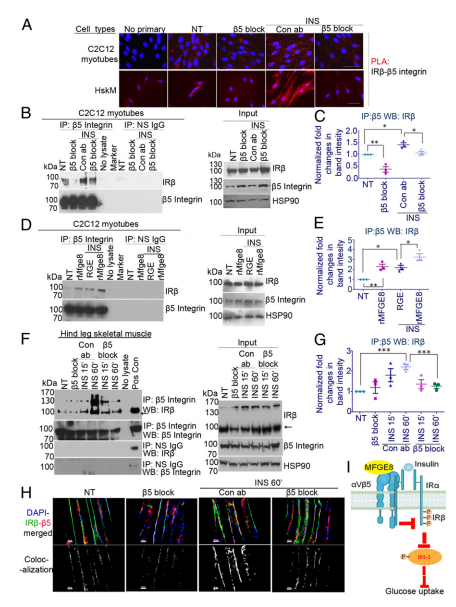


Fig. 5. Interaction between IR β and the β 5 integrin. (A) PLA showing proximity between IR β and β 5 in C2C12 myotubes and primary HskM treated with β 5 blocking or isotype control antibody and with or without 30 min of insulin (INS, 100 nM) treatment. n = 3 for C2C12, and n = 2 for HsKM myotubes. (Scale bar, 5 µm.) Magnification of 60× for C2C12 myotubes; 40× magnification for HsKM. (B-E) Coimmunoprecipitation experiments and respective densitometric analyses demonstrating the effect of β 5 blockade (B and C) and rMFGE8 treatment (D and E) on the interaction between IR β and β 5 in the presence or absence of insulin (INS, 100 nM) using an antibody recognizing the cytosolic domain of β 5 antibody for pulldown and subsequent Western blot for IR β . Western blots are representative of three independent experiments. Densitometric data represented as fold changes relative to no treatment group (NT). (F) Coimmunoprecipitation studies of skeletal-muscle lysates from WT mice treated with IP insulin (1 U/kg) for 15 min (INS 15') and 60 min (INS 60') in presence of β 5 blocking or control antibody. Western blots are representative of three independent experiments (three mice total per condition). (G) Densitometric analysis of Western blots (including F) showing fold changes in IR $\beta-\beta$ 5 interaction relative to no treatment (NT). (H) Immunostaining showing colocalization of IR β and β 5 integrin in the hind-leg skeletal muscles of mice 60 min after insulin treatment (INS 60') in presence of β 5 blocking or control antibody. For all in vivo experiments, 7 to 8 wk old WT male mice were used. Data analyzed by ANOVA followed by Bonferroni's posttest. *P < 0.05, **P < 0.01, and ***P < 0.001. (I) Proposed model showing how enhanced interaction between IR β and β 5 impedes insulin signaling.

or isotype control antibody. Insulin treatment enhanced the colocalization of IR β and β 5 (Fig. 5*H*). β 5 blockade dampened the colocalization signal after insulin treatment (Fig. 5*H*), corroborating the data from coimmunoprecipitation experiments (Fig. 5*F* and *G*). Taken together, these data indicate that insulin and MFGE8 enhance the interaction between the $\alpha\nu\beta$ 5 integrin and IR β . (Fig. 5*I*).

Insulin Causes Cell-Surface Enrichment of MFGE8. We were next interested in investigating whether insulin acts as an upstream modulator of the MFGE8/ β 5 integrin pathway. We first examined whether insulin induces cellular secretion of MFGE8 in differentiated C2C12 myotubes. We found no differences in MFGE8 protein levels with insulin treatment with or without β 5 blockade (*SI Appendix*, Fig. S8 *A* and *B*). Furthermore, we did not observe any significant fluctuations in serum or skeletal-muscle MFGE8 levels in mice during the course of an insulin tolerance test (ITT) (*SI Appendix*, Fig. S8 *C* and *D*).

We then tested whether insulin promotes MFGE8 localization at the outer cell surface, where it can bind to and activate integrin receptors. We performed cell fractionation on skeletal muscles harvested from WT mice after insulin administration. Interestingly, we observed enriched levels of MFGE8 in the plasma membrane 30, 45, and 60 min after insulin stimulation and a corresponding decrease in the intracellular pool of MFGE8 (Fig. 6 *A* and *B*).). However, we found minimal differences in surface-membrane MFGE8 15 min after insulin treatment compared to the untreated group. Taken together, these data suggest why blocking the MFGE8/ β 5 pathway impacts later phases of the ITT without affecting the earlier time points. We confirmed the accuracy of our cellular fractionation technique by Western blotting individual fractions for markers of the endoplasmic reticulum (ER; BiP), Golgi (STX), cytoplasm (HSP90), and plasma membrane (CAV-1) (Fig. 6*C*).

To investigate the underlying mechanism of MFGE8 movement to the cell surface, we treated C2C12 myotubes with insulin for 30 min with or without administration of Golgicide A (GCA). GCA is a highly selective and reversible inhibitor of Cis-Golgi protein Golgi-specific brefeldin-resistance guanine nucleotide exchange factor-1 (GBF-1). GBF-1 coordinates the assembly and transport of vesicles through the ER-Golgi network (51). We then incubated cells with cell-impermeable biotin to selectively biotinylate cell-surface proteins, purified protein from lysates using a streptavidin column, and western blotted for MFGE8. Insulin increased the relative abundance of biotinylated and reduced the abundance of nonbiotinylated MFGE8 (Fig. 6 D-F). Moreover, pretreatment with GCA significantly reduced the cell-surface level of MFGE8 in the presence of insulin indicating that insulin stimulates MFGE8 translocation through the ER-Golgi pathway (Fig. 6 D-F). Successful inhibition of GBF-1 by GCA treatment was confirmed by reduced cell-surface level of ABCA1 transporter (51-53) (Fig. 6D).

Previous studies suggested that AMP-activated protein kinase (AMPK) regulates GBF-1 activity through phosphorylation of GBF-1 at its Thr-1333 residue (54). To understand whether insulin affects GBF-1-dependent transport of MFGE8 via regulation of AMPK, we pretreated cells with AMPK activator AICAR for 2 h and then treated cells with insulin for 30 min and performed cellsurface biotinylation assay described above. AICAR treatment dampened the insulin-mediated enrichment of cell-surface level of MFGE8 (Fig. 6 G-I), indicating that AMPK modulates GBF-1-dependent vesicular transport of MFGE8 from the ER-Cis-Golgi face to the cell membrane. AICAR-mediated activation of AMPK was confirmed by an AMPK activity assay (SI Appendix, Fig. S9). The AMPK activity assay also showed a reduction in AMPK activity in insulin-treated cells (SI Appendix, Fig. S9), suggesting that insulin-dependent reduction in AMPK activity activates GBF-1, which in turn, enhances the transport of MFGE8 to the cell surface. Taken together, these data indicate that insulin enriches MFGE8 at the plasma membrane of skeletal muscle cells, thus enhancing its ability to bind and activate the $\alpha\nu\beta5$ integrin (Fig. 6J).

Metabolic Regulation of MFGE8. We were next interested in understanding the physiological regulation of MFGE8 under different metabolic states. We therefore evaluated blood glucose, serum insulin, and serum MFGE8 levels in mice under fed, fasted (5 and 16 h), and refed states (1 h after 5 or 16 h of fasting). We found a direct correlation between serum glucose, insulin, and MFGE8

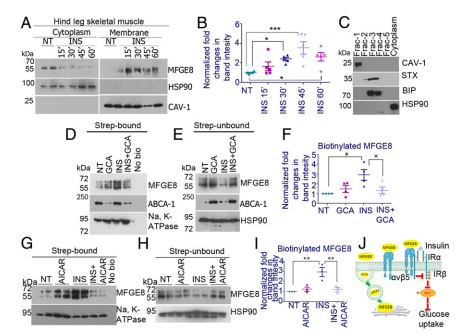


Fig. 6. Insulin induces cell-surface enrichment of MFGE8. (A-C) Cell fractionation of insulin-treated (1U/kg IP) skeletal-muscle tissue samples followed by Western blotting showing (A) MFGE8 expression in the cytoplasmic and cell-surface membrane fraction. Western blotting for HSP90 and CAVEOLIN-1 (CAV-1) confirmed cytoplasmic and membrane fractions, respectively. Western blots are representative of five independent experiments. (B) Densitometric analysis of Western blots (including A) showing relative fold changes in membrane MFGE8 in insulin-treated groups compared to NT. (C) Western blot showing surface membrane (CAV-1), ER (BIP), Golgi (STX) and cytoplasmic (HSP90) marker expression in different fractions from ultracentrifugation. Fraction 1 represents the cell-surface membrane used in A. WT male mice were used for all in vivo experiments. (D-F) Western blot showing cell-surface (streptavidin-bound) (D) and intracellular (streptavidin-unbound) (E) MFGE8 protein levels before and after 30 min of insulin (100 nM) treatment (INS) in C2C12 myotubes pretreated with either GCA (20 µM) or DMSO. Western blots are representative of four independent experiments. Western blotting for ABCA-1 confirmed successful inhibition of GBF-1 by GCA. (F) Densitometric analysis of Western blots (including D) showing fold changes in streptavidin-bound MFGE8 relative to NT. (G and H) Western blot showing cell-surface (streptavidin-bound, G) and intracellular (streptavidin-unbound, H) MFGE8 protein levels before and after 30 min of insulin (100 nM) treatment (INS) in C2C12 myotubes pretreated with either AICAR (10 mM) or DMSO. Data represents three independent experiments. (/) Densitometric analysis of Western blots (including G) showing fold changes in streptavidin-bound MFGE8 relative to NT. For all biotinylation experiments (D, E, G, and H), streptavidin-bound and unbound fractions were probed for HSP90 as a control for the intracellular (streptavidin-unbound) and Na,K-ATPase for the membrane-bound (streptavidin-bound) fractions. Total cell lysates not exposed to biotin reagent served a negative control (No bio). Densitometric data (B, F, and I) were analyzed by one-way ANOVA followed by Bonferroni's posttest. *P < 0.05, **P < 0.01, and ***P < 0.001. (J) Proposed model showing how cellsurface enrichment of MFGE8 impedes insulin receptor signaling via ligation of β 5 integrin.

levels (Fig. 7 A–C) with each decreasing in the fasting states and increasing in the fed states. To examine whether refeeding, which increased both serum insulin and MFGE8, impacted the interaction between β 5 and IR β in vivo, we performed coimmunoprecipitation studies of skeletal muscle tissue lysates (Fig. 7D). Refeeding markedly enhanced the interaction between β 5 and IR β as compared with the fasting state (Fig. 7D and E). These data indicate that MFGE8 and the interaction between β 5 and IR β are regulated by intermediary metabolic inputs linked to fasting and refeeding.

Discussion

Insulin resistance is an underlying causative factor that predicts an individual's progression toward T2D. T2D, in turn, is a major cardiovascular disease risk factor, and cardiovascular disease is the leading cause of death among those with T2D (55). Insulin remains the only uniformly effective treatment for type I diabetes and is also a critical component of the therapeutic arsenal for T2D, especially in the face of disease progression. However, chronic insulin treatment in the context of T2D, particularly in obese individuals, can be problematic. For one, insulin-resistant individuals often require exceptionally high doses of insulin, increasing the likelihood of serious consequences when dosing errors are made. Additionally, chronic insulin treatment is associated with weight gain, which can exacerbate the obesity that fuels an individual's insulin resistance in the first place. Finally, long-term treatment with high doses of insulin could theoretically produce untoward consequences through its mitogenic effects, especially in acknowledgment that obesity is a known risk factor for a variety of cancers. For all of these reasons, it would be useful to have more effective approaches to avoid the use of insulin in patients with T2D or at least the means to improve insulin sensitivity sufficiently to allow individuals to reduce their overall daily insulin dose.

Integrins are cell-surface receptors that mediate bidirectional signaling between the cell and the extracellular environment. The main body of literature examining the effect of integrins on insulin signaling has focused on $\beta 1$ integrin complexes. These data have identified important roles for these integrins in obesityassociated insulin resistance through mechanisms likely related to the physical consequences of fat expansion (26-30). By contrast, the data presented here demonstrate a direct effect of both MFGE8 and the $\alpha\nu\beta5$ integrin on insulin signaling that is independent of the effects of Mfge8 deletion on fat mass and that is apparent even in the absence of obesity (56). This phenotype is revealed by acute antibody-dependent $\beta 5$ or MFGE8 blockade in WT mice. Interestingly, the phenotype is not apparent in mice genetically lacking MFGE8 from birth (before the onset of differences in body composition at 10 wk of age), likely due to the induction of compensatory mechanisms.

Binding of insulin to the insulin receptor activates its intrinsic tyrosine kinase activity via receptor autophosphorylation and consequent phosphorylation of downstream signaling substrates. The delayed recovery of serum glucose levels we observed in the ITT after integrin blockade is strikingly similar to what is observed in ITT experiments in mice lacking the phosphatase PTP1B (10, 57). PTP1B is a well-established inhibitor of insulin receptor signaling that dephosphorylates IR^β and IRS-1. These data, coupled with the increase in the interaction between the $\beta 5$ and IR β after insulin stimulation, led us to consider whether $\beta 5$ dampens and/or terminates insulin receptor signaling. Our in vitro and in vivo data in skeletal muscle and the liver demonstrating persistent phosphorylation of IR β and IRS-1 with β 5 blockade are consistent with this hypothesis. Of note, the time course of these differences is somewhat different in each organ system, which is likely the result of the more complex role the liver plays in glucose homeostasis.

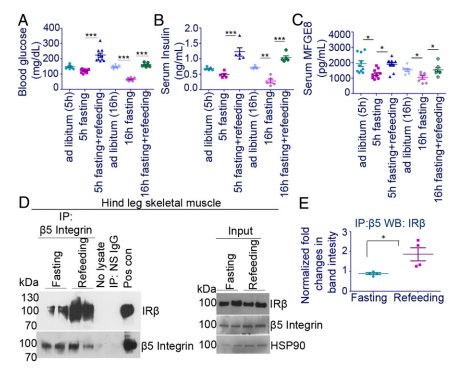


Fig. 7. Metabolic regulation of serum MFGE8 levels. (A-C) Blood glucose (A), serum insulin (B), and serum MFGE8 (C) levels in WT mice subjected to fasting for 5 and 16 h before a refeeding period of 1 h. n = 7 to 10 mice in A and C, n = 5 mice for B. Both male and female mice were used for these experiments. Data are from two to three independent experiments. (D) Coimmunoprecipitation experiments using hind-leg skeletal muscle lysates showing an interaction between IR β and β 5 in 16 h fasted mice and mice refed for 1 h after a 16 h fast. Data represents two independent experiments (four mice total per condition). WT male mice were used for this experiment. (E) Densitometric analysis of Western blots (including D) showing fold changes in IR β - β 5 interaction in the refed state relative to the fasted condition. Densitometry data were analyzed by Student's t test. Data in A-C were analyzed by one-way ANOVA followed by Bonferroni's posttest. *P < 0.05, **P < 0.01, and ***P < 0.001.

Whether these effects are through regulation of protein phosphatase or kinase activity remains to be determined. Alternatively, we can speculate the potential for a physical interaction between the integrin and the insulin receptor that, in the presence of ligand, induces structural changes in the insulin receptor that dampen its intrinsic tyrosine kinase activity.

An effect of insulin on MFGE8 protein production as a mechanism for our phenotype seems unlikely given the relatively rapid timeframe (30 min) within which we see MFGE8 impact insulin-stimulated glucose uptake. While we did not find an effect of insulin on MFGE8 secretion, insulin caused enrichment of cell membrane MFGE8 leading us to hypothesize that insulin modulates intracellular transport of MFGE8. GBF-1 coordinates the movement of proteins predominantly at the ER-cis-Golgi face (36, 37, 58, 59). Our data are consistent with work showing that GBF-1 facilitates anterograde trafficking of extracellular matrix proteins in mouse fibroblasts and human endothelial cells (60). Previous studies concerning how GBF-1 activity is regulated by AMPK have shown inconsistent results (54, 61). While AMPKmediated phosphorylation of GBF-1 at Thr-1333 deactivates GBF-1 leading to Golgi disassembly (54, 61), AMPK activation in human endothelial cells augments GBF-1 activity (60). In our study, AICAR-mediated AMPK activation phenocopies the effect of GBF-1 inhibition in insulin-treated myotubes suggesting that AMPK activation in the presence of insulin might have an inhibitory effect on GBF-1-dependent transport of MFGE8. As reported earlier, insulin treatment causes a reduction in AMPK activity (62-64) further supporting the hypothesis that insulin via suppression of AMPK activity triggers GBF-1-dependent transport of MFGE8 to the cell surface.

Our final sets of experiments were designed to examine whether MFGE8 is physiologically regulated by varying metabolic dietary states. Interestingly, serum MFGE8 levels closely track blood glucose and serum insulin levels in the fed, fasting, and refeeding state indicating metabolic regulation of serum MFGE8 that correlates tightly with fluctuations of endogenous insulin levels. Furthermore, the marked increase in β 5-insulin receptor interaction in skeletal-muscle lysates in the refeeding state suggests that metabolic regulation of MFGE8 directly impacts skeletal muscle integrin–insulin receptor interaction. One interesting question is what the cellular source of serum MFGE8 is, and to what extent the insulin-stimulated enrichment of cell membrane MFGE8 contributes to serum levels. This is a difficult question to answer experimentally since MFGE8 is produced and secreted by multiple cell types.

Normal glucose homeostasis is arguably one of the most essential vital functions of most living organisms. Decades of research have shown multiple, sometimes redundant, levels of regulation that control homeostatic glucose metabolism and provide tight regulation of this metabolism in multiple tissues under highly varying environmental and internal physiologic states. Our work adds to this understanding by demonstrating a previously unrecognized negative feedback loop of insulin signaling that is reliant on the MFGE8/65 pathway. This negative feedback of insulin signaling might be important at an organism-wide level, preventing dangerous excursions in glucose levels by excessive or prolonged insulin signaling. Alternatively, we might speculate this self-regulation at the receptor level of insulin signaling may be analogous to autoregulation that takes place in the excitatory receptors of neuronal synapses in which case the negative feedback is also particularly important for regional tissue physiology. In insulin signaling, negative feedback in one region of muscle may prevent deprivation of glucose from other nearby regions. Addressing such issues in the future will help us work toward a more complete understanding of all levels of the regulation of glucose metabolism in health and disease.

Whether direct targeting of MFGE8 or the β 5 integrin is a viable therapeutic option for patients with diabetes mellitus remains to be determined. MFGE8 impacts a heterogeneous array

of biological functions some of which, from a therapeutic viewpoint, are potentially favorable and others deleterious (65). For example, MFGE8 has multiple anti-inflammatory effects (46, 66–68), which could translate into the development of unwanted inflammation with prolonged therapeutic blockade. MFGE8 promotes wound healing, epithelial-cell proliferation, and tumor-cell progression (41, 43, 69–73). These effects could, in the setting of pathway blockade, lead to delayed wound repair in diabetics or alternatively inhibit malignant transformation and growth. The antifibrotic effects of MFGE8 in the setting of blockade could exacerbate fibrotic disease though these effects that seem to be independent of the β 5 integrin (74–76). The sum impact of blocking this pathway on cardiovascular health is difficult to define. MFGE8 deficiency accentuates cardiac hypertrophy and atherosclerosis in mice (39, 40), while higher coronary artery MFGE8 expression is associated with greater risk of developing coronary artery disease in humans (38). Our own work in metabolism indicates that blockade of the MFGE8/65 pathway leads to a reduction in fat absorption and postprandial lipemia (56, 77), two effects that would be especially beneficial for the cardiovascular health of obese, diabetic patients. Alternatively, targeting $\beta 5$ may be a more attractive therapeutic avenue. In fact, Cilengitide, a dual small-molecule inhibitor of $\alpha\nu\beta3$ and $\alpha\nu\beta5$, has already been used in clinical trials for glioblastoma (78). Though global $\beta 5^{-/-}$ mice develop normally (79), they are prone to age-dependent blindness (80). Whether this would be a limiting factor with biological or pharmacological blockade of $\beta 5$ in diabetes patients is unclear. Finally, our work provides a potential mechanism to explain the recent observations that correlate serum MFGE8 levels with multiple indices of insulin resistance (35, 36) and the increased risk of developing diabetes in a population, in which a missense variation of Mfge8 that increases plasma levels (37).

Materials and Methods

Mice. For all studies, 6 to 8 wk old age- and sex-matched mice in C57BL/6 background were used. *Mfge8^{-/-}* mice were purchased from RIKEN, are in the C57BL/6 background, and have been extensively characterized (66).

GTT, ITT, and PTT. The 7 to 8 wk old mice were fasted for 5 h and then injected IP with either 2 gm/kg glucose (GTT) or 1U/kg insulin (ITT). We treated mice IP with rMFGE8 or RGE protein (1 mg/kg), β 5 blocking antibody (5 mg/kg) (clone ALULA, provided by A. Atakilit, University of California San Francisco [UCSF]) or mouse isotype control antibody (clone 2H6-C2, ATCC CRL-1853), MFGE8 blocking antibody (2 mg/kg) (clone 2422; MBL life sciences, D161-3) or Armenian hamster isotype control antibody (clone eBio299Arm, ThermoFisher scientific, 14-4888-81) for 1 h before glucose or insulin administration. For PTT, 6 wk old male mice were fasted for 16 h followed by IP injection of pyruvate (2 gm/kg). Mice were treated with β 5 blocking or isotype control antibody 1 h before glucose or insulin or pyruvate injection and then again at regular time intervals for 2 h post glucose or insulin or pyruvate administration to measure blood glucose levels.

In Vivo 18-Fluoro-Deoxy-Glucose Deposition. Mice were imaged in a dedicated small animal PET/CT scanner (Inveon, Siemens Medical Solutions) under iso-flurane (2 to 2.5%) anesthesia 1 h after 18-Fluoro-Deoxy-Glucose (18FDG) and insulin injection in the presence of control or blocking antibody. All PET/CT data were normalized to an effective 18FDG dose for analysis and display within AMIDE (81, 82) (*SI Appendix, Supplementary Methods*).

In Vitro Glucose Uptake Assay. The 2NDBG uptake assay in C2C12 myotubes, HskM, and mouse primary skeletal-muscle myotubes was performed using a commercial assay kit (Cayman chemicals). Briefly, cells were preincubated with insulin (100 nM) for 10 min followed by the addition of nonhydrolyzable fluorescent glucose analog 2NBDG (10 mg/mL) for 20 min in the same media. We then measured the fluorescent intensity of cellular 2NBDG (excitation: 488 nm and emission: 535 nm) using a plate reader. Fluorescent intensities of 2NBDG were then normalized to wheat germ agglutinin (WGA 680 or WGA 594) staining intensities (*SI Appendix, Supplementary Methods*). Detailed methods of the treatment with function blocking antibodies, inhibitor, and recombinant integrin ligands are described in *SI Appendix, Supplementary Methods*.

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Treatment of Cells. C2C12 myotubes after 4 to 5 d of differentiation were washed in warm phosphate-buffered saline (PBS) and then incubated in serum-free Dulbecco's modified Eagle medium (DMEM) for 4 h before treating them with insulin (100 nM) for 15 or 30 min. For studying the dosedependent activation of the insulin receptor pathway, C2C12 myotubes were treated with different doses of insulin (25, 50, 100, and 200 nM) for 30 min. HepG2 and primary hepatocytes were treated with 100 nm insulin for 30 min. Treatment with $\beta 5$ blocking (5 $\mu g/mL)$ antibody or a control antibody was administered 1 h before insulin stimulation. rMFGE8 (10 µg/ mL) or RGE (10 µg/mL) control construct was added to the media concurrently with insulin. GCA (20 μ M) and AICAR (10 mM) were administered 2 h before insulin stimulation. An equivalent amount of dimethyl sulfoxide (DMSO) was added to control cells. Detailed methods of cell culture, protein isolation, Western blot, coimmunoprecipitation, PLA, and cell-surface biotinylation are described in SI Appendix, Supplementary Methods and Table S1.

Acute Treatment with β 5 Blocking Antibody In Vivo. We fasted mice for 4 h before treating them with β 5 blocking or control antibody for 1 h. Mice were then treated with IP insulin (1 U/kg) for 15 or 60 min before harvesting the hind-leg skeletal muscle and liver tissues. Tissues were then subjected to protein isolation, membrane fraction isolation, and immunostaining (*SI Appendix, Supplementary Methods*).

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Fasting and Refeeding of Mice. Mice were fasted for 5 or 16 h and then refed with normal chow diet for 1 h prior to obtaining samples (*SI Appendix, Supplementary Methods*).

Statistical Analysis. One-way ANOVA was used to compare data between multiple groups. When the ANOVA comparison was significant, further pair-wise analysis was performed using Bonferroni's posttest. For analysis of blood glucose and insulin levels over time during GTTs, ITTs, and PTTs, a two-way ANOVA for repeated measures followed by Bonferroni's posttest were used. Mann–Whitney U or a Student's t test was used to compare between two groups depending on whether the data passed normality tests. All statistical analysis was performed using GraphPad Prism 6.0. Data are represented as mean \pm SEM.

Study Approval. All experiments were approved by the Institutional Animal Care and Use Committee of UCSF and the UCSF Institutional Review Board.

Data Availability. All study data are included in the article and/or SI Appendix.

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