Protein Tyrosine Phosphatase 1B Regulates Pyruvate Kinase M2 Tyrosine Phosphorylation

Received for publication, December 3, 2012, and in revised form, April 24, 2013. Published, JBC Papers in Press, May 2, 2013, DOI 10.1074/jbc.M112.441469

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Background: Tyrosine phosphorylation of PKM2 inhibits its activity; however, the phosphatase(s) that regulates PKM2 phosphorylation remains unidentified.

Results: PKM2 is a novel PTP1B substrate and PKM2 Tyr-105 and Tyr-148 are key sites that mediate the interaction.

Conclusion: PTP1B regulates PKM2 tyrosine phosphorylation and activity in adipocytes.

Significance: These findings provide new insights into the regulation of adipose PKM2 activity.

Protein-tyrosine phosphatase 1B (PTP1B) is a physiological regulator of glucose homeostasis and adiposity and is a drug target for the treatment of obesity and diabetes. Here we identify pyruvate kinase M2 (PKM2) as a novel PTP1B substrate in adipocytes. PTP1B deficiency leads to increased PKM2 total tyrosine and Tyr-105 phosphorylation in cultured adipocytes and in vivo. Substrate trapping and mutagenesis studies identify PKM2 Tyr-105 and Tyr-148 as key sites that mediate PTP1B-PKM2 interaction. In addition, in vitro analyses illustrate a direct effect of Tyr-105 phosphorylation on PKM2 activity in adipocytes. Importantly, PTP1B pharmacological inhibition increased PKM2 Tyr-105 phosphorylation and decreased PKM2 activity. Moreover, PKM2 Tyr-105 phosphorylation is regulated nutritionally, decreasing in adipose tissue depots after high-fat feeding. Further, decreased PKM2 Tyr-105 phosphorylation correlates with the development of glucose intolerance and insulin resistance in rodents, non-human primates, and humans. Together, these findings identify PKM2 as a novel substrate of PTP1B and provide new insights into the regulation of adipose PKM2 activity.

The obesity epidemic and its associated comorbidities, including type 2 diabetes, cardiovascular disease, and many cancers, have increased interest in adipose tissue metabolism because this may present a novel therapeutic target (1). The adipose tissue integrates an array of homeostatic processes and is an important regulator of systemic insulin sensitivity and energy metabolism (2). There are two major types of adipose tissues in mammals: white and brown. White adipose tissue is the primary site for triglyceride storage or fatty acid release, whereas brown adipose tissue (BAT) is a thermogenic tissue with an established role in the defense against cold and emerging antiobesity properties (3, 4). Genetic and molecular studies identify tyrosine phosphorylation as a regulator of adipose tissue function, systemic glucose homeostasis, and energy balance (5–9).

Tyrosine phosphorylation is tightly controlled by the opposing actions of protein tyrosine kinases and protein tyrosine phosphatases (PTPs) (10). Protein tyrosine phosphatase 1B (PTP1B) is a ubiquitously expressed non-receptor tyrosine-specific phosphatase that is localized on the cytoplasmic face of the endoplasmic reticulum (11–13). PTP1B is a physiological regulator of glucose homeostasis and energy balance. Specifically, whole-body PTP1B KO mice are hypersensitive to insulin, lean, and resistant to high-fat diet (HFD)-induced obesity (14, 15). Mice with tissue-specific PTP1B deletion in the liver and muscle exhibit improved glucose homeostasis independent of body weight (16–18), whereas mice with neuronal deletion exhibit decreased body weight (19). However, the role of PTP1B in adipocytes is unresolved, with studies demonstrating detri-

*This work was supported, in whole or in part, by National Institutes of Health Grants R56DK084317 and R01DK090492 (to F. G. H.); AT-003645, HL-075675, HL-091333, and DK-087307 (to P. J. H.); CA-69202 (to Z. Y. Z.); and NCI and National Institutes of Health grants (to L. C. C.). This work was also supported by Juvenile Diabetes Research Foundation Research Grant 1-2009-337 (to F. G. H.), by a Training Program in Biomolecular Technology Fellowship T32-GM008799 (to J. B.), by an American Diabetes Association and multicampus award from the University of California, Office of the President (to P. J. H.), and by Damon Runyon Cancer Research Foundation Amgen Fellowship DRG-2056-10 (to C. A. L.). Dr. Cantley owns equity in and receives compensation from Agios Pharmaceuticals and serves on the Board of Directors and Scientific Advisory Board of Agios Pharmaceuticals. Agios Pharmaceuticals is identifying metabolic pathways of cancer growth and survival.

‡This article contains supplemental Figs. S1–S4.

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The abbreviations used are: BAT, brown adipose tissue; PTP, protein tyrosine phosphatase; PKM2, pyruvate kinase M2; HFD, high-fat diet; CII, compound II; D/A, PTP1B mutant D181A; IR, insulin receptor.
mental or beneficial effects of adipose PTP1B deficiency on body mass and insulin sensitivity (19, 20). Notably, PTP1B is a modulator of brown fat adipogenesis through a peroxisome proliferator-activated receptor γ-dependent mechanism, and PTP1B deficiency regulates PKR-like endoplasmic reticulum-regulated kinase phosphorylation and protein synthesis (21–23). Collectively, the salutary effects of PTP1B deficiency on obesity and diabetes have focused attention on it as a potential therapeutic target.

Pyruvate kinase (PK), a rate-limiting glycolytic enzyme, catalyzes the generation of pyruvate and ATP from phosphoenolpyruvate and ADP (24). In mammals, four pyruvate kinase isoforms exist: the L and R isoforms of the PKLR gene are expressed in liver and erythrocytes, respectively; the M1 splice isoform of the PKM gene is characteristic of more differentiated adult cells (e.g. muscle and brain); and the M2 isoform is expressed during embryonic development and in proliferative adult cell types (25). PKM2 expression facilitates aerobic glycolysis in tumors and provides a growth advantage (26, 27). Of note, tyrosine phosphorylation inhibits PKM2 activity (28). However, the phosphatase(s) that regulate(s) PKM2 remain(s) unidentified. In this study, we identify PKM2 as a novel PTP1B substrate and provide new insights into the regulation of adipose PKM2 activity.

**EXPERIMENTAL PROCEDURES**

**Reagents**—DMEM, G418, penicillin/streptomycin, puromycin, newborn calf serum, FBS, and trypsin were purchased from Invitrogen. PTP1B pharmacological inhibitor (compound II, CII) was prepared as described (29). Antibodies for human PTP1B (FG6), mouse PTP1B, and PKM1/2 were purchased from Abcam (Cambridge, MA). Tubulin antibodies were purchased from Upstate Biotechnology (Lake Placid, NY). PKM1 antibodies were purchased from Sigma. pPKM2 (Tyr-105), PKM2, PAKT (Ser-473), AKT, and FLAG antibodies were purchased from Cell Signaling Technology (Beverly, MA). pPR (Tyr-1162/Tyr-1163), and IR antibodies were purchased from Santa Cruz Biotechnology (Santa Cruz, CA). HRP-conjugated secondary antibodies were purchased from BioResources International (Carlsbad, CA). Unless indicated otherwise, chemicals were purchased from Sigma.

**Cell Culture**—Brown adipose cell lines were generated from whole-body PTP1B KO mice and reconstituted with human PTP1B WT and substrate-trapping mutant (D181A) as described previously (21). 3T3-L1 cells were maintained in DMEM containing 25 mM glucose, 10% newborn calf serum, 50 units/ml penicillin, and 50 μg/ml streptomycin. PKM silencing in 3T3-L1 cells was achieved by testing three different hairpins (Open Biosystems). Because the hairpins target regions that are shared for mouse PKM1 and PKM2, expression of both proteins was evaluated with one shRNA yielding a significant decrease in PKM2 without significant alterations in PKM1. Lentiviruses were generated by cotransfection of vectors in HEK293FT cells using Lipofectamine 2000 (Invitrogen) following the guidelines of the manufacturer and then used to infect 3T3-L1 cells. Cells were selected using puromycin (2 μg/ml), and drug-resistant pools were propagated. Knockdown cells were reconstituted by transient transfection of PKM2 WT or mutants. Stable cell lines were selected using G418 at a concentration of 400 μg/ml.

To induce cell differentiation, brown and 3T3-L1 preadipocytes were grown to confluence in culture medium containing 10% FBS or 10% newborn calf serum, respectively. Confluent cells were then switched to differentiation medium containing 20% FBS, 20 mM insulin, and 1 mM triiodothyronine (T3) for 48 h. Adipocyte differentiation was induced by incubating cells for 48 h in differentiation medium further supplemented with 5 μM dexamethasone, 0.5 mM isobutylmethylxanthine, and 0.125 mM indomethacin (induction medium). After induction, cells were cultured in differentiation medium until they exhibited a differentiated phenotype with accumulation of fat droplets. For confocal microscopy, cells were grown on coverslips and fixed with 100% ice-cold methanol for 10 min and then blocked with 3% BSA in PBS for 60 min. After incubation with primary antibodies and fluorescent-conjugated secondary antibodies, cells were mounted in Mowiol (Sigma) and examined using Olympus Fluoview laser-scanning confocal microscopy. Adipose tissue from different fat pads were fixed in 10% formalin and embedded in paraffin. Sections were stained using PKM2 and pPKM2 (Tyr-105) antibodies. Detection of protein specific bands was performed with appropriate fluorescein-conjugated secondary antibodies and visualized using a Leica DMi3000B inverted microscope.

**Biochemical Analyses**—Tissues/cells were homogenized in radioimmunoprecipitation assay buffer. Lysates were clarified by centrifugation at 13,000 rpm for 10 min, and protein concentrations were determined using a bichinonic acid assay kit (Pierce Chemical). Proteins (30–90 μg) were resolved by SDS-PAGE and transferred to PVDF membranes. For substrate-trapping experiments, cells were lysed in 1% Nonidet P-40 buffer with a protease inhibitor mixture (without sodium orthovanadate). Immune complexes were collected on PureProteome beads (Millipore) and washed with lysis buffer. Immunoblotting of lysates was performed with primary antibodies, and after incubation with secondary antibodies, proteins were visualized using enhanced chemiluminescence (Amersham Biosciences). Pixel intensities of immunoreactive bands were quantified using FluorChem Q imaging software (Alpha Innotech).

For site-directed mutagenesis, the indicated tyrosine (Y) to phenylalanine (F) mutations in mouse PKM2 cDNA were generated using a QuickChange Lightning site-directed mutagenesis kit (Agilent) following the instructions of the manufacturer and confirmed by DNA sequence analysis.

**PTP1B Enzymatic Activity**—3T3-L1 preadipocytes were treated with PTP1B inhibitor (compound II, 200 mM) (29) for 12 h and then lysed in Nonidet P-40 buffer (10 mM Tris-HCl (pH 7.4), 150 mM NaCl, 1 mM EDTA, 1% Nonidet P-40, 5% glycerol, and proteases inhibitors). 1 mg of total protein lysates was used to immunoprecipitate PTP1B using PureProteome beads (Millipore). The precipitate was washed twice with ice-cold lysis buffer and then resuspended in 50 μl of reaction buffer (50 mM HEPES (pH 5.0), 1 mM EDTA, 100 mM NaCl and 5 mM DTT) for 10 min, and the reaction was initiated by the addition of p-nitrophenyl phosphate (5 mM). PTP1B activity
was monitored by measuring the absorbance of p-nitrophenyl phosphate at 405 nm using a Wallac Victor 2 plate reader (PerkinElmer Life Sciences, Waltham, MA).

**Measurement of Total Pyruvate Kinase Activity**—Cells were lysed in 10 mM Tris–HCl (pH 7.5), 1.5 mM MgCl₂, 20 mM NaCl, 1 mM DTT, 1 mM PMSF, and a protease inhibitor mixture, incubated on ice for 20 min, and lysed with a glass Dounce homogenizer. Cell debris and unbroken cells were removed by centrifugation at 10,000 rpm for 15 min, and total pyruvate kinase activity was measured in the supernatant by a lactate dehydrogenase-coupled assay as described previously (30) with modifications. Briefly, proteins (1 μg) were added to 200 μl of reaction buffer containing 100 mM Tris–HCl (pH 8.0), 100 mM KCl, 10 mM MgCl₂, 0.5 mM EDTA, 0.2 mM β-NADH, 1.5 mM ADP, 5 mM phosphoenolpyruvate, and 0.6 units/ml lactate dehydrogenase. Pyruvate kinase activity was calculated at 25 °C by monitoring the absorbance at 340 nm using a Wallac Victor 2 plate reader. To determine the effects of PTP1B inhibition on pyruvate kinase activity cells were pretreated with CII for 12 h, and then pyruvate kinase activity was determined as described. CII was added to the lysis and reaction buffers.

**LC-MS/MS Mass Spectrometry**—Brown preadipocytes were cultured and differentiated as indicated. Adipocytes were homogenized in Nonidet P-40 buffer (10 mM Tris–HCl (pH 7.4), 150 mM NaCl, 1% IGEPAL™CA-630, and protease inhibitors). Lysates were clarified by centrifugation at 13,000 rpm for 10 min, and protein concentrations were determined using a bicinchoninic acid assay kit (Pierce Chemical). Proteins were resolved by SDS-PAGE, and gels were stained using Coomassie Brilliant Blue. Then, the 60- and 160-kDa bands were excised and processed by in-gel trypsin digestion to generate tryptic peptides. Tryptic peptides were resolved by reverse-phase chromatography (Cys-18) on an integrated nano-LC system (Easy-nLC, Proxeon Biosystems) and analyzed on a Linear Quadrupole ION Trap/Finnigan Polaris Q Mass Spectrometer (Thermo Scientific). Tandem mass spectra were extracted using ReadW.exe in centroid mode and searched with X!Tandem using the international protein index. Scaffold (Proteome Software) was used to validate MS/MS-based peptide and protein identifications as described previously (31).

**Mouse Studies—C57Bl/6** (The Jackson Laboratory) male mice were maintained on a 12-hour light-dark cycle and fed ad libitum. Mice were fed standard lab chow (Purina Lab Chow, catalog no. 5001) or switched to a HFD (60% kcal from fat, 15% fructose-sweetened beverages with their usual ad libitum diets). For the final 2-week intervention period, subjects consumed diets at the baseline energy level and macronutrient composition, except that 30% of the energy was from complex carbohydrates and 25% was provided by fructose- or glucose-sweetened beverages. The study was approved by the University of California Davis Institutional Review Board, and subjects provided informed consent to participate.

**Statistical Analyses**—Data are expressed as means ± S.E. Statistical analyses were performed using the JMP program (SAS Institute). Comparisons between groups were performed using unpaired two-tailed Student’s t test. *p ≤ 0.05; **p ≤ 0.01.

**RESULTS**

**Identification of Adipose PTP1B Substrates**—To gain insight into the molecular mechanisms underlying PTP1B action, we utilized substrate-trapping and mass spectroscopy approaches to identify novel adipose PTP1B substrates. PTP1B WT has a high catalytic constant (32), rendering its interaction with substrates transient and difficult to detect. However, the steady-state interaction of PTP1B with its substrates is enhanced in the substrate-trapping PTP1B mutant D181A (D/A) that retains substrate binding but is catalytically impaired (33). Thus, PTP1B D/A forms stable complexes with tyrosine-phosphorylated substrates. The substrate-trapping approach has been utilized successfully to identify PTP1B substrates (13, 22, 33–35). Brown preadipocyte cell lines were generated from PTP1B KO mice and reconstituted with PTP1B D/A as described previously (21). PTP1B was immunoprecipitated from starved or insulin-stimulated (10 and 30 min) PTP1B D/A adipocytes, resolved using SDS-PAGE, and then immunoblotted with anti-phosphotyrosine antibodies. Upon insulin stimulation, two prominent hyperphosphorylated bands at ~60 and ~160 kDa were detected corresponding to “trapped” putative PTP1B substrates (Fig. 1A, arrowheads). Lysis under stringent conditions (radioimmunoprecipitation assay buffer) significantly attenuated tyrosine phosphorylation, likely because of disruption of PTP1B D/A-substrates coassociation. Proteins were stained...
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PTP1B Regulates PKM2 Tyrosine Phosphorylation—Initially, we examined the expression of the M1 splice isoform (PKM1) of the PKM gene and M2 isoform (PKM2) in brown (21) and white (3T3-L1) adipose cell lines. Immunoblot analyses revealed the expression of both PKM1 and PKM2 in preadipocytes and adipocytes (differentiation days 4, 10, and 18) (Fig. 1B). Because mass spectroscopy identified peptides that are common to PKM1 and PKM2, we investigated the coassociation of each isoform with PTP1B. PTP1B was immunoprecipitated from starved or insulin-stimulated PTP1B WT- and PTP1B D/A-reconstituted adipocytes (21), resolved using SDS-PAGE, and then immunoblotted with PKM1- and PKM2-specific antibodies (Fig. 1C). Significant coassociation of PTP1B D/A with PKM1 and PKM2 was observed in adipocytes upon insulin stimulation. Given the reported regulation of PKM2 by tyrosine phosphorylation (28), and because PKM2 appears to be the major isoform in adipocytes (Fig. 3), we further investigated PKM2-PTP1B interaction. We reasoned if PKM2 is a PTP1B target, then PTP1B deficiency should lead to increased PKM2 tyrosine phosphorylation. To test this, PKM2 was immunoprecipitated from starved or insulin-stimulated KO and reconstituted (WT and D/A) adipocytes under basal and insulin-stimulated conditions (Fig. 1D). PKM2 total tyrosine phosphorylation was elevated significantly in KO and D/A compared with WT adipocytes under basal and insulin-stimulated conditions (Fig. 1D). PKM2 Tyr-105 phosphorylation was elevated significantly in KO and D/A compared with WT adipocytes (Fig. 1D). In addition, we determined PKM2 tyrosine phosphorylation in interscapular BAT of control (fl/fl) and adipose-specific PTP1B KO (AdPTP1B-KO) mice. AdPTP1B KO mice fed regular chow or an HFD exhibited increased PKM2 Tyr-105 phosphorylation compared with control mice (Fig. 1E). Together, these data demonstrate elevated both total and Tyr105 phosphorylation of PKM2 upon PTP1B deficiency in adipocytes and in vivo suggesting that PTP1B plays an important role in regulating PKM2 activity.

Mapping of PKM2-PTP1B Interaction—Elevated PKM2 tyrosine phosphorylation upon PTP1B deficiency prompted us...
to investigate whether PKM2 is a direct substrate of PTP1B. We examined the subcellular localization of PTP1B D/A and PKM2. GFP-tagged PTP1B D/A was transiently expressed in preadipocytes, and colocalization with endogenous PKM2 was monitored using confocal microscopy (Fig. 2A). Consistent with previous reports, PTP1B D/A localized to the endoplasmic reticulum network (11–13). Importantly, a subfraction of endogenous PKM2 colocalized with PTP1B. In addition, biochemical studies were performed to determine PTP1B-PKM2 coassociation. To that end, lentiviral shRNA was used to generate 3T3-L1 preadipocytes with stable knockdown of PKM (sh-PKM) as described under “Experimental Procedures.” Knockdown cells were reconstituted with non-targetable (shRNA-resistant) cDNA encoding mouse PKM2 WT, K433E mutant, and tyrosine (Y)-to-phenylalanine (F) mutants of individual PKM2 tyrosines (Fig. 2B). PTP1B D/A was transiently expressed in the generated cell lines and then immunoprecipitated, and coassociation with PKM2 was determined. Consistent with the proposed hypothesis, coassociation of PKM2 WT and PTP1B D/A was observed (Fig. 2C, first lane). Treatment with pervanadate, a strong inhibitor of tyrosine phosphatases that, as a phosphate mimetic, localizes at the catalytic pocket and oxidizes the essential cysteinyl residue in the catalytic center of the enzyme (36), disrupted PKM2-PTP1B D/A association (Fig. 2C, second lane). This suggested that the association is consistent with enzyme-substrate interaction that is mediated by the active site of PTP1B. Of note, PKM2 K433E (which attenuated total tyrosine phosphorylation) (28) exhibited minimal coassociation with PTP1B D/A (Fig. 2C). Strikingly, mutation of single tyrosine residues Tyr-105 or Tyr-148 nearly abolished PKM2-PTP1B D/A coassociation, whereas mutations of other tyrosine residues did not significantly alter coassociation (Fig. 2C and D). No association was observed between PKM2 and PTP1B WT, indicating that like many of its substrates, PTP1B WT does not form a stable complex with PKM2 (data not shown). Collectively, these data demonstrate that PKM2 is a
direct substrate of PTP1B and that the Y105F and Y148F mutants disrupt PKM2-PTP1B D/A association significantly.

Tyrosine Phosphorylation Regulates Adipose PKM2 Activity—To investigate the role of PKM2 Tyr-105 and Tyr-148 in regulating adipose PKM2 activity, we utilized 3T3-L1 preadipocytes expressing shRNA targeted to PKM (sh-PKM) or a scrambled (SCR) control and knockdown cells reconstituted with WT (M2R), Y105F, and Y148F mutants. B, the bar graph represents normalized data for PKM2 (Tyr-105)/PKM2 and presented as mean ± S.E. *, p ≤ 0.05 and **, p ≤ 0.01, significant difference between different cells versus control (sh-SCR). A.U., arbitrary unit. C, total pyruvate kinase activity in sh-PKM, sh-SCR, and knockdown preadipocytes reconstituted with PKM2 WT (M2R) or mutants (Y105F and Y148F). Cells were treated with PTP1B inhibitor (CII), and total pyruvate kinase activity was determined (dotted lines). D, the bar graph represents normalized data for pyruvate kinase activity (amount of β-NADH consumed using 1 µg of protein from total cell lysates/1 min of enzymatic reaction) and presented as mean ± S.E from two independent experiments. *, p ≤ 0.05 and **, p ≤ 0.01, significant difference between PTP1B inhibitor-treated versus non-treated for each cell type; #, p ≤ 0.05; ##, p ≤ 0.01, significant difference between control (sh-SCR) versus other cells for the same treatment.

FIGURE 3. Tyrosine phosphorylation of PKM2 regulates its activity. Immunoblot analyses of PKM2 (Tyr-105), PKM2, PKM1, PKM1/2, and tubulin in 3T3-L1 preadipocytes expressing shRNA targeted to PKM (sh-PKM) or a scrambled (SCR) control and knockdown cells reconstituted with WT (M2R), Y105F, and Y148F mutants. B, the bar graph represents normalized data for PKM2 (Tyr-105)/PKM2 and presented as mean ± S.E. *, p ≤ 0.05 and **, p ≤ 0.01, significant difference between different cells versus control (sh-SCR). A.U., arbitrary unit. C, total pyruvate kinase activity in sh-PKM, sh-SCR, and knockdown preadipocytes reconstituted with PKM2 WT (M2R) or mutants (Y105F and Y148F). Cells were treated with PTP1B inhibitor (CII), and total pyruvate kinase activity was determined (dotted lines). D, the bar graph represents normalized data for pyruvate kinase activity (amount of β-NADH consumed using 1 µg of protein from total cell lysates/1 min of enzymatic reaction) and presented as mean ± S.E from two independent experiments. *, p ≤ 0.05 and **, p ≤ 0.01, significant difference between PTP1B inhibitor-treated versus non-treated for each cell type; #, p ≤ 0.05; ##, p ≤ 0.01, significant difference between control (sh-SCR) versus other cells for the same treatment.
tiated adipocytes (supplemental Fig. S1A). To further establish regulation of PKM2 phosphorylation and activity by PTP1B, preadipocytes were treated with PTP1B inhibitor (CII) as described under “Experimental Procedures.” CII decreased PTP1B activity by 69.7 ± 5.4% and increased PKM2 Tyr-105 phosphorylation (supplemental Fig. S1, B and C). Notably, CII significantly decreased total pyruvate kinase activity in control (sh-SCR) and Y148F cells (Fig. 3, C and D). M2R cells treated with CII exhibited a trend for decreased pyruvate kinase activity but did not reach statistical significance, whereas knockdown and Y105F CII-treated cells exhibited comparable activity to non-treated cells (Fig. 3, C and D). Together, these data are consistent with previous findings that PKM2 tyrosine phosphorylation at Tyr-105 inhibits its pyruvate kinase activity.

Adipose PKM2 Tyr-105 Phosphorylation Is Regulated Nutritionally—To investigate whether adipose PKM2 phosphorylation is nutritionally regulated in vivo, we determined the effects of high-fat feeding on Tyr-105 phosphorylation in various adipose tissue depots. Mice were fed regular chow diet or a HFD (60% kcal from fat) starting at 7 weeks of age and then sacrificed after 3, 7, and 11 weeks. Immunoblot analyses revealed comparable PKM2 Tyr-105 phosphorylation in mice fed regular chow diet at 3 and 11 weeks in all examined adipose depots (Fig. 4). Notably, high-fat feeding for 3 weeks significantly attenuated PKM2 Tyr-105 phosphorylation in subcutaneous and epididymal depots (Fig. 4, B, C, and F). Prolonged high-fat feeding (7 and 11 weeks) led to significant attenuation of PKM2 Tyr-105 phosphorylation in all examined adipose depots, and this correlated with increased PTP1B expression (Fig. 4). In line with these biochemical studies, immunostaining of adipose tissue sections revealed comparable results with PKM2 Tyr-105 phosphorylation decreasing upon prolonged high-fat feeding (supplemental Fig. S2). Of note, HFD-induced attenuation of PKM2 Tyr-105 phosphorylation correlated with decreased insulin receptor (IR) (Tyr-1162/Tyr-1163) phosphorylation and decreased Akt (Ser-473) phosphorylation in adipose depots upon HFD feeding (supplemental Fig. S3). Decreased IR and Akt phosphorylation indicates decreased insulin sensitivity in HFD-fed mice compared with those fed chow. Together, these data demonstrate that adipose PKM2 Tyr-105 phosphorylation is regulated nutritionally and correlates with the development of insulin resistance in HFD-fed mice.

FIGURE 4. Regulation of adipose PKM2 Tyr-105 phosphorylation by high-fat feeding. Immunoblot analyses of PKM2 (Tyr-105), PKM2, PTP1B, and tubulin in lysates of brown (A), subcutaneous (S.Q.) (B), epididymal (Epi.) (C), mesenchymal (Mes.) (D), and retroperitoneal (Ret.) (E) adipose tissue depots of mice fed chow or an HFD for the indicated times. Each lane represents a sample from a separate animal. F, the bar graph represents normalized data for PKM2 (Tyr-105)/PKM2 presented as mean ± S.E. *, p ≤ 0.05 and **, p ≤ 0.01, significant difference between all groups versus mice fed a chow diet for 3 weeks; #, significant difference between mice fed an HFD for 7 and 11 weeks versus mice fed a HFD for 3 weeks.
We investigated whether adipose PKM2 Tyr-105 phosphorylation directly correlates with an altered metabolic state (independently of HFD-induced obesity) and whether this is conserved among species. To that end, we utilized different experimental models including rats, non-human primates, and human subjects who were subjected to different interventions that altered their metabolic state.

Firstly, we used UCD-T2DM diabetic rats, which exhibit adult onset polygenic obesity and insulin resistance and subsequently develop type 2 diabetes (37). Leptin administration to diabetic UCD-T2DM rats normalizes fasting plasma glucose concentrations and markedly improves insulin sensitivity independently of food intake (38). Diabetic UCD-T2DM rats received leptin (murine recombinant, 0.5 mg/kg) or saline injections (control). Importantly, saline-injected animals were pair-fed to the leptin-treated animals to control for food intake, and tissues were collected after one month of treatment. Immunoblot analyses revealed increased PKM2 Tyr-105 phosphorylation in white adipose tissue of the leptin-treated rats compared with controls (Fig. 5A). Consistent with the improved insulin sensitivity in leptin-treated rats (38), they exhibited increased IR and Akt phosphorylation compared with controls (supplemental Fig. S4A). Thus, adipose PKM2 Tyr-105 phosphorylation increases upon improved systemic insulin sensitivity independently of significant alterations in body weight.

Next, we investigated the regulation of PKM2 Tyr-105 phosphorylation in a non-human primate model of insulin resistance, fructose-fed rhesus macaques (39). Base-line adipose tissue lysates from rhesus monkeys that were monitored for 6 months (Control) and then divided into two groups: one fed fructose (Fructose) and the other fed fructose supplemented with fish oil (Fructose + F. Oil) for 6 months. The bar graph represents normalized data for PKM2 (Tyr-105)/PKM2, and data are presented as mean ± S.E., p = 0.01, significant difference between chow (Ctr.) versus fructose diets; ##, p = 0.01, significant difference between fructose (Fruct.) versus fructose with fish oil (Fruct + F. Oil) diets. C, immunoblots of PKM2 (Tyr-105) and PKM2 in lysates of subcutaneous adipose tissue biopsies from human subjects before and after consuming 25% of energy requirements as fructose-sweetened (upper panel) or glucose-sweetened (lower panel) beverage for 10 weeks. Subjects were separated into two groups: body mass index < 30 and body mass index ≥ 30. The bar graph represents normalized data for PKM2 (Y105)/PKM2 and presented as mean ± S.E., **, p = 0.01, significant difference between the same subjects before and after fructose-sweetened beverage consumption.
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months as detailed under “Experimental Procedures.” Animals fed fructose progressively developed glucose intolerance and decreased insulin sensitivity (39). Immunoblot analyses of subcutaneous adipose tissue revealed that PKM2 Tyr-105 phosphorylation was decreased significantly in the fructose-fed group compared with controls (Fig. 5B). On the other hand, the animals supplemented with fish oil exhibited PKM2 Tyr-105 phosphorylation that was comparable with controls. In addition, PKM2 Tyr-105 phosphorylation correlated with IR and Akt phosphorylation, which decreased in the fructose-fed group compared with controls and increased with fish oil supplementation (supplemental Fig. S4B).

Lastly, to determine whether findings from these animal models translate to humans, we analyzed PKM2 Tyr-105 phosphorylation in human subcutaneous adipose tissues from subjects that developed insulin resistance while consuming fructose-sweetened beverages. Subjects were provided 25% of energy requirements from glucose- or fructose-sweetened beverages for 10 weeks. Subjects consuming fructose, but not those consuming glucose beverages, exhibited increased fasting plasma glucose and decreased insulin sensitivity (40). Immunoblot analyses of subcutaneous adipose tissue in fructose-fed overweight (body mass index < 30) or obese (body mass index ≥ 30) subjects exhibited decreased PKM2 Tyr-105 phosphorylation after fructose feeding (Fig. 5C). In contrast, Tyr-105 phosphorylation did not change significantly in subcutaneous adipose tissue samples from subjects consuming glucose-sweetened beverages (Fig. 5C). Collectively, these results demonstrate that adipose PKM2 Tyr-105 phosphorylation directly correlates with the metabolic state in different species. Whether alterations in Tyr-105 phosphorylation are causative or consequences of the metabolic state remain to be determined.

DISCUSSION

The coordinated actions of protein tyrosine phosphatases and protein tyrosine kinases regulate phosphotyrosine-mediated signaling under physiological and pathophysiological conditions. Understanding the dynamic modulation of phosphotyrosine signaling in adipose tissue may provide avenues for therapeutic intervention in metabolic diseases. In this study, we identified PKM2 as a novel PTP1B substrate in adipocytes and adipose depots in vivo and determined PKM2 Tyr-105 and Tyr148 as key residues that mediated the interaction. In addition, we demonstrated regulation of PKM2 pyruvate kinase activity in adipocytes by tyrosine phosphorylation. Further, we uncovered direct correlation of adipose PKM2 Tyr-105 phosphorylation with the systemic metabolic state in rodents, nonhuman primates, and humans. Collectively, these findings provide new insights into the regulation of adipose PKM2 by tyrosine phosphorylation and its metabolic function.

Several lines of evidence established PKM2 as a novel PTP1B substrate. First, mass spectrometry of adipocytes expressing the PTP1B-trapping mutant identified PKM as a PTP1B-interacting protein. Second, adipocytes and adipose tissue lacking PTP1B exhibited increased PKM2 total and Tyr-105 phosphorylation. Third, substrate trapping demonstrated a direct association between PKM2 and PTP1B D/A. Importantly, per-
non-human primates, and humans. Additional studies are warranted to delineate the role of adipose PKM2 in metabolic regulation and to determine whether alterations in PKM2 Tyr-105 phosphorylation are causative or a consequence of the metabolic state in vivo.

The metabolic functions of adipose PTP1B are unresolved, with studies demonstrating detrimental, beneficial, or no effects of adipose PTP1B deficiency on body mass regulation and insulin sensitivity. Antisense oligonucleotides that lower PTP1B levels only in adipose tissue and liver reduce diet-induced obesity and enhance insulin signaling in obese (ob/ob) and diabetic (db/db) mice (48, 49). Similarly, administration of PTP1B antisense oligonucleotides to monkeys reduces PTP1B in adipose tissue and liver and improves insulin sensitivity (50). The observed effects could reflect a loss of PTP1B in liver and/or adipose tissue. On the other hand, adipose-specific PTP1B KO mice generated using the adipocyte protein 2 (aP2)-promoter to drive Cre recombinase expression exhibit increased body weight when fed an HFD (19). However, if the observed weight gain is caused only by adipose-specific PTP1B deficiency remains to be determined because the aP2 promoter can delete the target gene in other cell types (51, 52). Indeed, adipose-specific PTP1B KO mice generated using the adiponectin promoter exhibit comparable body weight to controls when fed an HFD but display larger adipocytes and increased basal lipogenesis (20). These studies identify PKM2 as one of several PTP1B substrates that can mediate PTP1B functions in adipose tissue. Presumably PTP1B can engage different substrate(s) in adipocytes, depending, at least in part, on the nature of the challenge/stimulus and its magnitude and duration. Intriguingly, adipose PTP1B function could vary, depending on the adipose depot. PTP1B inhibition in 3T3-L1 adipocytes inhibits adipogenesis (53). In contrast, PTP1B deficiency in brown adipocytes promotes differentiation and adipogenesis (21, 54).

The salutary effects of PTP1B deficiency and inhibition make it a therapeutic target for obesity, type 2 diabetes, and some types of cancer (55, 56). In addition, PKM2 plays important roles in cancer cells signaling and metabolism and is proposed as an attractive therapeutic target. PKM2 deficiency or inhibition elicits anticancer effects, such as impairment of cancer growth, induction of cell death, and increased sensitivity to chemotherapy (27, 57–59). On the other hand, compounds that activate PKM2 catalytic function may also have therapeutic implications to suppress tumorigenesis (60–62). The regulation of PKM2 activity in adipose tissue by tyrosine phosphorylation as presented here highlights the need to evaluate PKM2 functions in normal tissues before inhibitors and activators are deployed for therapeutic interventions. Dissecting the intracellular signaling events elicited by PTP1B and PKM2 will likely yield valuable insights and may lead to new avenues for therapeutic intervention.

Acknowledgments—We thank Dr. Benjamin Neel (Ontario Cancer Institute) for PTP1B floxed mice and Dr. Evan Rosen (BIDMC/Harvard University) for Adiponectin-Cre mice.

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SUPPLEMENTARY FIGURE LEGENDS

Figure S1: Regulation of pyruvate kinase activity by tyrosine phosphorylation. Total pyruvate kinase activity in differentiated 3T3-L1 adipocytes expressing shRNA targeted to PKM (sh-PKM) or a scrambled (SCR) control and knockdown cells reconstituted with WT (M2R), Y105F and Y148F mutants. Bar graph represents normalized data for pyruvate kinase activity and presented as means ± SEM from two independent experiments. (*) indicates significant difference between control (sh-SCR) versus other cells. B) 3T3-L1 preadipocytes treated were with PTP1B inhibitor (CII; 200 nM) for 12 h and PTP1B activity was determined. (*) indicates significant difference between treated and untreated cells. C) Immunoblots of PKM2 (Tyr105), PKM2 and Tubulin in lysates of sh-PKM and sh-SCR preadipocytes without (-) or with (+) PTP1B inhibitor treatment.

Figure S2: Immunostaining of PKM2 and Tyr\textsuperscript{105} in adipose depots. Brown (BAT), epidydimal (Epi.) and mesenchymal (Mes.) adipose sections of wild type male mice fed regular chow or HFD for 3 and 11 weeks were immunostained using PKM2 and Tyr105 antibodies.

Figure S3: Decreased IR and Akt phosphorylation in adipose depots of mice fed HFD. Immunoblots of pIR (Y1162/Y1163), IR, pAkt (S473), and Akt in lysates of (A) brown (BAT), (B) subcutaneous (S.Q.), (C) epidydimal (Epi.), (D) mesenchymal (Mes.), and (E) retroperitoneal (Ret.) adipose tissue depots of mice fed chow or HFD for the indicated times. Each lane represents a sample from a separate animal. Bar graphs represent normalized data for pIR (Y1162/Y1163)/IR (F) and pAkt (S473)/Akt (G) presented as means ± SEM. (*) indicates significant difference between all groups versus mice fed chow diet for 3 weeks, (#) indicates significant difference between mice fed HFD for 7 and 11 weeks versus mice fed HFD for 3 weeks.

Figure S4: IR and Akt phosphorylation in adipose tissue of rats and rhesus monkeys. A) Immunoblots of pIR (Y1162/Y1163), IR, pAkt (S473) and Akt in lysates of adipose tissues from leptin treated and non-treated UCD-T2DM diabetic rats. Bar graphs represent normalized data for pIR (Y1162/Y1163)/IR and pAkt (S473)/Akt, and presented as means ± SEM. (*) indicates significant difference between leptin treated and control rats. B) Immunoblots of pIR (Y1162/Y1163), IR, pAkt (S473) and Akt in lysates of adipose tissues from rhesus monkeys that were divided into two study groups; one fed fructose (Fructose) and the other fed fructose supplemented with fish oil (Fructose + F. Oil) for 6 months. Bar graphs represent normalized data for pIR (Y1162/Y1163)/IR and pAkt (S473)/Akt and presented as means ± SEM. (*) indicates significant difference between chow (Ctr.) and fructose diets. (#) indicates significant difference between fructose (Fruct.) and fructose with fish oil (Fruct + F. Oil) diets.
Figure S1
Figure S2
Figure S3
Figure S4