Snf1 Phosphorylates Adenylyl Cyclase and Negatively Regulates Protein Kinase A-dependent Transcription in Saccharomyces cerevisiae*

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Background: The Snf1/AMPK and PKA pathways are crucial for nutrient sensing and utilization in yeast.

Results: A novel cross-talk mechanism between Snf1/AMPK and PKA is proposed.

Conclusion: Snf1 and Cyr1 interact in a nutrient-independent manner. Active Snf1 phosphorylates Cyr1 and negatively regulates cAMP content and PKA-dependent transcription.

Significance: This is the first evidence of regulation of PKA pathway by Snf1/AMPK.

In eukaryotes, nutrient availability and metabolism are coordinated by sensing mechanisms and signaling pathways, which influence a broad set of cellular functions such as transcription and metabolic pathways to match environmental conditions. In yeast, PKA is activated in the presence of high glucose concentrations, favoring fast nutrient utilization, shutting down stress responses, and boosting growth. On the contrary, Snf1/AMPK is activated in the presence of low glucose or alternative carbon sources, thus promoting an energy saving program through transcriptional activation and phosphorylation of metabolic enzymes. The PKA and Snf1/AMPK pathways share common downstream targets. Moreover, PKA has been reported to negatively influence the activation of Snf1/AMPK. We report a new cross-talk mechanism with a Snf1-dependent regulation of the PKA pathway. We show that Snf1 and adenylyl cyclase (Cyr1) interact in a nutrient-independent manner. Moreover, we identify Cyr1 as a Snf1 substrate and show that Snf1 activation state influences Cyr1 phosphorylation pattern, cAMP intracellular levels, and PKA-dependent transcription.

All organisms need to match growth and metabolism with the availability of nutrients. In the budding yeast Saccharomyces cerevisiae, several mechanisms of sensing of the carbon source present in the environment have evolved (see Refs. 1–3 for extensive review). Nutrients, and in particular glucose, influence the yeast cell metabolic behavior through the activation of signaling pathways, which in turn finely tune transcription and enzymatic activities (4), as well as cell growth and cell cycle (5).

The main signaling pathway activated by glucose in yeast is the PKA pathway, which shows extensive involvement in growth, proliferation, and metabolism (2, 6). The second messenger responsible for the activation of PKA is cAMP, which is synthesized by adenylyl cyclase (Cyr1) upon stimulus from the G-proteins Gpa2 and the RAS pathway (1, 7). cAMP binds to a homodimer of PKA regulatory subunits Bcy1, causing its dissociation from the dimer of catalytic subunits (composed by two of the partially redundant proteins Tpk1, Tpk2, and Tpk3), thus activating the kinase (8). Active PKA has a broad influence on transcription. During exponential growth and in the absence of stress, PKA phosphorylates the transcription factors Msn2,4, thus causing their confinement to the cytoplasm (9). Msn2,4 activate the transcription of genes with promoters containing stress response element sequences, comprising genes coding for chaperones, antioxidant proteins, and proteases and, more in general, for elements necessary to respond to various stress types (10). Besides Msn2,4, PKA controls several other transcription factors including Rim15 (11) and Rap1 (12). Furthermore, PKA directly regulates important metabolic pathways such as glycolysis (6). It phosphorylates proteins catalyzing key enzymatic steps, such as both isoforms of pyruvate kinase, Cdc19 and Pyk2 (13–15).

On the contrary, the signaling pathway centered on the kinase Snf1 is required to adapt to nutrient limitation and to utilize alternative carbon sources, such as sucrose or ethanol (16). Snf1 shares evolutionary conserved functions with the AMP-activated kinase (AMPK) in higher eukaryotes, constituting the well defined family of regulators of cellular energy homeostasis in eukaryotes (17). Snf1 is activated through the phosphorylation of Thr210 by one of the three constitutively active upstream kinases Sak1, Tos3, and Elm1 (18, 19). In

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response to high glucose concentrations, Snf1 is inactivated through dephosphorylation of Thr\(^{210}\) by the Gln7 protein phosphatase (also known as PP1), which is targeted to Snf1 by the adaptor subunit Reg1 (20, 21). However, recent studies highlighted Snf1 function also in glucose-repressed conditions, as well as its partial activation and its role in the regulation of cell cycle (22–25). The most studied function of Snf1 is the regulation of transcription, involving more than 400 genes (26). Active Snf1 causes the translocation to the cytoplasm of the inhibitor Mig1, responsible for the repression of over 90 genes (27). Besides Mig1, Snf1 regulates the activity of various transcription factors. Cat8 and Sip4, which bind carbon source response elements, mostly regulate the expression of gluconeogenic genes (28), and are activated by Snf1-dependent phosphorylations (29, 30). Furthermore, Snf1 regulates through phosphorylation several metabolic enzymes, such as Pkl2, Gpd2, and Acc1 (6), to promote carbon saving in conditions of nutrient scarceness.

In S. cerevisiae, several examples of cross-talk between the Snf1 and PKA pathways have been reported. PKA phosphorylates and contributes to the regulation of the Snf1-activating kinase Sak1 (31) and the β subunit Sip1, preventing the localization of the Snf1 complex to the vacuole (32). Snf1 and PKA share common downstream targets. PKA phosphorylates and deactivates Adr1, the transcriptional activator of several glucose-repressed genes, whereas Snf1 indirectly causes its dephosphorylation and activation (33, 34). Surprisingly, both PKA and Snf1 act as repressors of the transcription factor Msn2, but, whereas PKA is the main regulator of this factor, Snf1 targets it only to shut off transcription of the stress response element regulon after prolonged stationary phase (9, 35, 36). Remarkably, a direct regulation of the Snf1 orthologue AMPK by PKA has been demonstrated in human adipocytes, where PKA phosphorylates Ser\(^{173}\) of AMPK preventing the phosphorylation of residue Thr\(^{172}\), requested for activation (37). Both residues are conserved in S. cerevisiae (Thr\(^{210}\) and adjacent Ser\(^{211}\)), thus a similar mechanism has been hypothesized in yeast (31).

Here, we propose a novel cross-talk mechanism between Snf1/AMPK and PKA pathways. Seeking for new Snf1 targets, we identify adenylate cyclase (Cyr1) as a Snf1-interacting protein. We show that Cyr1 is a Snf1 target in vitro and that its phosphorylation pattern is altered in vivo in the snf1Δ mutant compared with the wild type. Strikingly, we demonstrate that constitutive activation of Snf1 reduces cAMP intracellular content and functionally impairs PKA activity as a regulator of glucose-repressed genes.

### Experimental Procedures

#### Yeast Strains and Growth Conditions

**S. cerevisiae** strains used in this study are listed in Table 1. Synthetic medium contained 2, 5, or 0.05% glucose or 2% ethanol (as indicated in the figures), 6.7 g/liter of yeast nitrogen base (Difco), 50 mg/liter of required nutrients at standard pH (5.5). Cell density of liquid cultures grown at 30 °C was determined with a Coulter counter on mildly sonicated and diluted samples or spectrophotometrically at 600 nm. All experiments were performed with cells in exponential phase of growth, at cell densities between \(A_{600\text{ nm}} = 0.1 (2 \times 10^6 \text{ cells/ml})\) and \(A_{600\text{ nm}} = 0.7 (1.3 \times 10^7 \text{ cells/ml})\).

**Recombinant and Genetic Techniques**—DNA manipulation and yeast transformations were carried out according to standard techniques. To obtain the tagged strains, proteins were tagged with a C-terminal 4HA or 9myc epitope by an in locus 3’ in-frame insertion. *Escherichia coli* DH5α and BL21 (DE3/pLysE) were used in cloning experiments and for expression of recombinant proteins, respectively. The Hpal-Xhol

### Table 1

| Yeast strains used in this work | Strain | Genotype | Origin |
|--------------------------------|--------|----------|--------|
| **BY wt** | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 | Open       | Biostems |
| Snf1Δ | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 snf1::HPH | This study | Ref. 25 |
| Snf1−9myc | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 SNF1-9myc:LEU2 | This study | Biostems |
| Cyr1-TAP | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 CYR1-TAP:His3 | Open       | Biostems |
| Snf1-myc Cyr1-TAP | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 SNF1-9myc:LEU2 CYR1-TAP:His3 | This study | Biostems |
| Snf1-HA T100 | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 SNF1-9myc:LEU2 CYR1-TAP:His3 | Reg1::Ura3 | Biostems |
| Snf1-HA G53R | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 SNF1-9myc:LEU2 CYR1-TAP:His3 | This study | Biostems |
| Snf1-G53R-HA | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 SNF1-9myc:LEU2 CYR1-TAP:His3 | Reg1::Ura3 | Biostems |
| Snf1-G53R-HA | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 SNF1-9myc:LEU2 CYR1-TAP:His3 | This study | Biostems |
| Snf1-9myc Cyr1 | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 SNF1-9myc:LEU2 CYR1-TAP:His3 | Reg1::Ura3 | Biostems |
| Snf1-T510A-4H | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 SNF1-9myc:LEU2 CYR1-TAP:His3 | Reg1::Ura3 | Biostems |
| Snf1-G53R-HA | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 SNF1-9myc:LEU2 CYR1-TAP:His3 | Reg1::Ura3 | Biostems |
| Snf1-9myc Cyr1 | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 SNF1-9myc:LEU2 CYR1-TAP:His3 | Reg1::Ura3 | Biostems |

* a Isogenic to BY4741.
* b Isogenic to W303-1A.
fragment of CYR1 from the YCplac33-CYR1 plasmid (38) was cloned in the HincII-Xhol site of pIVEX2.4a plasmid, originating plasmid pIVEX-CYR1(335–1066). Mutant CYR1-4E (S435E,S527E,T631E,S645E) gene was obtained by synthesis of a custom Hpal-Xhol CYR1 fragment (Eurofins) and subcloning in the YCplac33-CYR1 plasmid.

Protein Extraction, Immunoblotting, and Immunoprecipitation—Samples of cells were harvested and lysed using ice-cold lysis buffer (50 mM Tris, pH 7.5, 150 mM NaCl, 0.1% Nonidet P-40, 10% glycerol) plus 1 mM PMSF, proteases inhibitor mix (Complete EDTA free protease inhibitor mixture tablets; Roche), and phosphatase inhibitor mix (Sigma). An equal volume of acid-washed glass beads (Sigma) was added, and cells were broken by 20 vortex/ice cycles of 1 min each. Extracts were transferred to new tubes and clarified by centrifugation. Protein concentration was determined using the Bio-Rad protein assay. Crude protein extracts were incubated 2 h at 4 °C with anti-HA affinity matrix (Roche) to immunoprecipitate Snf1-HA protein complexes or with IgG-agarose beads (Sigma) to immunopurify Cyr1-TAP. The immunocomplexes obtained after removal of complexes or with IgG-agarose beads (Sigma) to immunopurify Snf1-HA protein affinity matrix (Roche) to immunoprecipitate Snf1-HA protein complexes were immunopurified from 270 mg of total protein extract and resolved by SDS-PAGE followed by staining with GelCode™ blue stain reagent (Pierce) for MS analysis. Bands were excised and treated as reported in the section “Sample Preparation and Mass Spectrometry Techniques.” For the analysis of Cyr1 phosphorylation, Cyr1-TAP was immunopurified from 9 mg of total protein extract and resolved by SDS-PAGE followed by silver staining, according to the manufacturer’s instructions (GE Healthcare).

Recombinant Protein Purification—E. coli strain BL21 (DE3)/pLYsE/ was transformed with pIVEX2.4a-CYR1(335–1066) or pH13-MIG1 (39), cultured in Luria-Bertani broth with 100 mg/liter of ampicillin and 34 mg/liter of chloramphenicol at 37 °C (A600 nm = 0.3), and induced for 2 h with 1 mM isopropyl thio-beta-D-galactoside at 30 °C. His6-Cyr1(335–1066) and His6-Mig1(207–413) were purified on Ni2+–nitrilotriacetic acid beads (Qiagen) and eluted serially with 200, 400, and 800 mM imidazole. Protein concentration was measured by Bradford method using Bio-Rad protein assay kit (Bio-Rad). The purified proteins were dialyzed against a buffer containing 20 mM HEPES, pH 7, 100 mM NaCl, 0.5 mM EDTA, 0.5 mM DTT, 5 mM MgAc, 5% glycerol and stored at −80 °C.

In Vitro Phosphorylation Assays—In vitro phosphorylation of recombinant His6-Mig1(207–413) was carried out as in Ref. 39. In vitro phosphorylation of recombinant His6-Cyr1(335–1066) (4 µg of purified protein) was performed in a buffer containing 20 mM HEPES, pH 7, 100 mM NaCl, 0.5 mM EDTA, 0.5 mM DTT, 5 mM MgAc, using protein kinase Snf1-HA or Snf1-T210A-HA immunopurified from yeast cells growing in 2% glucose. The reaction was started by adding 0.24 µM [γ-32P]ATP (specific radioactivity, 2000 cpm/pmol) and incubated at 30 °C for 30 min. The reaction was stopped by adding 4× SDS sample buffer and then heated for 5 min at 95 °C, and proteins were separated by SDS-PAGE. Phosphorylated bands were visualized by autoradiography.

Sample Preparation and Mass Spectrometry Techniques—The protein content of IP Snf1-HA sample and IP untagged control sample were resolved by SDS-PAGE, analyzed, and compared using a GS-800™ densitometer and Quantity One® analysis software. Each protein band was quantified by densitometric analysis. Only bands exclusively present on IP Snf1-HA sample or bands whose A value differed at least 0.1 in the comparison between IP Snf1-HA sample and IP untagged control sample were excised and analyzed by mass spectrometry. These data generate the list of Snf1-interacting proteins reported in Table 2.

Bands were excised after staining, reduced with DTT, and alkylated with 55 mM iodoacetamide at room temperature for 45 min. Bands were dried, soaked with ammonium bicarbonate 0.1 M, and digested overnight with trypsin sequence grade, at 37 °C using a protease:protein ratio (1:10). Tryptic digests were extracted with 50% acetonitrile (ACN) in 0.1% TFA, desalted/ concentrated on a µZipTipC18 (Millipore), and analyzed by mass spectrometry. LC-electrospray ionization-MS/MS analysis was performed on a Dionex UltiMate 3000 HPLC System with a PicoFrit ProteoPrep C18 column (200 mm, internal diameter of 75 µm) (New Objective). Gradient: 1% ACN in 0.1% formic acid for 10 min, 1–4% ACN in 0.1% formic acid for 6 min, 4–30% ACN in 0.1% formic acid for 147 min, and 30–50% ACN in 0.1% formic acid for 3 min at a flow rate of 0.3 µl/min. The eluate was electrosprayed into an LTQ Orbitrap Velos (Thermo Fisher Scientific) through a Proxeon nanoelectrospray ion source (Thermo Fisher Scientific). Data acquisition was controlled by Xcalibur 2.0 and Tune 2.4 software (Thermo Fisher Scientific). The LTQ-Orbitrap was operated in positive mode in data-dependent acquisition mode to automatically alternate between a full scan (m/z 350–2000) in the Orbitrap (at resolution, 60,000; AGC target, 1,000,000) and subsequent CID MS/MS in the linear ion trap of the 20 most intense peaks from full scan (normalized collision energy of 35%, 10–ms activation). Data Base searching was performed using the Sequest search engine contained in the Proteome Discoverer 1.1 software (Thermo Fisher Scientific). The following parameters were used: 10 ppm for MS and 0.5 Da for MS/MS tolerance, carbamidomethylation of Cys as fixed modification; oxidation of Met and phosphorylation of Ser, Tyr, and Thr, as variable modifications, trypsin (two misses) as protease. To generate the list of phosphosites reported in Table 3, we considered only the sites with the highest X Correlation value (Xcorr) (≥1.5), the rank value of 1 and the best fragmentation pattern, selected manually after visual inspection of the MS/MS spectra. Three different tools for phosphorylation site prediction where applied: NetPhos Yeast 1.0, NetPhosK 1.0 Server-CBS, and Phosida.

cAMP Assay—cAMP was measured using the cAMP Biotrak EIA Assay (GE Healthcare) following the manufacturer’s instructions. Sample preparation was carried out as in Ref. 40.
**Results**

*Snf1 Interacts with Adenylate Cyclase*—Snf1 is known to regulate glucose metabolism and to promote the utilization of alternative carbon sources mainly through the derepression of a large transcription regulon and the phosphorylation of metabolic enzymes (16). However, because the cellular metabolic behavior could be the result of the action of different and somehow opposite signaling pathways, we wondered whether Snf1 could directly regulate such pathways. To detect possible new targets of Snf1, we chose to use a co-immunoprecipitation/MS approach, immunoprecipitating myc-tagged Snf1 and detecting co-immunoprecipitated proteins with mass spectrometry after resolution with SDS-PAGE. We performed this experiment with protein extracts of exponentially growing cells in 2% glucose, because we already demonstrated that in this condition Snf1 is at least partially functional (22, 25).

Interestingly, we identified a total of 50 potentially interacting proteins of which only 16 have been already identified as Snf1 interactors (Table 2) (41–43). We detected most of the components of the complex of Snf1 and of the Snf1/AMPK Regulates cAMP/PKA Pathway

### TABLE 2

| Interactor | Description | Previously identified | Sequence coverage | Peptides matched |
|------------|-------------|-----------------------|------------------|-----------------|
| Snf1 complex | One of three β subunits of the Snf1 kinase complex | Yes | 39.9 | 18 |
| Snf2 | One of three β subunits of the Snf1 kinase complex | Yes | 51.6 | 16 |
| Gal83 | One of three β subunits of the Snf1 kinase complex | Yes | 54.7 | 20 |
| Snf4 | Activating γ subunit of the AMP-activated Snf1p kinase complex | Yes | 76.4 | 24 |
| Sak1 | Upstream serine/threonine kinase for the SNF1 complex | Yes | 35.6 | 30 |
| Reg1 | Regulatory subunit of type 1 protein phosphatase Glc7p | Yes | 39.4 | 27 |
| Glucose metabolism | | | | |
| Cdc19 | Pyruvate kinase | Yes | 48.8 | 14 |
| Hxk2 | Hexokinase isoenzyme 2 | Yes | 18.4 | 7 |
| Pfk1 | α subunit of hetero-octameric phosphofructokinase | Yes | 38.3 | 20 |
| Pfk2 | β subunit of hetero-octameric phosphofructokinase | Yes | 44.9 | 11 |
| Tdh2 | Glyceraldehyde-3-phosphate dehydrogenase, isozyme 2 | No | 79.2 | 21 |
| Tdh3 | Glyceraldehyde-3-phosphate dehydrogenase, isozyme 3 | No | 38.5 | 12 |
| Pdc1 | Major of three pyruvate decarboxylase isozymes | Yes | 47.6 | 14 |
| Eno2 | Enolase II | No | 78.7 | 24 |
| Ald6 | Cytosolic aldehyde dehydrogenase | Yes | 62.8 | 26 |
| Atp2 | β subunit of the F1 sector of mitochondrial F1F0 ATP synthase | Yes | 58.9 | 17 |
| Tal1 | Transaldolase, enzyme in the nonoxidative pentose phosphate pathway | Yes | 45.7 | 13 |
| Ribosomes and translation | | | | |
| Rps8a | Protein component of the small (40S) ribosomal subunit | No | 35.5 | 5 |
| Rpl1a | Ribosomal 60S subunit protein L1A | No | 67.1 | 18 |
| Rpl3 | Ribosomal 60S subunit protein L3 | Yes | 53.8 | 21 |
| Rpl4a | Ribosomal 60S subunit protein L4A | Yes | 48.3 | 13 |
| Rpl6a | Ribosomal 60S subunit protein L6A | Yes | 17.1 | 2 |
| Rpl8a | Ribosomal 60S subunit protein L8A | Yes | 60.9 | 17 |
| Rpl17b | Ribosomal 60S subunit protein L17B | Yes | 8.7 | 1 |
| Rpo9 | Conserved ribosomal protein P0 of the ribosomal stalk | No | 43.6 | 12 |
| Eef1 | Elongation factor 2, also encoded by EFT2 | No | 37.0 | 10 |
| Eef2 | Elongation factor 2, also encoded by EFT1 | No | 62.7 | 41 |
| Tef1 | Translational elongation factor EF-1α, also encoded by TEF2 | No | 23.9 | 4 |
| Tef2 | Translational elongation factor EF-1α, also encoded by TEF1 | Yes | 63.8 | 21 |
| Sro9 | Cytoplasmic RNA-binding protein; associates with translating ribosomes | Yes | 24.7 | 7 |
| Cft1 | RNA-binding subunit of the mRNA cleavage and polyadenylation factor | No | 35.0 | 28 |
| Nip1 | elf3c subunit of the eukaryotic translation initiation factor 3 (elf3) | No | 22.2 | 13 |
| Stam1 | Protein required for optimal translation under nutrient stress | Yes | 42.5 | 12 |
| Amino acid biosynthesis and transport | | | | |
| Lys20 | Homocitrate synthase isozyme | Yes | 61.2 | 20 |
| Lys21 | Homocitrate synthase isozyme | Yes | 47.5 | 19 |
| Cpa2 | Large subunit of carbamoyl phosphate synthetase | Yes | 50.1 | 37 |
| Npr1 | Protein kinase that stabilizes several plasma membrane amino acid transporters | Yes | 58.7 | 28 |
| Fatty acid synthesis | | | | |
| Fas1 | β subunit of fatty acid synthase | Yes | 47.0 | 66 |
| Fas2 | α subunit of fatty acid synthase | Yes | 52.5 | 70 |
| IMP dehydrogenases | | | | |
| Imd1 | Nonfunctional protein with homology to IMP dehydrogenase | Yes | 30.6 | 8 |
| Imd2 | Inosine monophosphate dehydrogenase | Yes | 45.3 | 18 |
| Imd3 | Inosine monophosphate dehydrogenase | Yes | 42.2 | 15 |
| Imd4 | Inosine monophosphate dehydrogenase | Yes | 42.8 | 16 |
| Chaperones | | | | |
| Ssa2 | ATP-binding protein involved in protein folding and vacuolar import of proteins | Yes | 64.8 | 30 |
| Ssb1 | Cytoplasmic ATPase that is a ribosome-associated molecular chaperone | Yes | 49.8 | 21 |
| Ssb2 | Cytoplasmic ATPase that is a ribosome-associated molecular chaperone | Yes | 49.8 | 21 |
| Bmh1 | 14–3–3 protein, major isoform | Yes | 58.4 | 13 |
| Others | | | | |
| Cyr1 | Adenylate cyclase, required for cAMP production and protein kinase A signaling | No | 19.7 | 19 |
| Cka1 | α catalytic subunit of CK2 | Yes | 31.3 | 3 |
| Cop1 | α subunit of COPI vesicle coatamer complex | No | 26.2 | 20 |
## Table 3: Identified phosphopeptides

| Sequence | P site(s) | WT | snf1Δ | Modification(s) | Xcorr | m/z | MH⁺ | Charge | Xcorr | m/z | MH⁺ | Charge |
|----------|-----------|-----|-------|-------------|------|-----|-----|-------|------|-----|-----|-------|
| Ser26    | ×         | Ser26 (phospho) | 2.62 | 775.9883 | 2325.9503 | 3 | 4.73 | 775.9901 | 2325.9558 | 3 |
| Thr30    | ×         | Thr30 (phospho) | 3.11 | 775.9844 | 2325.9386 | 3 | 4.21 | 775.9915 | 2325.9398 | 3 |
| Thr228   | ×         | Thr228 (phospho) | 1.77 | 787.3881 | 2360.1496 | 3 | 2.07 | 787.3915 | 2360.1599 | 3 |
| Ser218   | ×         | Ser218 (phospho) | 2.18 | 816.3416 | 2361.6682 | 3 | 4.03 | 816.3440 | 2361.6689 | 3 |
| Ser287   | ×         | Ser287 (phospho) | 3.83 | 816.3416 | 2361.6682 | 3 | 3.97 | 816.3440 | 2361.6689 | 3 |
| Ser288, Ser291 | (phospho), (phospho) | 3.83 | 816.3416 | 2361.6682 | 3 | 3.97 | 816.3440 | 2361.6689 | 3 |
| Thr350   | ×         | Thr350 (phospho) | 1.76 | 882.0539 | 2644.1471 | 3 | 4.13 | 882.0574 | 2644.1576 | 3 |
| Thr370   | ×         | Thr370 (phospho) | 2.68 | 803.3379 | 2301.9991 | 2 | 2.23 | 803.3417 | 2302.0105 | 3 |
| Ser435   | ×         | Ser435 (phospho) | 1.76 | 774.6774 | 2322.0176 | 3 | 1.51 | 774.6856 | 2322.0423 | 3 |
| Ser527   | ×         | Ser527 (phospho) | 6.29 | 1192.4958 | 2383.9844 | 2 | 5.90 | 1192.4950 | 2383.9827 | 2 |
| Ser1053  | ×         | Ser1053 (phospho) | 3.15 | 940.7603 | 2820.2662 | 3 | 3.06 | 940.7603 | 2820.2662 | 3 |
| Ser1306  | ×         | Ser1306 (phospho) | 1.80 | 803.0618 | 2407.1707 | 3 | 1.52 | 803.0770 | 2407.2165 | 3 |
| Ser1661  | ×         | Ser1661 (phospho) | 4.14 | 857.8657 | 2438.4411 | 4 | 2.91 | 857.8685 | 2438.4523 | 4 |

**Notes:**
- Snf1/AMPK regulates the cAMP/PKA pathway.
- Loss of Snf1 causes an alteration in the phosphorylation pattern.
- Snf1-AKD interacts with Cyr1 in vivo.
- The interaction was not dependent on glucose concentration.
- Detectable in extracts from reg1Δ cells (Fig. 2B) and was detected in the presence or the absence of Snf1.
- Phosphorylation status of adenylate cyclase in vivo was determined by TAP-tagged Cyr1 from WT and snf1Δ strains.
- Extracts and after resolution by SDS-PAGE and silver staining were analyzed.
- Phosphoproteomics chips were conducted.
- Seven phosphosites were identified.
- Snf1 is a phosphoprotein, and several phosphorylated residues have been identified in proteomic studies (45–48).

**References:**
- This finding was of particular interest, because it was previously shown that Snf1, Mig1, and Mig2 are Snf1-dependent, not being detectable in the absence of Snf1.
- In a WT strain, we identified a set of phosphorylated sites.
- Strikingly, we also identified several additional sites.
- One of the newly identified interactors was adenylate cyclase (Cyr1), the essential metabolic processes, such as glycolytic enzymes, and in such terms of cAMP, activator of PKA (Fig. 1 and Table 2). To validate this finding, we performed Co-IP experiments, density degrading TAP tagged Snf1, and K84R or the presence of a constitutively active mutant, Snf1-G53R (Fig. 2D). Furthermore, we identified TAP tagged Snf1, K84R or in the presence of a constitutively active mutant, Snf1, G53R (Fig. 2E). Although, we were able to demonstrate that the interaction was particu-
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we checked whether the identified sites were contained in consensus sequences of a specific kinase, we failed to identify any obviously responsible for the phosphorylation events. These data provide a substantial advance in mapping possible regulatory post-translational modifications of adenylate cyclase and demonstrate the necessity of Snf1 to preserve the proper phosphorylation pattern of Cyr1.

Snf1 Phosphorylates the RAS-associating Domain of Adenylate Cyclase in Vitro—Because we demonstrated that a lack of Snf1 could influence the phosphorylation pattern of Cyr1 in vivo, we sought for direct kinase activity of Snf1 on adenylate cyclase in vitro. Because we wanted to demonstrate a direct kinase activity of immunopurified Snf1, discerning from phosphorylation events dependent on kinases co-immunoprecipitate with Snf1 (Table 2), we searched for the most suitable catalytic-deficient mutant of Snf1 to be included as a control. We therefore tested different Snf1 mutants for their ability to phosphorylate recombinant Mig1 in vitro. Strikingly, the unphosphorylatable mutant Snf1-T210A turned out to be more impaired than the Snf1-K84R mutant (Fig. 4A) and was chosen as a control for subsequent analysis. We then purified from E. coli a His6-tagged region of Cyr1 containing the whole RAS-associating domain (RAD), four of five (Ser435, Ser527, Thr631, and Ser645) residues whose phosphorylation was found to be Snf1-dependent (Fig. 3A) and two residues contained in Snf1 phosphorylation consensus sites (Ser554 and Thr736). We then performed an in vitro kinase assay with 32P-labeled ATP, using the purified protein as substrate and immunoprecipitated HA-tagged Snf1 and Snf1-T210A as kinases. In the presence of the WT kinase, but not in the presence of the Snf1-T210A mutant, the His6-Cyr1(335–1066) recombinant protein incorporated ATP, demonstrating that this domain of adenylate cyclase is a bona fide substrate of Snf1 in vitro (Fig. 4B).

To determine which residues of His6-Cyr1(335–1066) were phosphorylated in vitro by Snf1, we carried out a phosphorylation assay, followed by SDS-PAGE resolution and MS analysis. Remarkably, all the sites whose phosphorylation was Snf1-dependent in vivo (Fig. 3A and Table 3) turned out to be phosphorylated by Snf1 in vitro (Ser435, Ser527, Thr631, and Ser645) (Fig. 4C and Table 4), and among these only Ser435 and Ser527 were slightly phosphorylated in the presence of Snf1-T210A, possibly evidencing a residual activity of the mutant kinase (Table 4). Remarkably, two phosphosites that turned out to be Snf1-independent in vivo (Ser483 and Ser524) and two sites that were phosphorylated upon SNF1 deletion (Thr1043 and Thr1046) were also found to be phosphorylated in vitro by Snf1 (Fig. 4C and Table 4). These data demonstrate that although the in vivo Snf1-dependent phosphosites are not contained in perfect Snf1 consensus sequences, they are potential targets of direct phosphorylation by Snf1.

Catalytic Activation of Snf1 Negatively Regulates cAMP Content—The function of adenylate cyclase is to catalyze the synthesis of cAMP, the second messenger responsible for the activation of PKA, in response to stimuli from the RAS and GPCR systems (1). Having shown that a lack of Snf1 influences the phosphorylation pattern of Cyr1 and that Snf1 can phosphorylate Cyr1 in vitro, we wondered whether activation of Snf1 could influence the activity of adenylate cyclase. cAMP is rapidly synthesized at high levels after glucose refeeding to starved cells but is present at detectable basal levels also in exponentially growing cells (40). Therefore we measured cAMP levels in log phase cells growing in 2% glucose, either lacking Snf1 or...
harboring WT Snf1 or a Snf1-G53R mutant, which is constitutively activated because of the persistent phosphorylation of Thr210 (49).

Strikingly, although a lack of Snf1 did not influence the cAMP content of cells growing in glucose-repressed conditions, the presence of the Snf1-G53R mutant nearly halved the cAMP levels, as compared with the WT (Fig. 5). These data demonstrate that the catalytic activity of Snf1 can reduce the intracellular content of cAMP in glucose-repressed condition.

Catalytic Activation of Snf1 Negatively Influences PKA-dependent Transcriptional Expression—PKA regulates several cellular processes including nutrient sensing, energy metabolism, cell cycle progression, thermotolerance, osmotic shock tolerance, sporulation, pseudohyphal growth, aging, and autophagy (1). Some of these processes, such as thermotolerance, are usually used to indirectly measure the activity of the PKA pathway (8). However, Snf1 regulates these processes independently from PKA (50), and the use of these methods to measure the influence of Snf1 catalytic activity on the PKA pathway was therefore discarded.

Thus, we chose to indirectly measure the activity of the PKA pathway through the effect on the transcription of genes which are regulated by transcription factors that are PKA targets. Some transcription factors are shared downstream targets of Snf1 and PKA (9, 32–36); thus the strategy required the identification of Snf1-independent target genes.

Several PKA-regulated genes have been mapped (51), and we checked the dependence of selected genes (HXT1, HXT7, and ATR1) during a nutritional shift-up (from ethanol to glucose containing medium) using a cyr1Δpde2Δyak1Δ strain, in which the disruption of adenylate cyclase impairs the activation of PKA (35) (Fig. 6A). Because HXT1, whose expression was PKA-dependent, is also known to be regulated by Snf1 (52), it was included in subsequent analyses as a control.
Snf/AMPK Regulates cAMP/PKA Pathway

To assess whether Snf1 catalytic activity influenced the expression of PKA-regulated genes, we tested the expression of HXT1, HXT7, and ATR1 in the presence of the aforementioned Snf1-G53R mutant. HXT1 turned out to be down-regulated in a Snf1-G53R strain grown in 2% glucose (Fig. 6B), thus proving that this mutant is constitutively active. The PKA-activated gene ATR1 was slightly down-regulated in the presence of a Snf1-G53R mutant, but more strikingly, in the same strain the PKA-repressed gene HXT7 was 5-fold derepressed (Fig. 6B). HXT7 is known to be Snf1-independent (4), and we also checked its independency from the Snf1-inhibited repressor Mig1 using a mig1Δ strain (Fig. 6C), demonstrating that this gene is a useful PKA-dependent/Snf1-independent reporter gene. To further characterize the effect of Snf1 activity on PKA-dependent genes, we tested the expression of HXT1, HXT7, and ATR1 in the presence of a Snf1-G53R mutant in 0.05, 2, and 5% glucose. In cells growing in 0.05% glucose, where PKA activity is already low and Snf1 activity is high, all three genes were equally expressed in both the WT and the Snf1-G53R strain (Fig. 6D). Strikingly in 5% glucose, where PKA activity should be maximal, activation of Snf1 through the G53R mutation still causes the deregulation of PKA-dependent genes (Fig. 6D). Altogether, these data demonstrate that Snf1 catalytic activity functionally influences the PKA pathway, determining an alteration in the expression of PKA-dependent genes.

Mutation of Snf1-dependent Phosphorylation Sites of Cyr1 Influences PKA-related Phenotypes—We showed that in the presence of active Snf1, cAMP levels were low (Fig. 5). In yeast, two phosphodiesterases, Pde1 and Pde2, degrade cAMP, and Pde1 is subjected to activation by PKA phosphorylation, generating a negative feedback on the pathway (53). Thus, to assess whether the observed effect of Snf1 on cAMP content and PKA-dependent gene expression was due to activation of the phosphodiesterases, we tested the expression of HXT1, HXT7, and ATR1 in a pde1Δpde2Δ strain in the presence of a Snf1-G53R mutant. Interestingly, deregulation of PKA-dependent genes by Snf1 activation was observed even in the absence of the phosphodiesterases (Fig. 7A). Only the derepression of HXT7 was somehow reduced, but the basal levels of the mRNA were already high in the pde1Δpde2Δ strain compared with a WT (data not shown).

We therefore tested the possibility that the PKA transcriptional activity was instead dependent upon phosphorylation of adenylyl cyclase. Thus, we generated a mutant Cyr1 in which the four Snf1-dependent phosphosites near the RAD (Ser635, Ser527, Thr631, and Ser645), identified in vivo (Fig. 3A and Table

### TABLE 4

| Phosphorylatable residue | Immunoprecipitated kinase | Snf1-HA | Snf1-T210A-HA |
|--------------------------|---------------------------|---------|---------------|
| Found                    | Xcorr                     | 5.98    | 3.44          |
| Ser835                   | X                         | 1.84    |               |
| Ser483                   | X                         | 3.57    | 1.98          |
| Ser497                   | X                         | 4.06    | 1.65          |
| Ser224                   | X                         | 3.24    |               |
| Thr209                   | X                         | 2.24    |               |
| Thr1043                  | X                         | 1.71    |               |
| Thr1046                  | X                         | 1.50    |               |
| Ser1052                  | X                         |         |               |

FIGURE 4. Snf1 phosphorylates the Cyr1 RAS-associating domain in vitro. A, autoradiography of the in vitro kinase assay with γ-32P-labeled ATP of recombinant His6-Mig1(207–413). The different lanes indicate the strain from which the kinase was immunopurified. A kinase reaction without substrate is shown as control. B, autoradiography of the in vitro kinase assay with γ-32P-labeled ATP of recombinant His6-Cyr1(335–1066). The different lanes indicate the vector contained by host E. coli from which purification of the substrate protein was performed. A kinase reaction without substrate is shown as control. C, schematization of the phosphosites identified by MS analysis on recombinant His6-Cyr1(335–1066) after in vitro phosphorylation with immunopurified Snf1-HA.

FIGURE 5. A constitutively activated Snf1 mutant reduces intracellular cAMP levels. Histogram showing the intracellular cAMP content of the strains expressing Snf1 or Snf1-G53R, growing in exponential phase. The data are normalized on cellular dry weight. *, p value < 0.05.

FIGURE 6. Effect of Snf1 catalytic activity on PKA-related phenotypes. A, expression of PKA-regulated genes in the presence of a Snf1-G53R mutant in 0.05, 2, and 5% glucose. B, PKA activity in the presence of a Snf1-G53R mutant. C, immunoprecipitation of Snf1 and Snf1-T210A with anti-HA epitope.
3) and confirmed to be Snf1 targets in vitro (Fig. 4C and Table 4), were mutated to glutamate to mimic constitutive phosphorylation. The resulting mutant, Cyr1-4E, had no effect on duplication time (WT 117 ± 13 min, Cyr1-4E 134 ± 5 min), cell volume (WT 58.8 ± 0.8 fL, Cyr1-4E 58.3 ± 1.1 fL) and protein content (WT 400 ± 40 a.u., Cyr1-4E 417 ± 37 a.u.). Although the mutant protein was expressed at the same level as WT Cyr1 (Fig. 7B), it induced, in a cyr1Δpde2Δyak1Δ background, a higher resistance to heat shock (Fig. 7C), a typical hallmark of PKA activity impairment (54). Furthermore, we tested the expression of HXT7 in a Cyr1-4E mutant compared with the WT. Interestingly, we observed a more than 2-fold derepression of HXT7 in the presence of the mutated Cyr1, indicating a partial down-regulation of the pathway (Fig. 7D).

Taken together, these data indicate that Cyr1 phosphorylation, rather than phosphodiesterase activation, is important for the control of HXT7 expression. In particular, the phosphorylation status of four Snf1-dependent residues seems to directly influence PKA activity.

Discussion

A major challenge in the understanding of a signaling pathway is the discovery of cross-talk with other pathways. Because Snf1 is mainly known for its role in shaping metabolism to guarantee cell growth in nutritional scarceness, it seemed likely that it could regulate other signaling pathways that cause opposite outcomes. In search for new targets of Snf1, we identified a set of previously unknown Snf1-interacting proteins (Fig. 1 and Table 2), including adenylate cyclase (Cyr1), suggesting a possible cross-talk between Snf1 and PKA pathways. PKA was a possible candidate as a Snf1-regulated pathway, because it is activated in conditions of nutrient abundance. The two pathways share common downstream targets, such as transcription factor Adr1 (9, 33–36), and PKA has been reported to regulate...
the recruitment of Snf1 to the vacuole and to contribute to the inactivation of the Snf1-activating kinase Sak1 (31, 32). This regulation seems to be evolutionarily conserved, because in mammalian cells PKA phosphorylates and inactivates the Snf1 orthologue AMPK (37). Thus, showing the converse regulation of the PKA pathway by Snf1, we suggest a novel cross-talk mechanism. Although we showed that the interaction of Snf1 and Cyr1 is independent from nutritional conditions (Fig. 2, C and D) and from Snf1 activity (Fig. 2, E and F), we demonstrated that the lack of Snf1 could alter the phosphorylation status of the PKA pathway by Snf1, we suggest a novel cross-talk mechanism. Although we showed that the interaction of Snf1 and Cyr1 is independent from nutritional conditions (Fig. 2, C and D) and from Snf1 activity (Fig. 2, E and F), we demonstrated that the lack of Snf1 could alter the phosphorylation status of Snf1-dependent phosphorylation sites in the Cyr1 RAD affects PKA-dependent transcription. A, real time PCR quantification of mRNAs of the indicated genes in pde1Δpde2Δsnf1Δ cells expressing Snf1 or Snf1-G53R in exponential phase of growth in 2% glucose. B, Western blot showing the detection of Cyr1 in the cyr1Δpde2Δyak1Δ strain transformed with the indicated plasmid. Swi6 was used as loading control. C, drop test on YPD plates of 10⁴ cells of the cyr1Δpde2Δyak1Δ strain transformed with the indicated plasmid after heat shock at 51 °C for the indicated time. D, real time PCR quantification of HXT7 mRNA in cyr1Δpde2Δyak1Δ cells transformed with the indicated plasmid in exponential phase of growth in 2% glucose.

FIGURE 8. Map of the cross-talk between the Snf1 and PKA pathways. Schematization of the basic circuitry of the cross-talk between the Snf1 and PKA pathway, applied to the expression of the PKA-repressed gene HXT7. Subsidiary pathways were omitted for simplification. Dotted lines and solid lines indicate inefficient and efficient processes respectively. In high glucose condition, Cyr1 produces high cAMP levels, which activate PKA that can inhibit Msn2,4 translocation into the nucleus and Sak1 activity. Therefore, expression of HXT7 is low and Sak1 inefficiently phosphorylates Snf1. In low glucose, Snf1 is active and inhibits Cyr1, thus resulting in lower cAMP levels and lower PKA activity; Msn2,4 is nuclear and activates HXT7 expression.
Cyr1 in vivo and that Snf1 can phosphorylate an important regulatory domain of adenylate cyclase in vitro (Figs. 3 and 4B). Moreover, by mapping the in vitro phosphorylated sites, we demonstrated that Snf1 directly phosphorylates the phosphosites, which turned out to be Snf1-dependent in vivo (Fig. 4C and Table 4). Strikingly, by using a constitutively activated Snf1 mutant, we highlighted an inverse correlation between the activation state of the kinase and the intracellular content of cAMP (Fig. 5). To determine whether the activity of Snf1 had actual influence on PKA activity, we established a method based on the expression of an endogenous gene used as a reporter. The gene HXT7, identified as a target of the PKA-repressed transcription factors Msn2 and Msn4 (51), was chosen for this purpose. In fact, it showed PKA-dependent repression in a strain lacking adenylate cyclase (Fig. 6A), and it was independent from direct Snf1 activity in a miglΔ strain (Fig. 6C). Moreover, HXT7 was already reported to be Snf1-independent (4), and no interaction of Snf1-regulated transcription factors with the promoter of HXT7 was reported in specialized databases. HXT7 turned out to be dramatically derepressed in a Snf1-G53R strain (Fig. 6B), thus linking Snf1 activation to the transcription of PKA-dependent genes. Moreover, the phosphomimetic mutation of Snf1-dependent Cyr1 phosphosites induced alteration of heat shock resistance and caused HXT7 derepression. HXT7 encodes for a high affinity glucose transporter, specifically expressed in low glucose media (55), and the dependence of its expression from an interplay between PKA and Snf1, which act as opposite pathways, seems therefore biologically consistent. The regulation of PKA pathway by Snf1 seems likely to occur in opposite pathways, seems therefore biologically consistent.  

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