Inhibition of the metabolic regulator AMP-activated protein kinase (AMPK) is increasingly being investigated for its therapeutic potential in diseases where AMPK hyperactivity results in poor prognoses, as in established cancers and neurodegeneration. However, AMPK-inhibitory tool compounds are largely limited to compound C, which has a poor selectivity profile. Here we identify the pyrimidine derivative SBI-0206965 as a direct AMPK inhibitor. SBI-0206965 inhibits AMPK with 40-fold greater potency and markedly lower kinase promiscuity than compound C and inhibits cellular AMPK signaling. Biochemical characterization reveals that SBI-0206965 is a mixed-type inhibitor. A co-crystal structure of the AMPK kinase domain/SBI-0206965 complex shows that the drug occupies a pocket that partially overlaps the ATP active site in a type IIB inhibitor manner. SBI-0206965 has utility as a tool compound for investigating physiological roles for AMPK and provides fresh impetus to small-molecule AMPK inhibitor therapeutic development.

The ability to maintain energy homeostasis during acute or chronic periods of nutrient shortfall is an essential characteristic of all living organisms. A direct molecular link between nutrient supply and energy demand is provided by AMP-activated protein kinase (AMPK), a key regulator of cellular and whole-body metabolism. AMPK is an evolutionarily conserved serine/threonine kinase that senses, and is activated by, low adenylate charge (elevated AMP/ATP and ADP/ATP ratios) resulting from energetically demanding processes such as muscle contraction or reduced energy supply caused by hypoxia or nutrient deprivation. AMPK redirects cellular metabolism by down-regulating numerous ATP-consuming anabolic processes (e.g. protein, cholesterol, and fatty acid synthesis) and up-regulating similarly diverse ATP-producing catabolic processes (e.g. fat oxidation, glycolysis, and autophagy) to restore energy balance. It does this by direct phosphorylation, either modulating the activities of rate-limiting enzymes in multiple metabolic processes or regulating transcriptional activities of the factors governing their expression. For example, AMPK phosphorylation of cytosolic acetyl-CoA carboxylase 1 (ACC1) and mitochondrial-associated ACC2, inhibits de novo lipogenesis and promotes fat oxidation, respectively. AMPK signaling has also been associated with a range of non-metabolic regulatory roles (e.g. circadian rhythm, mitochondrial fission, and appetite control).

AMPK is an αβγ heterotrimeric complex, consisting of a catalytic α subunit (isoforms α1 and α2) and regulatory subunits β (isoforms β1 and β2) and γ (isoforms γ1, γ2, and γ3). The α subunit contains a canonical, bi-lobed kinase domain at the N terminus, followed by autoinhibitory (AID) and scaffolding domains, and a Ser/Thr-rich loop region (ST loop). The β subunit contains a mid-molecule carbohydrate-binding module (CBM) and C-terminal scaffolding domain, whereas the γ subunit consists of four cystathionine β-synthase domains and three adenine nucleotide binding sites (termed γ-sites 1, 3, and 4) that endow AMPK with its energy-sensing capabilities. AMPK signaling is dynamically and tightly controlled by numerous activating/inhibiting and localization mechanisms. Spatially, AMPK is targeted to various subcellular organelles...
and compartments in response to leptin and circadian rhythm (nucleus) (3, 4), glucose starvation (lysosomes, plasma membranes) (5, 6), mitochondrial damage (mitochondria), and increased autophagic flux (autophagosomes) (7, 8). Activation is triggered primarily by binding of AMP or ADP to exchangeable γ-sites 1 and 3, stimulating phosphorylation of the kinase activation loop residue Thr-172 (pThr-172) by upstream kinases LKB1 and Ca2+/calmodulin-dependent protein kinase 2 (CaMK2) and co-localizing AMPK with substrates on intracellular membranes (6, 9). pThr-172–independent activity has been demonstrated, although the contribution this makes to overall AMPK metabolic control is presently unclear (8, 10).

AMPK signaling is negatively regulated by multiple mechanisms, including exchange of AMP/ADP for ATP, AID-mediated autoinhibition, pThr-172–dephosphorylating actions of phosphatases, phosphorylation of suppressive regulatory sites in the α-ST loop, and a variety of nutrients (glucose, amino acids, and lipids), hormones (insulin, leptin, resistin), and cytokines (tumor necrosis factor α, ciliary neurotrophic factor, and interleukin-6) (11).

The metabolic dimensions associated with major human diseases, such as type 2 diabetes, cancer, and inflammatory disorders, have encouraged efforts to develop small-molecule AMPK activators. Patented examples now number in the hundreds. One of the first pharmacological AMPK activators discovered was 5-aminimidazole-4-carboxamide ribonucleoside (AICAR) (12). AICAR is taken up by cells via the adenosine transport system (13) and converted to the monophosphorylated derivative ZMP, which functions as an AMP-mimetic. Other AMPK agonists can be broadly classified as direct activators (those that bind to drug sites located either between the kinase domain and β-CBM (e.g. A-769662, salicylate, 991, and PF-937141) or within the γ subunit (e.g. C2 (18, 19)) or as indirect activators (those that commonly induce energy imbalance through mitochondrial toxicity, including metformin, xenobiotics, and other natural products).

Pharmacological AMPK inhibition provides potential strategies to treat obesity (appetite suppression), type 2 diabetes (enhanced insulin secretion), and stroke (neuroprotection) (11), and AMPK hyperactivity has also been linked to pathogenesis of neurodegeneration (20). The role of AMPK in cancer is complex; whereas initial studies demonstrated a tumor-suppressive role, AMPK signaling also contributes to the metabolic adaptations associated with tumor growth (e.g. increased glycolytic flux (the Warburg effect) and maintenance of ATP and NADPH) and promotes anchorage-independent proliferation (21–23). AMPK also promotes autophagic processes, via phosphorylation of ULK1, to maintain homeostasis in the neoplastic cell (24). Thus, AMPK is considered pro-tumorigenic under certain circumstances, underpinning the attraction of AMPK inhibition as a strategy for cancer treatments.

Current availability of small-molecule AMPK inhibitors, either for clinical application or as research tools to delineate AMPK’s physiological roles, is extremely limited. By far the most widely applied AMPK inhibitor, the pyrazolopyrimidine derivative compound C (dorsomorphin), was originally selected from a high-throughput screen and used to confirm AMPK-dependent effects of AICAR and metformin in cultured hepatocytes (25). Compound C is an ATP-competitive inhibitor and binds to the highly conserved active site of AMPK (26). However, in vitro screening has shown that compound C is promiscuous, inhibiting multiple kinases with similar or greater potency than AMPK (27). Numerous studies have since described off-target or AMPK-independent cellular effects, including inhibition of bone morphogenetic protein type I receptors ALK2, ALK3, and ALK6 (28), hypoxia-induced HIF-1 activation (29), preadipocyte proliferation (30), and macrophage chemotaxis (31). Compound C also blocks AICAR cellular uptake through competition for adenosine transporter binding sites, which largely accounts for its suppressive effects on AICAR-mediated AMPK activation (13). In light of these undesirable properties, conclusions drawn from using compound C as an AMPK inhibitor are now viewed with extreme caution, and recommendations are to avoid its application altogether (27). Other characterized direct antagonists include MT47-100, a low-potency, β2-AMPK allosteric inhibitor that intriguingly also activates β1-complexes (32), and SU6656, that paradoxically stimulates net cellular AMPK signaling by promoting phosphorylation of Thr-172 by LKB1 (33).

Here, we report the discovery of the small-molecule SBI-0206965 as a direct, type IIb AMPK inhibitor that demonstrated improved potency and kinase selectivity relative to compound C. A crystal structure of the AMPK α2 kinase domain/SBI-0206965 complex revealed a binding pocket that partially overlapped the ATP site; however, SBI-0206965 displayed mixed-competitive kinetics. SBI-0206965 inhibited AMPK signaling in a variety of cell types. Our study also highlights limitations pertaining to SBI-0206965 when used in conjunction with AICAR.

**Results**

**SBI-0206965: A potent, small-molecule AMPK inhibitor**

Using an active site competitive screen, Egan et al. (34) recently reported the 2-aminopyrimidine derivative SBI-0206965 (Fig. 1A) was a potent and highly selective inhibitor of the autophagy initiator ULK1. α1- and α2-AMPK complexes were also reported as hits in this screen. Subsequent studies in MEFs showed that SBI-0206965 (50 μM) produced a disconnect between AMPK-pThr-172 (elevated) and pACC (unchanged), indicative of direct inhibition of AMPK activity (34). Using cell-free phosphorylation of the AMPK and ULK1 synthetic peptide substrate S108tide as a measure of kinase activity (8), we found that SBI-0206965 (1 μM) was a more effective inhibitor of AMPK α1β1γ1 (80% inhibition) compared with ULK1 (63% inhibition) at 200 μM ATP (Fig. 1B). Conversely, compound C (Fig. 1A) was more effective as an inhibitor of ULK1 (36% inhibition) than AMPK (19% inhibition) at 200 μM ATP (Fig. 1B). Under our assay conditions, SBI-0206965 inhibited both α1β1γ1 and α2β1γ1 with nanomolar IC50 (Fig. 1C), whereas compound C inhibited both ULK1 and α1β1γ1 with micromolar IC50 (Fig. 1D). Using purified enzymes, we were unable to detect an effect of SBI-0206965 on the rate of LKB1-mediated phosphorylation of Thr-172 (Fig. 1E).

To assess kinase selectivity in terms of activity inhibition, we profiled SBI-0206965 (0.25 μM) against a diverse panel of 50
High-potency AMPK inhibitor

A

SBI-0206965

compound C

B

untreated
compound C (1 μM)
SBI-0206965 (1 μM)

kinase activity (% vs untreated)

α1β1γ1
ULK1

C

D

ULK1
α1β1γ1
α2β1γ1

kinase activity (% vs untreated)

log[SBI-0206965] (mM)

log[compound C] (mM)

SBI-0206965 IC_{50} (μM)

vs. ULK1
1.05 ± 0.07
vs. AMPK α1
0.40 ± 0.05
vs. AMPK α2
0.33 ± 0.07

compound C IC_{50} (μM)

vs. ULK1
3.09 ± 0.09
vs. AMPK α1
15.89 ± 0.10

E

basal
+ SBI-0206965

ptT172/α (arbitrary units)

LKB1 (min)
5
15

LKB1 (min)
SBI-0206965 (1 μM)
- 5 5 15 15
- - + - +

pT172
α

kD
75
50
S1). SBI-0206965 inhibited LKB1 and CaMKK2 activities by 40.001; ****, (compared with 31 kinases with compound C) (Fig. 2 and Fig. 3). SBI-0206965 displayed a preferable selectivity profile, inhibiting activity of only five other kinases by 0.06 to 0.59 μM with 100-fold elevation in [ATP] (from 20 to 2000 μM), whereas compound C displayed conventional ATP-competitive inhibitor kinetics, with IC_{50} increasing 8-fold from 1.91 ± 0.04 to 15.36 ± 0.06 μM across a 10-fold elevation in [ATP] (from 20 to 200 μM) (Fig. 3A). Whereas ATP (2000 μM) and compound C (20 μM) blocked binding of Thr-172–phosphorylated AMPK α2 kinase domain to ATP-agarose, SBI-0206965 (≤20 μM) was ineffective (Fig. 3B). However, SBI-0206965 (1 μM) inhibited the activity of phosphorylated α2 kinase domain to a similar extent as the AMPK heterotrimer at 200 μM ATP (Fig. 3C), confirming that the kinase domain alone encapsulates the SBI-0206965 binding site. We next tested the effect of varying ATP concentrations on the activity of phosphorylated α2 kinase domain across a range of [SBI-0206965]. SBI-0206965 displayed kinetic properties indicative of mixed-type inhibitor modality, as the K_s for ATP increased and V_max decreased with increasing [SBI-0206965] (Fig. 3D and Fig. S2, A and B). A secondary plot displaying the relationship between K_{i} / V_{max} (slopes on the Lineweaver–Burk plot (Fig. 3D)) and [SBI-0206965] generated a K_i of 261.4 nM (Fig. S2C). This is in contrast to the apparent αK_i (also commonly termed K_{i}') of 892.3 nM (Fig. S2D), which indicates that SBI-0206965 has a lower affinity for AMPK when ATP is bound. Equally, ATP has a lower affinity for AMPK when SBI-0206965 is bound (K_{i} (ATP) = 82.1 μM (Fig. 3D) versus αK_{i} (ATP) = 280.5 μM (derived from the equation, αK_i/K_j = αK_i/K_j)). In the context of AMPK regulation, these kinetic properties of SBI-0206965 are consistent with mixed-type inhibition.

To confirm the binding mode of SBI-0206965, we solved a 2.9 Å resolution crystal structure of the inhibitor complexed to AMPK (Table S1). A similar crystallization construct was used previously to visualize the compound C–binding pocket (26) (PDBe entry 3AQV) and adopted a structure essentially identical to that of the Thr-172–phosphorylated WT kinase domain in the heterotrimeric complex (16, 19). SBI-0206965 was found to occupy a pocket located between the kinase N- and C-lobes and hinge region, which partially overlaps the compound C–binding site (Fig. 4A and Fig. S3A). Specifically, the trimethoxyphenyl and pyrimidinyl rings of SBI-0206965 lie approximately within the same plane as, and overlap substantially with, the phenyl and pyrazolo[1,5-a]pyrimidinyl rings, respectively, of the compound C core. The positioning of the electronwith the negative bromine atom of SBI-0206965, probably contributing to increased potency relative to compound C, was confirmed by anomalous scattering at 13.6 keV (Fig. 4B) that revealed a large single peak for bromine (Table S1). The bromine moiety was found to occupy a large cavity bordered by residues Val-30, Lys-45, Ile-77, Met-93, Ala-156, and Asn-162. We could not unambiguously place the N-methylbenzamide group of SBI-0206965 within the electron density; therefore, we

Figure 1. Kinase selectivity profile for SBI-0206965 (0.25 μM) and compound C (2.5 μM). Profiling (n = 2 per kinase, with or without inhibitor) was performed using the MRC-PPU Express Screen service. MKK1 (activated 180% by SBI-0206965) is excluded from the left-hand panel for clarity.

SBI-0206965 is a mixed-type AMPK inhibitor.

We investigated the inhibitory mechanism of SBI-0206965 by performing dose-response assays at different ATP concentrations using α1β1γ1 AMPK. SBI-0206965 IC_{50} increased <4-fold from 0.16 ± 0.06 to 0.59 ± 0.09 μM with 100-fold
modeled two distinct conformations with equal occupancy (Fig. 4C). In one position (conformation A), the aromatic ring is rotated almost 80° with respect to the compound plane, and the N-methyl group extends into the space occupied by the catalytic loop, Mg2+/H11001-chelating residue Asn-144, and the ATP/β-phosphate in structures of active kinases (35). In the alternate position (conformation B), the benzamide ring is rotated 180° relative to conformation A, and the N-methyl group is directed toward P-loop residues Leu-22 and Gly-23 (Fig. 4C). In both conformations, SBI-0206965 makes two electrostatic contacts with the main chain of Val-96 in the kinase hinge region; otherwise, drug binding is mainly stabilized by hydrophobic contacts with hinge (Tyr-95), N-lobe (Leu-22, Ala-43, and the gatekeeper residue Met-93), and C-lobe (Ile-77, Gly-99, Glu-100, Leu-146, and Ala-156) residues.

Comparisons with either apo- (PDB entry 2YZA), compound C-complexed (26), or activated (PDB entry 4ZHJ) (19) kinase domain structures reveal the α2/SBI-0206965 complex contains many of the hallmarks of an unproductive kinase. In both compound C- and SBI-0206965-bound structures, the C-α-helix adopts a “swung-out” position, ensuring that the glutamate–lysine salt bridge, required for efficient phosphoryl transfer, is unformed (Fig. S3A). These residues (Lys-45 and Glu-64) are instead interdigitated by the activation loop residue Ser-161, hydrogen bonding with Glu-64. Glu-64 forms a further hydrogen bond with the backbone of Leu-160. Compared with the
compound C-bound structure, the DFG motif (Asp\textsuperscript{157}-Phe-Gly\textsuperscript{159}) in the activation loop of our SBI-0206965-bound structure is displaced away from the ATP active site by a maximum of 2.9 Å (Phe-158/H9251 C), presumably as a consequence of the nonplanar structure of SBI-0206965 relative to compound C. However, in common with the compound C structure, this motif adopts a unique conformation that is neither “DFG-in,” in which Asp-157 is appropriately positioned to chelate Mg\textsuperscript{2+} ions required for coordination of ATP phosphate groups, nor classical “DFG-out,” in which the motif flips by 180° such that the Phe side chain now occludes the ATP pocket and Asp-157 is no longer able to coordinate Mg-ATP for catalysis (Fig. S3B). Instead, the inhibited α2 DFG motif adopts an intermediate conformation; Asp-157 is sterically hindered from adopting the “DFG-in” conformation by the compound N-methylbenzamide and is oriented toward the N-lobe (Fig. S3B), with Phe-158 vacating the regulatory R-spine to displace His-137 in the kinase HRD motif (Fig. 4D). A consequence of this displacement is rearrangement of the HRD backbone and positioning of the Arg-138 side chain to prevent coordination with the phosphate group of pThr-172.

**SBI-0206965 inhibits cellular AMPK signaling**

We explored the efficacy of SBI-0206965 to inhibit AMPK signaling in a range of cell lines. In HEK293 cells, glucose starvation triggered a 1.9-fold increase in pThr-172, indicative of...
energy stress, and this was accompanied by a 3.8-fold increase in pACC (Fig. 5A). The addition of SBI-0206965 to glucose-free medium significantly suppressed pACC at concentrations of $5 \mu M$, probably due to direct AMPK inhibition because SBI-0206965 did not induce reductions in pThr-172, reflecting suppressed LKB1 activity (Fig. 5A). SBI-0206965 concentrations of $5 \mu M$ also blocked AMPK signaling in HEK293 cells treated with AICAR (Fig. 5B); however, this was probably due to inhibition of AICAR cellular uptake, reminiscent of compound C, because SBI-0206965 at $30 \mu M$ did not significantly affect ratios of adenine nucleotides in HEK293 cells, indicative of mitochondrial toxicity; however, we note a trend toward reduced adenylate energy charge at $30 \mu M$ (Fig. S4A). $5 \mu M$ SBI-0206965 also significantly suppressed pACC in the neuronal cell line SH-SY5Y, in which CaMKK2-mediated AMPK signaling was triggered by the addition of the calcium ionophore ionomycin (Fig. 5C).

Discussion

SBI-0206965 was originally identified as an ATP-competitive inhibitor of the autophagy initiator kinase ULK1, with the ability to suppress cellular ULK signaling and ULK1-mediated survival of lung cancer and glioblastoma cells when coupled with nutrient stress (34). In this initial report, AMPK signaling in MEFs (evidenced as ACC phosphorylation) was unchanged following incubation with $50 \mu M$ SBI-0206965, despite increased Thr-172 phosphorylation, which may have arisen from fluctuations in adenine nucleotide ratios at high concentrations (Fig. S4B). SBI-0206965 has subsequently been shown to suppress non-small-cell lung cancer cell growth (36). We now show that SBI-0206965 can be repurposed as a potent and kinase-selective inhibitor of the metabolic coordinator AMPK, providing a useful alternative to the almost singularly used, yet promiscuous, compound C. Because AMPK is a major positive regulator of autophagy through direct phosphorylation of ULK1 (37) and other autophagy-associated proteins (e.g. VSP34 and Beclin-1), simultaneous inhibition of both AMPK and ULK1 signaling supports the application of SBI-0206965 as a highly effective suppressor of prosurvival autophagic responses in tumor cells. We previously used SBI-0206965 as an agent to confirm ULK1-mediated phosphorylation of the drug-sensitizing AMPK $\beta_1$-subunit residue Ser-108, until recently regarded as an autophosphorylation site (8). Our demonstration now that SBI-0206965 also inhibits AMPK does not detract from our previous conclusions, because Ser-108 autophosphorylation is a cis event (10), and in our study, kinase-inactive AMPK enzyme was used as the cellular substrate.

Our biochemical characterization of SBI-0206965 as an AMPK antagonist highlights several limitations of use that...
High-potency AMPK inhibitor

We have confirmed that SBI-0206965 partially shares its binding site on AMPK kinase domain with ATP but under our assay conditions is not ATP-competitive, instead displaying a mixed-type inhibition profile (Fig. 3D). Mixed-type inhibition, by definition, is a conceptual mixture of competitive (increased \( K_m \) substrate) and uncompetitive inhibition (reduced \( V_{max} \)), in which binding of the inhibitor affects substrate binding, and vice versa (44). From a structural perspective, mixed-type inhibition would appear incompatible with overlapping inhibitor/substrate binding sites. However, numerous studies have reported mixed-type inhibition by agents that have either been demonstrated to bind at the enzyme’s active site or most likely do (i.e. transition state inhibitor analogues) (45–50). Specifically, a type II, active site binding inhibitor of insulin-like growth factor 1 receptor tyrosine kinase activity has been reported to display mixed-type kinetics with respect to ATP (45), whereas the tyrphostin inhibitor AG1296 displayed either competitive or mixed-type characteristics, depending on the activation state of its target kinase in platelet-derived growth factor receptor (46). Additionally, crystal structures of adenylate cyclase complexed to P-site inhibitors (adenine nucleosides/adenine nucleoside 3’-phosphates) show active site binding; these inhibitors display either noncompetitive (a special from of mixed inhibition) or uncompetitive kinetic properties (47). Active site binding by mixed inhibitors has also been inferred by direct methods (51, 52) or from close structural relationship to the substrate (53, 54). Type II kinase inhibitors such as SBI-0206965 probably stabilize the “DFG-out” conformation, an arrangement less favorable for ATP binding (55). Thus, the binding sites occupied by SBI-0206965 or ATP represent distinct conformational arrangements, each reducing the binding affinity of the competing ligand.

SBI-0206965 increased \( K_m(\text{ATP}) \), indicating a preference for binding to the free enzyme rather than the enzyme-substrate complex. This is consistent with the location of the inhibitor-binding site resolved in our structure and our observation that SBI-0206965 cannot displace AMPK from ATP-agarose (Fig. 3C). This indicates that SBI-0206965 can only bind to an actively cycling AMPK, as the binding site must first become available following release of the ADP product from the active site. Furthermore, for the inhibitor-bound AMPK complexes, [ATP] in the activity assay may be below \( K_m \) ATP, precluding an increase of IC_{50} at the same rate as increase of [ATP], as dictated by the Cheng–Prusoff equation (56). This probably also explains why SBI-0206965 is very effective at inhibiting AMPK in cells; type II inhibitors typically display high cellular potency, whereas millimolar [ATP] often prevents type I inhibitors (e.g. compound C) from maintaining the potency observed in cell-free activity assays (43).

In summary, we show here that SBI-0206965 displays preferable characteristics, relative to compound C, as an AMPK inhibitor in vitro. Further studies are required to reveal its efficacy in vivo; however, biochemical and structural data provided in this study suggest that SBI-0206965 is a promising lead for the development of a new class of AMPK inhibitors with therapeutic potential.

Experimental procedures

Reagents

Antibodies for pan-AMPK \( \alpha \) (catalog no. 2793, 1:1,000 dilution), and phosphospecific antibodies for \( \alpha \)-p Thr-172 (catalog no. 2535, 1:1,000 dilution) and pACC (catalog no. 3661, 1:1,000 dilution) were from Cell Signaling Technology. Streptavidin-IRDye 680RD (1:20,000 dilution) and anti-rabbit IgG IRDye 680 and anti-mouse IgG IRDye 800 fluorescent-labeled secondary antibodies (1:10,000 dilution) were from LI-COR Biosciences. GSH-Sepharose 4B (catalog no. 17075601) and streptavidin-Sepharose were from GE Life Sciences. Recombinant LKB1/ MO25/STRAD was from Sigma (catalog no. SRP0246). SBI-0206965 was from ApexBio. ATP-agarose was from Innova Biosciences. Complete protease inhibitor mixture was from Roche Applied Science. DNA oligonucleotides and all other reagents were from Sigma.

Kinase inhibitor profiling

Inhibitor profiling (Express screen) was performed by the International Centre for Kinase Profiling, Medical Research Council Protein Phosphorylation and Ubiquitylation Unit (MRC-PPU), University of Dundee, UK.
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Expression constructs

Assays—pET28b plasmid for expression of N-terminal His6 tag α2 kinase domain (residues 6–279) fusion protein was a generous gift from Dr. Dean Littler (Monash University). pET DUET constructs for Escherichia coli expression of AMPK heterotrimer αβγ1 have been described (10, 19).

Crystallization—pET21b plasmid for expression of α2 kinase domain (residues 6–278) incorporating T172D mutation, with a PreScission protease cleavable N-terminal His6 tag, was synthesized by General Biosystems. Constructs were sequence-verified, and expressed constructs were mass-verified by TOF MS.

Protein production

Heterotrimeric human AMPK GST-α1β1γ1 and GST-α2β1γ1, and human FLAG-ULK1 were produced in mammalian COS7 cells as described (8). AMPK and ULK1 were purified using GSH-Sepharose and anti-FLAG M2 affinity gel, respectively. FLAG-CaMKK2 was produced in SF121 insect cells as described (6) and purified using anti-FLAG M2 affinity gel. Thr-172–phosphorylated AMPK His-α2(6–279) kinase domain was produced in bacteria as described (10). Briefly, the pET28b-α2(6–279) construct was transformed into E. coli strain Rosetta (DE3) and expressed by isopropyl 1-thio-β-D-galactopyranoside induction. Cells were ruptured using a precooled EmulsiFlex-C5 homogenizer (Avestin) and protein immobilized on nickel-Sepharose before phosphorylation with CaMKK2. Protein was eluted using 400 mM imidazole and buffer-exchanged with buffer A (50 mM Tris HCl, pH 7.4, 150 mM NaCl, 0.02% Tween 20) using a PD10 desalt column. A similar procedure was used to produce unphosphorylated AMPK α2β1γ1, except the CaMKK2 incubation step was omitted.

Protein crystallization

His-α2(6–278) (T174D) was generated as described above for His-α2(6–279), except the CaMKK2 incubation step was omitted. His tag was removed with GST-tagged PreScission protease treatment (overnight, 4 °C) after the buffer exchange step. PreScission protease was removed using GSH-Sepharose and α2(6–279) (T174D) repurified by SEC. Concentrated α2(6–279) (T174D) (4 mg/ml) was incubated with 0.5 mM SBI-0206965 on ice for 30 min and centrifuged at 10,000 rpm for 3 min before setting crystallization experiments. Protein was mixed equally 1:1 at room temperature with a reservoir solution containing 9–14% ethanol, 5 mM MgCl2, 10 mM tris-(2-carboxyethyl)phosphine, and 3–7 mM MnCl2. Diffraction quality crystals were obtained through streak seeding with a cat’s whisker. Crystals appeared after 2–3 days and reached full size after 1–2 weeks. Crystals were then incubated in a cryoprotectant containing reservoir solution with an addition of 25% ethylene glycol. Data were collected on both MX1 and MX2 beamlines at the Australian Synchrotron (Melbourne, Australia). Data were processed and integrated using XDS (57) and scaled using AIMLESS from the CCP4 suite (58). The structure was solved by molecular replacement using Phaser from the CCP4 suite (59) and 3AQV as the search model. Iterative rounds of model building and refinement were performed using Coot (60) and Buster (https://www.globalphasing.com/buster/)6 (61), respectively. Data for the bromine anomalous map was collected on the MX2 beamline at the Australian Synchrotron near the bromine edge at 13.6 KeV, calculated on the X-ray anomalous scattering website (http://skuld.bmsc.washington.edu/scatter/AS_form.html). An anomalous map was generated by Phenix.refine (62). SBI-0206965 molecular coordinates and restraints were generated using the PRODRG web server (63). Structural validation was performed using Molprobity (64). Omit maps were generated using Buster, and figures were created using PyMOL.

S108tide activity assay

AMPK and ULK1 activities were determined by phosphorylation of the AMPK β1 Ser-108 peptide (S108tide) using 100 μM S108tide, 5 mM MgCl2 and [γ-32P]ATP in a 25-μl reaction volume at 30 °C. Reactions were terminated after 10 min by spotting 15 μl onto P81 phosphocellulose paper (Whatman) and washing in 1% phosphoric acid. Radioactivity was quantified by scintillation counting.

LKB1 phosphorylation assays

200 ng of unphosphorylated His-α2β1γ1 was incubated with 20 ng of LKB1/STRADα/MO25 in the presence of buffer A supplemented with 2 mM MgCl2, 200 μM ATP, and compounds (final 1% DMSO) or 1% DMSO vehicle, as indicated, at 32 °C. Reactions were terminated by the addition of SDS sample buffer and immunoblotted for pThr-172.

Immunoblotting

Samples were separated by SDS-PAGE on a 12% gel (7% for ACC), previously enriched using streptavidin-Sepharose (10), and transferred to Immobilon-FL polyvinylidene difluoride membrane (EMD Millipore). Membrane was blocked in PBS + 0.1% Tween 20 (PBST) with 2% nonfat milk for 30 min at 22 °C and then incubated for either 1 h or overnight with primary antibodies (dilutions in PBST). After washes with PBST, membranes were incubated with anti-rabbit or anti-mouse IgG secondary antibodies, fluorescein-labeled with IR680 or IR800 dye, for 1 h. Immunoreactive bands were visualized on an Odyssey® IR imaging system with densitometry analyses determined using ImageStudioLite software (LI-COR Biosciences).

ATP-agarose immobilization

20 μg of Thr-172–phosphorylated His-α2(6–279) kinase domain was incubated with ATP/compounds for 30 min at room temperature, before the addition of 10 μl of ATP-agarose suspended in 200 μl of buffer B (50 mM HEPES, pH 7.4, 100 mM NaCl, 10% glycerol, 5 mM MgCl2). Beads were incubated on a rotating wheel at 4 °C for 2 h, before washing three times with 500 μl of buffer A supplemented with 0.1% Tween 20. Beads were resuspended in 15 μl of SDS sample buffer, boiled, and immunoblotted for AMPK α.

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Cell culture and incubations

HEK293 and SH-SY5Y mammalian cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM) containing 10% fetal bovine serum and 1% penicillin/streptomycin antibiotics, at 37 °C with 5% CO₂. Cells were incubated with fresh DMEM for 3 h, before simultaneous incubation with glucose-free DMEM and or/compounds as indicated. Glucose-starved cells were washed 4 h post-treatment with ice-cold PBS and harvested by rapid lysis using ice-cold lysis buffer (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 50 mM NaF, 1 mM sodium pyrophosphate, 1 mM EDTA, 1 mM EGTA, 1 mM DTT, 1% Triton X-100, and Complete protease inhibitor mixture). AICAR/ionomycin-treated cells were harvested similarly, except this was performed 1 h post-treatment. Lysates were clarified by centrifugation (14,000 rpm; 3 min; 4 °C) and flash-frozen in liquid N₂ until processed.

Nucleotide measurements

Adenine nucleotides and adenylate energy charge were measured by LC-MS from HEK293 perchlorate extracts, as described previously (32).

ZMP measurements

All ZMP measurements were acquired using LC-MS from perchlorate extracts, using a method similar to that described previously for nucleotides. Briefly, LC conditions were optimized for a 150-mm (length) and 0.5-mm (inner diameter) Hypercarb column (3 μm; Thermo Fisher Scientific Australia Pty. Ltd.). The LC solvent system was (A) 25 mM triethylammonium bicarbonate buffer at pH 7.8 and (B) acetonitrile with 0.1% TFA. ZMP was eluted at a flow rate of 500 μl/min in a gradient program consisting of 100% A (5 min), 0–25% B (10 min), 50–80% B (5 min), and 100% B (5 min). Data were analyzed with Multiquant 2.0, utilizing the area under the LC chromatogram for the corresponding ZMP peak. Calibration curves were obtained by linear regression of the peak area ratio of a ZMP standard. The MS conditions and MRM values for ZMP were optimized by separate infusion of 1 μg/ml solution in 25 mM triethylammonium bicarbonate buffer at a flow rate of 50 μl/min.

Statistical analysis

The data are presented as mean values ± S.E. of at least three independent experiments unless stated. Statistical analyses were performed using GraphPad Prism version 7.0c. Ordinary one-way analysis of variance (ANOVA) with Dunnett’s multiple-comparison test was used for all comparisons unless stated.

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