Mitotic regulators and the SHP2-MAPK pathway promote IR endocytosis and feedback regulation of insulin signaling

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Insulin controls glucose homeostasis and cell growth through bifurcated signaling pathways. Dysregulation of insulin signaling is linked to diabetes and cancer. The spindle checkpoint controls the fidelity of chromosome segregation during mitosis. Here, we show that insulin receptor substrate 1 and 2 (IRS1/2) cooperate with spindle checkpoint proteins to promote insulin receptor (IR) endocytosis through recruiting the clathrin adaptor complex AP2 to IR. A phosphorylation switch of IRS1/2 orchestrated by extracellular signal-regulated kinase 1 and 2 (ERK1/2) and Src homology phosphatase 2 (SHP2) ensures selective internalization of activated IR. SHP2 inhibition blocks this feedback regulation and growth-promoting IR signaling, prolongs insulin action on metabolism, and improves insulin sensitivity in mice. We propose that mitotic regulators and SHP2 promote feedback inhibition of IR, thereby limiting the duration of insulin signaling. Targeting this feedback inhibition can improve insulin sensitivity.

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The pancreatic hormone insulin controls glucose homeostasis and promotes cell growth and proliferation. Dysregulation of insulin signaling is linked to human metabolic syndromes and cancer\(^1\). Insulin binds to the insulin receptor (IR) on the plasma membrane (PM), and triggers phosphorylation-mediated activation of crucial enzymes that regulate glucose and lipid metabolism, and cell growth and division\(^2,3\). Activated IR phosphorylates itself and the IR substrate (IRS) proteins on tyrosines. Phosphorylated IRS proteins bind to multiple downstream effectors and adaptors, and activate two major branches of insulin signaling: the phosphatidylinositol 3-kinase (PI3K)-protein kinase B (AKT) and mitogen-activated protein kinase (MAPK) pathways. The PI3K-AKT pathway mainly governs metabolic homeostasis, whereas the MAPK pathway controls cell growth and proliferation. Src homology phosphatase 2 (SHP2, also known as PTPN11) binds to the C-terminal phosphotyrosine sites of IRS1/2 and promotes the activation of the MAPK pathway\(^4,5\). Mutations of IR cause severe inherited insulin resistance syndromes\(^6\), but the molecular mechanisms underlying insulin resistance in type 2 diabetes are complex and multifactorial\(^1\). One common theme is that insulin resistance in diabetic animals or patients causes ectopic accumulation of diacylglycerol and abnormal activation of novel Akt, which suppress insulin signaling at the level of IRS1 and possibly IR\(^1,7\).

The spindle checkpoint monitors kinetochore-microtubule attachment during mitosis and prevents chromosome missegregation\(^8\). In response to unattached kinetochores, the mitosis arrest deficiency 2 (MAD2) and budding uninhibited by benomyl 1-related 1 (BUB3) proteins, as subunits of the mitotic checkpoint complex (MCC), inhibit the anaphase-promoting complex/cyclosome bound to its mitotic activator, the cell division cycle 20 (CDC20) protein, to delay chromosome segregation\(^9-13\). When all kinetochores are properly attached by microtubules, the MAD2-binding protein p31\(^{comet}\) (also called MAD2L1BP) prevents the conformational activation of MAD2 and collaborates with the ATPase TRIP13 to disassemble MCC, thus promoting chromosome segregation\(^11,14-17\).

We have recently discovered a critical role of MAD2, BUBR1, and p31\(^{comet}\) in insulin signaling during interphase (Fig. 1a)\(^18,19\). MAD2 and BUBR1 are required for clathrin-mediated endocytosis of IR. MAD2 directly binds to the C-terminal MAD2-interacting motif (MIM) of IR, and recruits the clathrin adaptor AP2 to IR through BUBR1. p31\(^{comet}\) prevents spontaneous IR endocytosis through blocking the interaction of BUBR1-AP2 with IR-bound MAD2. Adult liver-specific p31\(^{comet}\) knockout

**Fig. 1** CDC20 is required for insulin-activated IR endocytosis. **a** Model of the regulation of IR endocytosis by spindle checkpoint proteins. MIM, MAD2-interacting motif. **b** Representative images of liver specimens from human non-diabetic and diabetic patients stained with DAPI (blue) and anti-IR (red) and anti-ZO-1 (green) antibodies. Scale bars, 40 µm. **c** Quantification of the percentage of cells with positive IR PM signals in liver specimens in **b** (non-diabetes, n = 51; diabetes, n = 19; mean ± s.d.; ****p < 0.0001; Mann-Whitney test, a non-parametric test). The liver specimens from insulin-treated patients were presented as orange triangles. **d** Western blot analysis of lysates of HepG2 cells stably expressing IR-GFP WT that had been transfected with the indicated siRNAs. **e** Cells in **d** were serum starved, treated without or with 100 nM insulin for 5 min, and stained with anti-GFP antibodies. **f** Quantification of the ratios of PM and IC IR-GFP signals of cells in **e** (siLuc, n = 39; siLuc + Ins, n = 14; siBUBR1, n = 18; siBUBR1 + Ins, n = 29; siCDC20, n = 17; siCDC20 + Ins, n = 34; siBUB1, n = 23; siBUB1 + Ins, n = 12; siBUB3, n = 15; siBUB3 + Ins, n = 10; siMAD1, n = 19; siMAD1 + Ins, n = 18; mean ± s.d.; ****p < 0.0001; two-tailed unpaired t test). Source data are provided as a Source Data file.
(KO) mice exhibit premature IR endocytosis in the liver and whole-body insulin resistance. Conversely, BUBR1 deficiency delays IR endocytosis and enhances insulin sensitivity in mice. These findings implicate dysregulation of IR endocytosis as a potential mechanism of insulin resistance.

In this study, we show that IRS1/2 cooperate with several spindle checkpoint proteins to promote insulin-stimulated IR endocytosis. In addition to MAD2 and BUBR1, CDC20 is also required for IR endocytosis. We further identify feedback regulation of IR endocytosis through a phosphorylation switch on IRS1/2 that is dependent on SHP2 and the MAPK pathway. Finally, we present evidence to suggest that targeting SHP2 might be a viable strategy to increase insulin sensitivity and treat diabetes.

Results

Reduced IR PM localization in diabetes patients. Previous biochemical studies have shown that insulin binding to liver PMs is reduced in obese and diabetic mice and humans20,21, suggesting that the level of IR at the PM might be reduced in people with diabetes. To confirm these findings, we examined the IR PM levels in liver samples from human patients using immunofluorescence staining. Because of the challenges of collecting liver sections from normal healthy individuals, we used surgical resection samples from patients with hepatocellular carcinoma that contained normal (non-malignant) and malignant tissues, and analyzed only normal hepatocytes. We performed immunofluorescence with anti-IR and anti-ZO-1 (as a PM marker) antibodies on 51 non-diabetic and 19 type 2 diabetes patient samples, and analyzed IR PM levels. The percentage of cells with IR PM signals in the liver sections from type 2 diabetes patients was significantly lower than that in non-diabetic patients (Fig. 1b, c). There was no correlation between insulin treatment and IR PM levels. These findings suggest that reduced IR PM levels might be a contributing factor to insulin sensitivity in human patients.

CDC20 is required for proper IR endocytosis. We have previously shown that MAD2 and BUBR1 promote IR endocytosis through recruiting AP2 (Fig. 1a), and p31^comet inhibits this process through antagonizing the MAD2–BUBR1 interaction18. We further tested whether other well-known mitotic regulators also have a role in IR endocytosis. Depletion of CDC20, BUB1, BUB3, or MAD1 did not alter the ratio of PM and intracellular (IC) IR signals in the absence of insulin (Fig. 1d–f). Depletion of CDC20 or BUB1, but not BUB1, BUB3, or MAD1, increased the PM/IC ratio of IR in the presence of insulin and delayed insulin-activated IR endocytosis. Normalization of PM/IC ratios of insulin-treated samples to those of untreated samples produced similar results (Supplementary Fig. 1a). Thus, like BUBR1, CDC20 is required for insulin-activated IR endocytosis.

CDC20 binds to both MAD2 and BUBR1 and nucleates the formation of MCC that consists of BUB1, BUB3, MAD2, and CDC2011,13,22,23. BUB3 is not absolutely required for the integrity and function MCC, but enhances the efficiency of MCC formation by targeting BUB1 to kinetochores. Structures of MCC nicely explain the binding cooperativity among MAD2, CDC20, and BUBR1, and highlight the importance of the BUBR1 KEN1 motif in MCC assembly22. We hypothesized that the assembly of an MCC-like complex on IR might strengthen the weak MAD2–BUBR1 interaction and help to recruit AP2 to IR.

To test this hypothesis, we first examined the region of BUB1 sensitive for IR endocytosis. The C-terminal domain of BUB1 binds to the N-terminal trunk domain of AP2B1 (Fig. 2a–c and Supplementary Fig. 1b), consistent with a previous report24.

Next, we generated human 293FT cell lines stably expressing IR tagged with green fluorescent protein (GFP) at its C terminus (Supplementary Fig. 1c) and examined the localization of IR-GFP after transfection with the indicated small interfering RNAs (siRNAs) or siRNA-resistant Myc-BUBR1 constructs. Expression of wild-type (WT) BUBR1, but none of BUBR1 truncation mutants, restored IR endocytosis in cells depleted of BUBR1 (Fig. 2d, e and Supplementary Fig. 1d, e). BUBR1 interacts with CDC20 through multiple binding motifs11,13,25. We constructed BUBR1 mutants targeting KEN1 (ΔKEN1) or KEN2 (ΔKEN2) boxes in the N-terminal region, and Phe (ΔPhe, also known as ABBA3) and D boxes (ΔD2) in the middle region (Fig. 2a). The ΔKEN2, ΔPhe, and ΔD2 mutants restored insulin-activated IR endocytosis in cells depleted of BUBR1 (Fig. 2d, e and Supplementary Fig. 1d, e). The ΔPhe + ΔD2 mutant was, however, slightly defective. Consistent with this cellular phenotype, the middle region of BUB1 containing Phe and D boxes showed weak binding to AP2B1, and mutation of both Phe and D boxes in full-length BUB1 (ΔPhe + ΔD2) decreased the interaction, suggesting that BUB1 binds to AP2B1 through multiple binding sites (Fig. 2b). Strikingly, BUB1 ΔKEN1 could not restore IR endocytosis in BUB1-depleted cells (Fig. 2d, e and Supplementary Fig. 1d, e). Given the critical role of BUB1 KEN1 in MCC assembly, these data suggest that IR uses an MCC-like complex to recruit AP2, except that in this context MAD2 binds to the MIM on IR, not the MIM on CDC20.

IRS1/2 promote IR endocytosis. Insulin binding to IR promotes clathrin-mediated endocytosis of the insulin-IR complex, which regulates the intensity and duration of signaling. Two sequence motifs in IR—the NPEY960 and di-leucine (L986L) motifs—have previously been implicated in AP2 binding and IR endocytosis6–9 (Fig. 3a). We have shown that the MIM (R1333ILTL) of IR binds to MAD2, which in turn recruits AP2 with the help of BUB1 and CDC2018.

To study the relative contributions of these IR motifs to endocytosis, we generated HepG2 cell lines stably expressing IR-GFP WT, the MIM mutant (4A), Y960F, or L986A/L987A (AA), and examined the subcellular localization of these IR-GFP proteins (Fig. 3b, c). Without insulin treatment, IR WT, 4A, and Y960F localized to the PM, but IR AA was enriched in IC compartment. A large fraction of IR AA co-localized with the late endosome marker Rab7, indicating that IR AA underwent unscheduled endocytosis and accumulated in late endosomes (Fig. 3d). The di-leucine motif is located in a β strand of the N-terminal lobe of the IR kinase domain (Fig. 3e)30. Mutation of this motif is expected to alter the structural integrity or activity of the IR kinase domain.

The levels of ectopically expressed IR-GFP proteins were higher than that of the endogenous IR (Supplementary Fig. 2a, b). Furthermore, partial depletion of endogenous IR did not change the subcellular localization of these IR mutants. Thus, the behaviors of these mutants were not dependent on stoichiometric heterodimerization with the endogenous IR. On the other hand, because the IR depletion was incomplete, it remains possible that the residual endogenous IR might modify and regulate the ectopically expressed IR in a catalytic manner.

IR Y960F, 4A, and Y960F/4A mutants were less efficiently internalized after insulin stimulation (Fig. 3f and Supplementary Fig. 2c). As Y960 is phosphorylated in the activated IR31, defective endocytosis of IR Y960F suggests that phosphorylation of Y960 (pY960) might be required for timely IR internalization. The phosphotyrosine-binding domain of IRS1/2 directly binds to phosphorylated NPEY960 motif in activated IR32–35. Co-depletion of IRS1/2 blocked IR endocytosis induced by insulin, whereas depletion of either had no effect (Fig. 4a, b and Supplementary
Expression of RNAi-resistant IRS1 restored IR endocytosis in cells depleted of both IRS1/2. Thus, IRS1/2 act redundantly to promote IR endocytosis likely through binding to the phospho-NPEY960 motif. These results can explain why the activated IR is preferentially internalized.

IRS1 interacts with the adaptor-related protein complex 1 Mu subunit (AP1M1) and AP2M1 through multiple YXXΦ (Φ, bulky hydrophobic residues) motifs. The IRS1–AP2M1 interaction negatively regulates endocytosis of insulin-like growth factor 1 receptor (IGF1R). We confirmed that in vitro-translated (IVT) Myc-IRS1 full-length and the YXXΦ-containing central region (residues 449–679) bound to GST-AP2M1 (Fig. 4c and Supplementary Fig. 3c–f). IRS2 is highly homologous to IRS1 and also has conserved YXXΦ motifs (Fig. 4c and Supplementary Fig. 3f)37. AP2M1 binds to YXXΦ motifs and promotes clathrin-mediated endocytosis38. Thus, IRS1/2 contribute to IR endocytosis through bridging an interaction between AP2 and activated IR.

We tested whether mutations of YXXΦ motifs in IRS1 and IRS2 disrupted AP2M1 binding. In vitro-translated IRS1 (residues 449–864) bound to GST-AP2M1 (Supplementary Fig. 4a). Single YA mutant weakened the IRS1–AP2M1
**Fig. 2** The MAD2–CDC20–BUBR1 module is required for insulin-activated IR endocytosis. a Domains and motifs of human BUBR1 and AP2B1. TPR, tetratricopeptide repeat domain; GLEBS, GLE2-binding sequence; KARD, kinetochore attachment regulatory domain. Fragments that can or cannot bind to AP2B1 or BUBR1 are presented as + or −, respectively. NT, not tested. The KEN1/2 boxes, destruction boxes (D1/2), and Phe boxes (also called ABBA boxes; 1/2/3) are shown as red, blue, and green bars, respectively. b Binding of BUBR1 full length (WT) and mutants to AP2B1. 293FT cells were transfected with plasmids encoding GFP–AP2B1 and Myc–BUBR1 proteins. Total cell lysate (TCL) and anti-Myc-BUBR1 IP were blotted with anti-Myc and anti-AP2B1 antibodies. The relative band intensities are shown below (mean ± s.d.; n = 3 independent experiments). Asterisk indicates a cross-reacting band, which was due to the recognition of ectopically expressed Myc–BUBR1 proteins by the AP2B1 antibody. c Binding of AP2B1 full length (FL) and truncation mutants to BUBR1. 293FT cells were transfected with plasmids encoding GFP–AP2B1 and Myc–BUBR1 proteins. TCL and anti-Myc–BUBR1 IP were blotted with anti-GFP and anti-Myc antibodies. d 293FT cells stably expressing IR–GFP WT were transfected with the indicated siRNAs and Myc–BUBR1, serum starved, treated without or with 100 nM insulin for 5 min, and stained with anti-GFP (IR; green), anti-Myc (BUBR1; red), and DAPI (blue). Scale, 5 μm. e Quantification of the ratios of PM and IC IR–GFP signals of cells in d (siLuc, n = 96; siLuc + Ins, n = 29; siBUBR1, n = 29; siBUBR1 + Ins, n = 29; siBUBR1 + WT, n = 26; siBUBR1 + WT + Ins, n = 76; siBUBR1 + 1–525, n = 23; siBUBR1 + 1–525 + Ins, n = 51; siBUBR1 + 526–1050, n = 30; siBUBR1 + 526–1050 + Ins, n = 54; siBUBR1 + 1–350, n = 30; siBUBR1 + 1–350 + Ins, n = 32; siBUBR1 + 351–700, n = 26; siBUBR1 + 351–700 + Ins, n = 31; siBUBR1 + 701–1050, n = 33; siBUBR1 + 701–1050 + Ins, n = 34; siBUBR1 + ΔKEN1, n = 19; siBUBR1 + ΔKEN1 + Ins, n = 33; siBUBR1 + ΔKEN2, n = 21; siBUBR1 + ΔKEN2 + Ins, n = 29; siBUBR1 + ΔΔ2, n = 19; siBUBR1 + ΔΔ2 + Ins, n = 27; siBUBR1 + ΔPhe, n = 16; siBUBR1 + ΔPhe + Ins, n = 25; siBUBR1 + ΔPhe + ΔΔ2, n = 21; siBUBR1 + ΔPhe + ΔΔ2 + Ins, n = 20; mean ± s.d.; ***p < 0.001 and ****p < 0.0001; two-tailed unpaired t test). Source data are provided as a Source Data file.

The SHP2-MAPK pathway promotes IR endocytosis through IRS1/2. The tyrosine residues in YXXΦ motifs on IRS1/2 can be phosphorylated by the activated IR and be dephosphorylated by the tyrosine phosphatase SHP2.2,39 Strikingly, the serine residues immediately following the YXXΦ motifs are well conserved (Fig. 4c and Supplementary Fig. 3f), and can be phosphorylated by ERK1/2.40–42, ERK1/2-dependent phosphorylation has been proposed to reduce IRS1 tyrosine phosphorylation through negative feedback40–42, but the mechanism and function of this feedback regulation are unknown. We hypothesized that the MAPK pathway and SHP2 might regulate the IRS1/2–AP2 interaction and IR endocytosis through modulating IRS1/2 phosphorylation patterns.

To test this hypothesis, we examined the effects of inhibiting SHP2 or the MAPK pathway on insulin-activated IR endocytosis (Fig. 5a, b and Supplementary Fig. 5). The IR inhibitor (BMS359244) expectedly blocked IR endocytosis. Strikingly, the MEK inhibitors (U0126 and PD0325901) and the SHP2 inhibitor (SHP909) also inhibited IR endocytosis. By contrast, the AKT inhibitor (AKTI, VIII) did not affect IR endocytosis, indicating a specific requirement for the MAPK pathway and SHP2. Inhibitors of MPS1 (Reversine) and PLK1 (BI2566) did not appreciably inhibit IR endocytosis, ruling out the involvement of these mitotic kinases in this process.

The tyrosine residues in YXXΦ motifs on IRS1/2 also bind PI3K.43,44 The failure of IRS1 3YA mutant to restore IR endocytosis in cells depleted of IRS1/2 could be an indirect consequence of reduced PI3K activity. To test this possibility, we checked the effect of IRS1 3YA on insulin signaling (Supplementary Fig. 4g). IRS1/2 depletion was sufficient to inhibit IR endocytosis, but was insufficient to inhibit insulin signaling at insulin concentrations used in our IR endocytosis assays. Expression of IRS1 WT or the 3YA mutant did not appreciably alter insulin signaling. Furthermore, the PI3K inhibitor (GDC0941) did not affect insulin-activated IR endocytosis (Fig. 5a, b and Supplementary Fig. 5). This result argues against reduced PI3K activity as the underlying reason for the observed defects of insulin-activated IR endocytosis in IRS1/2-depleted cells, and further confirms that the PI3K-AKT pathway is not involved in this process.

We next chemically synthesized the unphosphorylated and phosphorylated IRS1 peptides containing YMPMS and examined their binding to AP2M1 (Fig. 5c). The unphosphorylated IRS1 peptide (YMPMS) bound to AP2M1, but the mutant peptide with Y612 and M615 replaced by alanine (AMPAS) did not. Phosphorylation of the serine in the YMPMS motif (YMPMPS) enhanced AP2M1 binding. Isothermal titration calorimetry (ITC) measurements confirmed that the phospho-serine IRS1 peptide (pS-IRS1) indeed bound to AP2M1 with higher affinity (K_D = 280 nM), as compared to the unphosphorylated peptide (K_D = 2.03 μM) (Fig. 5d). Thus, phosphorylation of IRS1 at S616, a known MEK/ERK site, augments the IRS1 endocytosis through multiple YXXΦ motifs.

RNAi-resistant IRS1 3YA mutants could not restore IR endocytosis in 293FT or HepG2 cells depleted of IRS1/2 (Fig. 4d, e and Supplementary Fig. 4c–f). Failure of these mutants to functionally complement indicates that the IRS1/2–AP2M1 interaction is required for insulin-activated IR endocytosis. We could not detect IRS1 in late endosomes during IR endocytosis, suggesting that IRS1 might dissociate from the IR complex prior to the fusion of the endocytic vesicles with the late endosome. Endogenous IRS1 interacted with the AP2 complex in 293FT cells and in primary mouse hepatocytes stimulated with insulin, but not in untreated cells (Fig. 4f, g). Thus, IRS1/2 bind to AP2M1 through canonical YXXΦ motifs in vitro and in mammalian cells, and promote insulin-activated IR endocytosis.
Taken together, our results support the following mechanism for insulin-activated IR endocytosis (Fig. 5g). The activated IR phosphorylates the tyrosine residues in YXXΦS motifs and the C-terminal SHP2-docking sites of IRS1/2, and stimulates the PI3K-AKT and MAPK pathways. In a negative feedback mechanism, activated ERK1/2 phosphorylate the serines in YXXΦS motifs on IRS1/2 and assist SHP2 to dephosphorylate IRS1/2. The IRS1/2 YXXΦS motifs with the serine phosphorylated and tyrosine dephosphorylated bind to AP2 with optimal affinities, promoting clathrin-mediated endocytosis of IR.

**Structural basis of the phospho-regulation of IR endocytosis.**
We next determined the crystal structure of AP2M1 (residues 160–435) bound to the serine-phosphorylated YXXΦS motif from IRS1 (pS-IRS1) (Table 1). The overall structure of the AP2M1–pS-IRS1 complex was similar to those of previously determined structures of AP2M1 bound to YXXΦ motifs. AP2M1 contained two interlinked β-sandwich subdomains: subdomain 1 (β1–6 and 17–19) and subdomain 2 (β7–16) (Fig. 6a). The pS-IRS1 peptide binds at the edges of strands β18 and β17 in subdomain 1, and interacts with residues from strands β1, β17, and β18 (Fig. 6a, b and Supplementary Fig. 6a). In particular, Y612 and M615 make extensive hydrophobic interactions with AP2M1.

The RNAi-resistant IRS1 3YF mutant with tyrosines in the YXXΦS motifs replaced by phenylalanines could not fully restore IR endocytosis in 293FT cells depleted of IRS1/2 (Fig. 4d, e). The hydroxyl group of Y612 forms a hydrogen bond with D176 in β1.

Fig. 3 The NPEY motif and MIM of IR are required for endocytosis. **a** Schematic illustration of sequence motifs and mutants of IR. **b** HepG2 cells stably expressing IR-GFP WT, 4A, Y960F, or AA were serum starved and stained with anti-GFP antibodies. Scale bars, 5 μm. **c** Quantification of the ratios of plasma membrane (PM) and intracellular (IC) IR-GFP signals of cells in **b** (WT, n = 21; 4A, n = 20; Y960F, n = 23; AA, n = 32; mean ± s.d.; ****p < 0.0001; two-tailed unpaired t test). **d** HepG2 cells stably expressing IR-GFP AA were serum starved and stained with anti-GFP and anti-RAB7 antibodies. Scale bar, 5 μm. The boxed region was magnified and shown on the right. Scale bar in the boxed region, 2.5 μm. **e** Ribbon diagram of the crystal structure of the IR kinase domain, with L986 and L987 shown as sticks. **f** HepG2 cells stably expressing IR-GFP WT, 4A, Y960F, or Y960F/4A were serum starved, treated without or with 100 nM insulin for the indicated durations, and stained with anti-GFP antibodies. Quantification of the ratios of PM and IC IR-GFP signals of cells was shown (WT, n = 40; WT + Ins, 10 min, n = 17; WT + Ins, 20 min, n = 33; Y960F, n = 32; Y960F + Ins, 10 min, n = 21; Y960F + Ins, 20 min, n = 19; 4A, n = 16; 4A + Ins, 10 min, n = 15; 4A + Ins, 20 min, n = 21; Y960F + 4A, n = 20; Y960F + 4A + Ins, 10 min, n = 16; Y960F + 4A + Ins, 20 min, n = 27; mean ± s.d.; ****p < 0.0001; two-tailed unpaired t test). Source data are provided as a Source Data file.
providing an explanation for why phenylalanines cannot functionally substitute for tyrosines. Phosphorylation of Y612 is expected to introduce both static hindrance and unfavorable electrostatic interactions with D176, explaining why tyrosine phosphorylation of YXXΦS motifs disrupts the IRS1–AP2 interaction.

We did not observe well-defined electron density for pS616 in IRS1, despite its ability to enhance the IRS1–AP2 interaction. pS616 is located in the vicinity of a positively charged patch on AP2M1 formed by residues K405, H416, and K420 (Fig. 6c), suggesting that the phospho-serine might engage in favorable electrostatic interactions with this basic patch. Mutations of H416 and K420 did not, however, reduce IRS1 binding (Supplementary Fig. 6b–d). Mutation of K405 destabilized the AP2M1 protein and reduced its binding to both the phosphorylated (pS616) and unphosphorylated IRS1 peptides. Thus, consistent with the lack of electron density, pS616 does not make defined electrostatic interactions with specific acceptor residues, and interacts with the positively charged patch as one structural entity.

SHP2 promotes IR endocytosis in mice. Liver-specific SHP2 KO mice show increased insulin sensitivity45,46, suggesting that SHP2 attenuates certain aspects of insulin signaling in the liver. We examined whether SHP2 inhibition could improve insulin sensitivity. The allosteric SHP2 inhibitor, SHP099, stabilizes the inactive conformation of SHP2, thus inhibiting its phosphatase activity4. WT mice maintained on high-fat diet (HFD) for...
5 weeks were treated with SHP099 (60 mg per kg body weight) by daily oral gavage for 6 days, and then tested for glucose and insulin tolerance. Strikingly, SHP099 administration markedly increased glucose tolerance and insulin sensitivity in HFD-fed mice (Fig. 7a, b). SHP099 did not change the body weight of mice fed HFD (Fig. 7c). Insulin stimulation caused IR endocytosis and reduced the IR staining at the PM in mouse liver sections (Fig. 7d, e). SHP099 delayed insulin-activated IR endocytosis. Finally, SHP099 inhibited the insulin-stimulated IRS1–AP2 interaction in primary hepatocytes (Fig. 4g).

To further confirm the requirement of SHP2 in promoting IR endocytosis in vivo, we introduced adeno-associated viruses (AAV) encoding control (Ctrl) or SHP2 short-hairpin RNAs (shRNA) into mice fed HFD via tail-vein injection. The SHP2 protein level in the liver from mice treated with AAV-SHP2 shRNA was reduced by ~70% as compared to that in mice treated with Ctrl shRNA (Supplementary Fig. 7a). By contrast, the SHP2 protein level in white adipose tissue (WAT) and skeletal muscle was not effectively depleted by SHP2 shRNA. Thus, as expected, SHP2 shRNA delivered by AAV was most effective in the liver. Consistent with the phenotypes of SHP099 administration, AAV-SHP2 shRNA treatment markedly increased glucose tolerance and insulin sensitivity in mice fed with HFD (Supplementary Fig. 7b, c). AAV-SHP2 shRNA treatment did not change body weight (Supplementary Fig. 7d). Importantly, AAV-mediated SHP2 silencing inhibited insulin-activated IR endocytosis in the liver (Supplementary Fig. 7e, f). These results confirm a role of SHP2 in promoting IR endocytosis and in metabolic homeostasis in vivo.

In addition to being a major organ for insulin signaling, liver is a major site for insulin clearance and defects in this process can cause hyperinsulinemia. Hepatic insulin clearance is mainly mediated by IR endocytosis, as liver-specific IR KO mice and mice deleted of the carcinoembryonic antigen-related cell adhesion molecule 1 (CEACAM1), a key regulator of IR endocytosis, develop severe hyperinsulinemia. We thus examined the effects of SHP099 on insulin clearance. To estimate insulin clearance, we examined the levels of insulin and C-peptide, a cleavage product of proinsulin, and determined the ratio of the serum levels of C-peptide and insulin. In HFD-fed mice, SHP099 slightly increased the fasting insulin level, but this increase was not significant (Fig. 7i). The serum levels of C-peptide were similar to those in vehicle-treated groups in both conditions (Fig. 7g), suggesting that SHP099 did not alter insulin secretion. As a result, the C-peptide:insulin ratio in mice fed HFD after SHP099 administration showed a mild reduction as compared to the control group (Fig. 7h). Therefore, despite causing defective IR endocytosis in the liver, SHP2 inhibition led to a mild defect in insulin clearance, but not severe hyperinsulinemia as observed in the CEACAM1 KO mice.

As shown in Fig. 5b, the inhibitory effect of SHP099 on IR endocytosis was not as complete as that of the IR inhibitor at 20 min after insulin stimulation, suggesting that there might be SHP2-independent IR endocytosis mechanisms. We have confirmed a requirement for CEACAM1 in IR endocytosis in our cellular assays (Fig. 7i, j). It is possible that CEACAM1 is required for all mechanisms of IR endocytosis, whereas SHP2 only regulates the IRS1/2 branch (Fig. 7k). In addition, other protein tyrosine phosphatases can contribute to IRS1/2 dephosphorylation, thereby regulating IR endocytosis. Future studies are needed to define the relationships and relative contributions of the various regulators of IR endocytosis.
SHP099-treated mice exhibited only a slight increase, and the levels of activating phosphorylation of AKT were similar to those in vehicle-treated mice. SHP099 did not appreciably alter the insulin-stimulated activation of the PI3K-AKT or MAPK pathways in WAT (Supplementary Fig. 8c, d). These data suggest that SHP2 inhibition has distinct effects on insulin signaling and possibly IR endocytosis in different tissues. The underlying reasons for these variations among tissues are unknown at present. Because SHP099 only increases AKT signaling in the liver, but not in skeletal muscle or WAT, our findings suggest that the liver is a major site of action for the insulin-sensitizing effect of the SHP2 inhibitor in vivo. We cannot, however, rule out the contributions of other tissues to the observed phenotypes.
Fig. 5  The SHP2-MAPK pathway promotes insulin-activated IR endocytosis. a HepG2 cells expressing IR-GFP WT were starved, treated with the indicated inhibitors for 2 h, treated without or with 100 nM insulin for 20 min, and stained with anti-GFP (IR; green) and DAPI (blue). Scale bar, 5 μm. b Quantification of the ratios of PM and IC IR-GFP signals of cells in a (DMSO, n = 47; DMSO + Ins, n = 56; IR, n = 50; IR + Ins, n = 65; U0126, n = 96; U0126 + Ins, n = 111; PD0325901, n = 26; PD0325901 + Ins, n = 54; SHPO99, n = 43; SHPO99 + Ins, n = 49; GDC0941, n = 57; GCO941 + Ins, n = 48; AKTi, n = 65; AKTi + Ins, n = 96; BI2536, n = 26; BI2536 + Ins, n = 24; Reversine, n = 37; Reversine + Ins, n = 27; mean ± s.d.; **** p < 0.0001; two-tailed unpaired t test). c Binding of IRS1 peptides to AP2M1 (residues 160–435). Input and proteins bound to IRS1 peptide beads were analyzed by SDS-PAGE and stained with Coomassie (CBB). The relative band intensities are shown below (mean ± s.d.; n = 5 independent experiments). d Isothermal titration calorimetry (ITC) analysis of binding between IRS1 peptides and AP2M1 (residues 160–435), with CBB indicated. e The IRS1 peptides were incubated with active SHP2 for the indicated durations, spotted onto membranes, and detected with the anti-pY612-IRS1 antibody. f Quantification of the relative SHP2 activity in e (mean ± s.d.; n = 4 independent experiments; **** p < 0.0001; two-tailed unpaired t test). e Model of the regulation of insulin-activated IR endocytosis by a phosphorylation switch on IR512/2. Insulin-bound IR phosphorylates itself and IRS1/2, and activates the PI3K-AKT and MAPK pathways. SHP2 acts upstream of RAS-RAF and promotes the activation of MAPK pathway. p31comet binds to the IR-bound MAD2 and blocks IR-AP2 association to prevent premature IR endocytosis. In feedback regulation, activated ERK1/2 phosphorylate S616 and other sites on IRS1. SHP2 binds to the C-terminal phosphotyrosine site on IRS1 and dephosphorylates Y612 of the doubly phosphorylated IRS1 (pY612/pS616), thus promoting IRS1–AP2M1 association. p31comet is released from MAD2 by an unknown mechanism, allowing the assembly of an MCC-like complex on IR. MAD2- and IRS1/2-dependent AP2 recruitment and clustering trigger clathrin-mediated IR endocytosis. Source data are provided as a Source Data file.

Table 1  Data collection and refinement statistics

| AP2M1-pIRS1          |           |
|----------------------|-----------|
| **Data collection**  |           |
| Space group          | P64       |
| **Cell dimensions**  |           |
| α, β, γ (°)          | 125.33, 125.33, 74.82 |
| α, β, γ (°)          | 90.00, 90.00, 120.00 |
| Resolution (Å)       | 32.6–3.20 (3.26–3.20) |
| Rmerge or Rsym       | 16.1 (29.2) |
| I/Δφ                | 13.2 (1.4) |
| Completeness (%)     | 100 (100) |
| Redundancy (%)       | 12.4 (13.3) |
| **Refinement**       |           |
| Resolution (Å)       | 27.14–3.20 (3.31–3.20) |
| No. of reflections   | 11,157 (1113) |
| Rwork/Rfree          | 23.3 (29.1)/23.6 (35.9) |
| No. of atoms         |           |
| Protein              | 2050      |
| Water                | 34        |
| B-factors            |           |
| Protein              | 45.5      |
| Water                | 47.3      |
| R.m.s. deviations    |           |
| Bond lengths (Å)     | 0.01      |
| Bond angles (°)      | 1.30      |

Data are collected from one crystal

*Values in parentheses are for highest-resolution shell

Discussion
We have elucidated a feedback regulatory mechanism of IR endocytosis and identified SHP2 as a molecular target whose inhibition delays insulin-activated IR endocytosis (Fig. 8c). Targeting this feedback regulation of IR endocytosis prolongs the metabolic branch of insulin signaling and improves insulin sensitivity in mice.

The mechanism of IR endocytosis was extensively studied for decades. Our recent discovery that mitotic checkpoint regulators, including MAD2 and BUBR1, are required for IR endocytosis promoted us to r-examine the mechanism of IR endocytosis. In addition to the recently discovered MIM, two other sequence motifs in IR, the NPXY960 motif and the di-leucine motif (L986/L987), had previously been implicated in AP2 binding and IR endocytosis. In this study, we have shown that the di-leucine motif is indeed required for IR endocytosis. Instead of directly binding to AP2, the phosphorylated NPXY motif binds to IRS1/2, which in turn recruits AP2 through multiple YXXΦ motifs. We propose that IRS1/2 bound to pY999 of IR provide one binding site for AP2, and the MAD2–CDC20–BUBR1 complex bound to the MIM of IR provides another AP2-binding site. These two modules collaborate to recruit AP2 to IR, triggering IR endocytosis. Alternatively, as IRS1/2 contain multiple functional YXXΦ motifs, each of these motifs and MAD2–CDC20–BUBR1 may mediate the recruitment of one AP2 complex, leading to the clustering of multiple AP2 molecules on IR and efficient assembly of the clathrin coat.

Our findings presented herein suggest that an MCC-like mitotic regulator module is assembled onto IR to control its endocytosis in interphase. These results raise the intriguing possibility that IR might reciprocally regulate MCC assembly and spindle checkpoint signaling during mitosis. It will be interesting to examine the effect of the connection between IR and mitotic...
checkpoint regulators on aneuploidy and spindle checkpoint activities. The levels of CDC20 and BUBR1 vary throughout the cell cycle and during aging. BUBR1 insufficiency causes premature aging and enhances insulin sensitivity in mice. Future studies are needed to test if the alteration of the levels of mitotic regulators directly contributes to aging through the regulation of insulin signaling.

Endocytosis of a cell surface receptor normally occurs after the receptor has been activated and has transduced its signals downstream. The mechanism ensuring that IR endocytosis only occurs after downstream signaling is unknown. In this study, we have shown that the MAPK pathway is required for IR endocytosis. Activated ERK1/2 phosphorylate the YXXΦS motifs in IRS1/2 and promote dephosphorylation of the tyrosine in these motifs by SHP2 in vitro and in vivo. The YXXΦS motifs
containing phospho-serine and unphosphorylated tyrosine are optimal ligands for AP2. Because ERK1/2 activation is a critical downstream event of insulin signaling, the requirement for an active MAPK pathway in IR endocytosis constitutes important feedback regulation that ensures proper insulin signaling. Chemical inhibition of SHP2 is an effective way to disrupt this feedback regulation, as it not only directly prevents the removal of the negative tyrosine phosphorylation but also inhibits the MAPK pathway and blocks the installation of the positive serine phosphorylation.

Our study suggests that inhibition of SHP2 delays IR endocytosis, prolongs IR signaling through the PI3K-AKT pathway, and improves insulin sensitivity in the mouse. A recent study showed that the SHP2 inhibitor 11a attenuated the activation of PI3K-AKT pathway upon hepatocyte growth factor stimulation in melanoma cell lines\(^5\). Thus, SHP2 inhibition selectively enhances the PI3K-AKT pathway upon insulin stimulation, but not other stimuli. SHP2 inhibition prolongs phosphorylation of YXXΦ motifs on IRS1/2. Because these phosphotyrosine motifs also interact with PI3K, the enhanced tyrosine phosphorylation of these motifs caused by SHP2 inhibition may directly promote PI3K activation. This direct effect may contribute to the effectiveness of SHP2 inhibitors in countering insulin resistance.

Although we have shown the reduction of IR on the cell surface in liver samples of human type 2 diabetes patients, it can be a consequence of insulin resistance, rather than the cause. It is also possible that the reduced IR localization at the PM is due to the decrease in overall IR expression. Despite these caveats, our immunofluorescence results are consistent with earlier findings, and suggest that mechanisms regulating IR PM levels, including endocytosis, might be defective in people with diabetes. The circulating insulin level is elevated with insulin resistance and prevents hyperglycemia. There is also evidence that insulin hypersecretion can cause insulin resistance\(^5,54\). Genetic suppression of insulin secretion improves insulin sensitivity in aging mice and extends lifespan\(^9\). It will be interesting to test if the hyperinsulinemia-mediated insulin resistance is associated with the reduction of IR on the cell surface.

Our current study monitored IR endocytosis and insulin signaling at fixed time points, but not in real time. Moreover, although the depletion of endogenous IR did not change the cellular localization of IR mutants, we cannot rule out contributions from heterodimers formed between the endogenous IR and the ectopically expressed IR mutants to the localization and endocytosis of IR mutants. Targeting and mutating the endogenous IR loci are needed to examine the real-time kinetics of IR endocytosis and the effects of various perturbations in cells and in vivo.

Future studies are required to clarify the potential role of premature IR endocytosis in the pathogenesis of human insulin resistance syndromes. Mutations of IR are known to cause inherited severe insulin resistance syndromes\(^8\), but the mechanisms by which these mutations affect IR function have not been systematically explored. It will be interesting to test whether these mutations cause premature IR endocytosis and whether SHP2 inhibitors can recover the IR PM levels and insulin sensitivity.

Hyperactivation of signaling through receptor tyrosine kinases (RTKs) drives cancer progression. Chemical inhibitors of RTKs are widely used in the clinic to treat human cancers. A common signaling pathway downstream of RTKs is the RAS-RAF-MAPK pathway, which promotes cell proliferation and survival. SHP2 acts upstream of RAS-RAF to activate the MAPK pathway\(^5,56\). Thus, intense efforts have been devoted to develop chemical inhibitors of SHP2, which are expected to have therapeutic potential in cancer chemotherapy. SHP099, a specific allosteric inhibitor of SHP2, has been shown to have efficacy in targeting RTK-driven cancers in animal models\(^4\). Although phenotypes of SHP2 KO mice suggest potential adverse effects of SHP2 inhibition\(^57\), SHP099 administration into mice did not show significant toxic effects in that previous study\(^4\) and in the present study. Our study shows that short-term SHP2 inhibition improves systemic insulin sensitivity in mice. Thus, SHP2 inhibitors, such as SHP099, can be potentially repurposed to treat type 2 diabetes. Obesity increases the risks of both diabetes and certain types of cancers in humans\(^60\). The prevalence of all three conditions (obesity, diabetes, and cancer) has increased in recent years. SHP2 inhibitors may be particularly beneficial to patients who suffer from both diabetes and cancer.

**Methods**

*Mice.* Animal work described in this manuscript has been approved and conducted under the oversight of the University of Texas Southwestern Institutional Animal Care and Use Committee. All animals were maintained in a specific antigen-free barrier facility with 12 h light/dark cycles (6 a.m. on and 6 p.m. off). Mice were fed a standard rodent chow (2016 Teklad Global 16% protein rodent diet, Harlan Laboratories). For inducing insulin resistance, C57BL/6 (Stock No. 000664, Jackson laboratory) were fed a HFD (60%) (OpenSource Diets, Cat. No. D12492). Glucose and insulin tolerance tests, and metabolic analysis were performed as described previously\(^18\). For in vivo pharmacological assays, 6- to 8-week-old male mice were fed HFD for 5 weeks. Two days before drug administration, mice were switched to normal chow. SHP099 (MedChem Express) was dissolved in dimethyl sulfoxide (DMSO) and diluted into a 0.5% hypromellose and 0.1% Tween-80 solution. A unit of 60 mg kg\(^{-1}\) of SHP099 was administered by daily oral gavage for 6 days. For glucose tolerance test, mice were fasted for 14 h, and their blood glucose levels (\(T = 0\)) were measured with tail bleeding using a glucometer (AlphaTRAK). Then, 2 g of glucose per kg of body weight was injected intraperitoneally. Blood glucose levels were measured at the indicated time points after glucose injection. For insulin tolerance test, mice fasted for 4 h were injected intraperitoneally with recombinant human insulin (Eli Lilly) at 1 U per kg body weight, and their blood glucose levels were measured at the indicated time points after injection.

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**Fig. 7** SHP2 inhibition delays IR endocytosis and improves insulin sensitivity in mice. **a, b** Glucose tolerance test (a) and insulin tolerance test (b) of male mice fed HFD for 6 days. The mice were administered vehicle or SHP099 for 6 days. At 1 day after the last drug administration, experiments were performed. For GTT (a), vehicle, \(n = 12\); SHP099, \(n = 10\); for ITT (b), vehicle, \(n = 11\); SHP099, \(n = 10\); mean ± s.e.m; two-tailed unpaired t test. **c** Body weight of mice administered vehicle or SHP099 at 7 days post administration. Mean ± s.d. **d** HFD-fed mice were administered vehicle or SHP099 for 5 days. The mice were fasted overnight and again administered vehicle or SHP099. At 2 h after the last administration, the mice were injected with or without 1 U insulin via inferior vena cava. The livers were collected at the indicated time points and the sections were stained with anti-IR (red) and DAPI (blue). Scale bars, 5 μm. **e** Quantification of the ratios of plasma membrane (PM) and intracellular compartments (IC) IR signals of the livers in **d** (vehicle, 0 min, \(n = 43\); 1 min, \(n = 54\); 5 min, \(n = 27\); 15 min, \(n = 36\) and SHP099, 0 min, \(n = 35\); 1 min, \(n = 43\); 5 min, \(n = 30\); 10 min, \(n = 33\); 15 min, \(n = 33\); mean ± s.d.; ****p < 0.0001; two-tailed unpaired t test). **f–h** The levels of fasting serum insulin (f) and C-peptide (g), and the ratio of C-peptide:insulin (h) in mice fed HFD for 5 weeks. The mice were administered vehicle or SHP099 for 6 days. Vehicle, \(n = 11\); SHP099, \(n = 12\). I HepG2 cells stably expressing IR-GFP were transfected with CEACAM1 siRNAs, serum starved, treated without or with 100 nM insulin for 5 min, and stained with anti-GFP and DAPI. Quantification of the ratios of PM and IC IR-GFP signals of cells was shown (siLUC, \(n = 28\); siLUC + Ins, \(n = 49\); siCEACAM1 #1 + Ins, \(n = 27\); siCEACAM1 #2 + Ins, \(n = 25\); siCEACAM1 #3 + Ins, \(n = 35\); siCEACAM1 #4 + Ins, \(n = 18\); mean ± s.d.; ****p < 0.0001; two-tailed unpaired t test). **j** Western blot analysis of cell lysates in **i**. **k** Model of the regulation of insulin-activated IR endocytosis. Source data are provided as a Source Data file.
Reagents. Generation of rabbit polyclonal antibodies against GST, MAD1, CDC20, and BUB1 was described previously18,61. The following antibodies were purchased from commercial sources: anti-ZO-1/TJP1 (Immunohistochemistry (IHC), 1:100; #402200) and anti-ACTIN (Western blotting (WB), 1:1000; MA137018, Thermo Scientific); anti-IR-pY1150/1151 (WB, 1:1000; 19H7; labeled as pY1152/1153 IR for mouse IR in this study), anti-IRS1-pS616 (WB, 1:1000; C15H5; labeled as pS612 IRS1 for mouse IRS1 in this study), anti-AKT (WB, 1:1000; 40D4), anti-pT308 AKT (WB, 1:1000; D25E6), anti-ERK1/2 (WB, 1:1000; L34F12), anti-pERK1/2 (WB, 1:1000; 197G2), anti-SHP2 (WB, 1:1000; D50F2), and anti-RAB7 (Immunofluorescence (IF), 1:200; D95F2, Cell Signaling); anti-IRS1-pY612 (WB, 1:1000; labeled as pY608 IRS1 for mouse IRS1 in this study), anti-IR-pY972 (WB, 1:1000; labeled as pY962 IR for mouse IR in this study), and anti-IR (WB, 1:1000; IF, 1:100; CT-3, Millipore); anti-AP2B1 (WB, 1:1000; BD Biosciences); anti-IRS2 (WB, 1:1000; EPR904) and anti-MYC (WB, 1:1000; IF, 1:500; 9E10; Roche); anti-IRS1 (WB, 1:1000; A301-158A, Bethyl Laboratory); anti-CEACAM1 (WB, 1:1000; 283340, R&D Systems); anti-ACTIN (WB, 1:1000; C-4), anti-IR (WB, 1:1000; IF, 1:100; CT-3, Santa Cruz); and anti-mCherry (IF, 1:100; 1C51, Novus).

The siRNAs were synthesized by Dharmacon (Lafayette, CO) and had the following sequences: human BUBR1 (GGACACAUUUAGAUGCACUtt); human CDC20 (AGAACAGACUGAAAGUACUUU); human MAD1 (GAGCAGAUCCGUUCGAAGUUU); human BUB3 (GAGUGGCAGUUGAGUAUUU); human BUB1 (GAGUGAUCACGAAUUUC); human IRS1 (GAACCUGAUUGGUAUCUAC); human IRS2 (On-TARGETplus human IRS2 (8660) siRNA-SMARTpool); human CEACAM1 #1 (CCAUCAUGUAGCGUGUA); human CEACAM1 #2 (GAUGUAGUGUCAUGAGGCU); human CEACAM1 #3 (CGAUUUGUGUGAGGUCUU); human CEACAM1 #4 (CCAUAA

**Fig. 8** Mitotic regulators and SHP2 promote feedback inhibition of IR. a Insulin signaling in the liver from mice fed HFD for 5 weeks. The mice were administered vehicle or SHP099 for 5 days, fasted overnight, and administered vehicle or SHP099 once more. At 2 h after the last administration, the mice were injected with or without 1 U insulin via inferior vena cava. The livers were collected at the indicated time points. Lysates were prepared from these tissues and subjected to quantitative immunoblotting with the indicated antibodies. b Quantification of the blots in a. Mean ± range; n = 2 independent experiments. c Targeting feedback regulation of IR endocytosis for diabetes treatment. Left panel depicts the feedback regulation of IR endocytosis by ERK1/2 and SHP2 during unperturbed insulin signaling. Right panel illustrates the mechanism by which SHP2 inhibitor (SHP099) or shRNA blocks growth-promoting IR signaling and IR endocytosis, and prolongs insulin signaling through the PI3K-AKT pathway, which controls metabolism. Source data are provided as a Source Data file.
10% (v/v) fetal bovine serum (FBS), 2 mM glutamine, and 1% penicillin/streptomycin (Chemicals), the SHP2 inhibitor SHP099 (10 μM; Medchem express), the AKT inhibitor GSK690693 (1 μM; Selleck Chemicals), the PKC inhibitor BI2536 (200 nM, Selleck Chemicals), and the MPS1 inhibitor Inverstat (1 μM, Sigma).

Cell culture, transfection, and viral infection. 293FT cells were obtained from Invitrogen (R70007). HepG2 cells were obtained from American Type Culture Collection (ATCC; ATCC HB-8065). 293FT and HepG2 cells were cultured in high-glucose Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% (v/v) fetal bovine serum (FBS), 2 mM l-glutamine, and 1% penicillin/streptomycin. No mycoplasma contaminations were detected in these cells. Plasmid transfections into 293FT and HepG2 cells were performed with Lipofectamine® 3000 (Invitrogen). siRNA transfections were performed with Lipofectamine RNAiMAX (Invitrogen).

293FT or HepG2 cells expressing IR-GFP WT, or mutants were generated previously. A list of primers used in this study is provided in Supplementary Table 1. Briefly, cDNAs encoding IR mutants were cloned into the pIRES-GFP vector. The viral vectors were co-transfected with viral packaging vectors into 293FT cells, and the viral supernatants were collected at 2 and 3 days after transfection. The concentrated viruses were added to HepG2 cells stably expressing IR-GFP WT. The cells were transfected with the indicated siRNA at 1 day after viral infection. Analysis was performed at 4 days after infection. The cells were serum starved for 14 h before insulin treatment.

Primary hepatocytes were isolated from 2-month-old male mice with a standard two-step collagenase perfusion procedure. Cells were plated on collagen-coated dishes and incubated in attachment medium (Williams’ E Medium supplemented with 5% (v/v) FBS, 10 mM insulin, 10 mM dexamethasone, and 1% (v/v) penicillin/streptomycin). After 2–4 h, the medium was changed to low-glucose DMEM supplemented with 5% (v/v) FBS, 10 mM dexamethasone, 10 mM insulin, 100 nM triiodothyronine, and 1% (v/v) penicillin/streptomycin. After 1 day, the cells were seeded at clonal density and treated with DMSO or SHP099 for 2 h.

AAV encoding SHP2 shRNA (AAV-GFP-U6-mPTPN11-shRNA) or control shRNA (AAV-GFP-U6-scrambled shRNA) were generated at Vector Biolabs. We injected 10- to 11-week-old male mice fed with HFD for 5 weeks with the indicated constructs. The central section of 4 μm z-stack containing three contiguous focal planes was used for quantification. The cell edges were defined with Image J. The whole cell signal intensity (WC) and signal intensity (IC) were measured. The PM signal intensity (PM) was calculated by subtracting IC from WC. Identical exposure times and magnifications were used for all comparative analyses.

Protein purification. The full-length human AP2M1 was cloned into a pGEX-4T-1, and the plasmid was transformed into Escherichia coli strain BL21 (DE3). Protein expression was induced by 0.2 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) at 25°C for 12 h. The harvested pellets were lysed in the lysis buffer I [20 mM Tris–HCl, pH 8.0, 150 mM NaCl, 1% (v/v) Triton X-100, 5% (v/v) glycerol, 1 mM phenylmethylsulfonyl fluoride (PMSF), EDTA, and 1 mM PMSE]. The lysates were cleared by centrifugation at 4°C overnight. The protein was further purified by Superdex 200 2000 sieve exclusion column (GE Healthcare). The resulting protein-bound beads were washed extensively with lysis buffer I.

The AF2M1 fragment (residues 160–435) was cloned into a modified pET28a that introduced an N-terminal His tag followed by a thrombin cleavage site. The plasmid was transformed into BL21 (DE3). Protein expression was induced by 0.2 mM IPTG at 20°C overnight. The harvested pellets were lysed in the lysis buffer I [20 mM Tris–HCl, pH 7.5, 500 mM NaCl, 20 mM imidazole, and 1 mM PMSE]. After sonication, lysates were cleared by centrifugation at 4°C. The supernatants were loaded on a 0.45 μm filter and incubated with pre-equilibrated Glutathione Sepharose 4B beads (GE Healthcare). The resulting protein-bound beads were washed extensively with lysis buffer I.

Crystallization of the AP2M1-pIRS1 complex. The purified AP2M1 (residues 160–435) was mixed with the p5-IRS1 peptide (CHTDDGYPMPSPGVA, residues 607–623) at a molar ratio of 1:3 and then crystallized with the hanging-drop vapor diffusion method. The crystals of the AP2M1–p5-IRS1 complex grew within several days after the protein solution was mixed with the reservoir solution [1.0 M sodium malonate, pH 5.0, 0.1 M sodium acetate tri-hydrate, pH 4.5, and 2% (w/v) PEG 20K]. All crystals were cryoprotected with the reservoir solution including 15% (v/v) glycerol for data collection.

Data collection and structure determination. X-ray diffraction datasets were collected at the Advanced Photon Source beamline Sector 19-ID at a wavelength of 0.97914 Å and at 100 K. HKL3000 was used to process the datasets. The crystal of the
the AP2M1–p65RS1 complex disfavored a minimum Bragg spacing of 3.2 Å and exhibited the symmetry of space group P2₃ with cell dimensions of a = b = 123.3 Å, c = 74.4 Å. There are two molecules in the asymmetric unit, with a 53.4% solvent content.

The structure was determined by molecular replacement with Phaser-MR63, using the structure of the AP2M1-IGN38 complex (PDB ID: 1BXX) as the search model. Structure refinement was performed with COOT and PHENIX64–66. The final Rwork and Rfree were 23.6% and 23.6%, respectively. The Ramachandran statistics indicated that 91%, 8.4%, and 0.4% of the residues were in the favored, the allowed, and outliers regions, respectively. Data collection and refinement statistics are provided in Table 1. The model quality was validated with Molprobity67.

Protein-binding assays. For GST pull-down assays of IVT IRS1 or IRS2 proteins, beads bound to GST-AP2M1 or GST were incubated with IVT products diluted in the cell lysis buffer at 4°C for 2 h. After incubation and washing, proteins bound to beads were eluted with the SDS loading buffer, resolved with SDS-PAGE, and detected with Coomassie staining or immunoblotting with the appropriate antibodies.

For peptide pull-down assays, 2 mg peptides were coupled to beads with the SulfoLink Immobilization Kit (Thermo Scientific). Beads were resuspended in Buffer A [50 mM Tris, pH 7.5, 0.2 M NaCl, 0.1% NP-40, 10 mM sodium pyrophosphate, 0.5 mM DTT, and 2 mM PMSF] supplemented with protease inhibitors (Roche), PhosSTOP (Roche), 25 U ml⁻¹ turbo nuclease (Acelagen), and 100 µM L-tyrochalin B (Sigma), homogenized with Minilys (Bertin Technologies), and then incubated on ice for 1 h. After centrifugation at 20,817 × g for 30 min, the supernatant was subjected to immunoblot analysis as described in the Tissue histology and immunohistochemistry section. Images of sections were acquired as a series of 0.4 µm stacks with a DeltaVision system (Applied Precision, Issaquah, WA). Images were analyzed as described in the Immunofluorescence section.

Statistical analyses. Prism was used for the generation of all curves and graphs and for statistical analyses. Results are presented as mean ± s.e.m. or mean ± S.E. Two-tailed unpaired t-tests were used for pairwise significance analysis. Mann-Whitney test, a non-parametric test, was performed for human sample analysis. Sample sizes were determined on the basis of the maximum number of mice that could be bred in similar ages to maintain well-matched controls. Power calculations for sample sizes were not performed. We monitored weight and health conditions of mice, and excluded mice from experiments if the animal was unhealthy and the body weight was more than two standard deviations from the mean. Randomization and blinding methods were not used, and data were analyzed after the completion of all data collection in each experiment.

Data availability The coordinates of the AP2M1-phRS1 structure have been deposited to the Protein Data Bank (PDB ID: 6BNT). All other data are available from the authors upon request. The source data underlyings Figs. 1c, d, f, 2b, c, 3c, f, 4d, e, 5b, c, e, f, 7a, c–e, and 8a, and supplementary Figs 1a, b, 2b, c, 3a, b, d, e, 4a–c, e–g, 5e, d, 7a–d, f and 8a are provided as a Source Data file.

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Additional information

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