Supplemental Material

Genome-wide cooperation by HAT Gcn5, remodeler SWI/SNF, and chaperone Ydj1 in promoter nucleosome eviction and transcriptional activation

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## Supplemental References

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Supplemental Methods

Yeast strain constructions

WT strain BY4741 and kanMX-marked deletion derivatives were described previously (Winzeler et al. 1999) and purchased from Research Genetics, and all deletions in these strains were verified by PCR analysis of genomic DNA either previously (Swanson et al. 2003) or in the current study. Double mutants harboring hphMX4-marked deletions were produced from the appropriate single mutants containing a kanMX4-marked deletion by gene replacement of the relevant WT chromosomal gene with a cassette containing hphMX4 in place of the CDS, amplified from plasmid pAG32 with the appropriate primers (Goldstein and McCusker 1999). The single mutants used in these gene replacements were complemented by a URA3 plasmid containing the WT allele of the kanMX-deleted gene, which was evicted by growth on 5-FOA after generating the second hphMX-marked second deletion. We verified that the slow-growth phenotypes of the ydj1Δsnf2Δ and ydj1Δgcn5Δ double mutants could be partially complemented by YDJ1 on plasmid pHQ2063 (construction of this plasmid described below). To construct HQY1602 and HQY1603, hsc82Δ::kanMX4 strain 771 was transformed with plasmid pHQ2062 (HSP82, URA3) and chromosomal HSP82 was replaced with hsp82Δ::hphMX4 to produce HQY1601; which was subsequently transformed with pHQ2066 (hsp82-T101I, LEU2) or pHQ2067 (hsp82-G170D, LEU2), and derivatives lacking pHQ2062 were selected on 5-FOA medium. Strains carrying P_{TET}-SNF2 were constructed by gene replacement with HIS3::P_{TET}-SNF2 DNA fragments PCR-amplified from plasmid pHQ2078 using forward primer: 5’-TGGA CCTTTTGT A T A T AAA TCA TCGGGAA GGTCA GCT A TTCTGTTGTTTCTCT- AA TCGCGA CTTTCTGCT A GCTGAA GCTTCGT A CGC 3’ (with SNF2 promoter sequences -100 to -31 italicized) and reverse primer: 5’-TCAAA TGCTGCCA TCTT AAA T A GCA GCGGTTG- A CCTCTTCGTTGCT AAA TTGA CGCTGTGGT A TGTTCA T A T A GGCCA CT A GTGGA TCTG 3’ (with SNF2 CDS sequences +70 to +1 italicized). Replacement of SNF2 promoter with the H13MX-tT A-tetO2 cassette was confirmed by PCR analysis and the Western blot analysis of Snf2 in Fig. S5.

Plasmids

A ~2.25 kb fragment of YDJ1 containing the 635 (5’) or 381bp (3’) of flanking sequences was PCR-amplified from WT genomic DNA with Asp718 and XbaI restriction sites added at the 5’ and 3’ ends, and cloned into YCplac33 to produce pHQ2063. A ~3.2 kb fragment of HSP82 containing the 2130bp CDS and 800 (5’) or 237 bp (3’) of flanking sequences was PCR-amplified from WT genomic DNA with BamHI and SalI restriction sites added at the 5’ and 3’ ends, and cloned into YCplac33 to produce pHQ2062. For construction of hsp82-T101I, a ~1.1 kb fragment containing the 5’ portion of HSP82 containing a mutation encoding the T101I substitution (introduced using the mutagenic oligonucleotide reverse primer) and BamHI and Eco47III restriction sites at the 5’ and 3’ ends, and a ~2.1 kb fragment of the 3’ portion of HSP82 with Eco47III and SalI restriction sites at the 5’ and 3’ ends, were PCR-amplified from WT genomic DNA. These two PCR fragments were digested with BamHI and Eco47III, or Eco47III and SalI, respectively, and inserted into YCplac111 between the BamHI and SalI sites to produce pHQ2066. Similarly, for construction of hsp82-G170D, a ~1.3 kb fragment containing the 5’ portion of HSP82 encoding the G170D mutation and BamHI and FspI restriction sites at the 5’ and 3’ ends, and a ~1.9 kb fragment of the 3’ portion of HSP82 with Eco47III and SalI restriction sites at the 5’ and 3’ ends, were PCR-amplified from WT genomic DNA. The PCR fragments were digested with BamHI and FspI or Eco47III and SalI, respectively, and inserted into YCplac111 between the BamHI and SalI sites to produce pHQ2067. A BglII-EcoRI
kanMX fragment of the kanMX-tTA-tetO2 cassette of pCM224 (Bellí et al. 1998) was replaced with the HIS3MX6 BglII-EcoRI fragment from pFA6a-3HA-His3MX6 (Longtine et al. 1998) to produce pHQ2078.

Paired-end sequencing of chromatin immunoprecipitates (PESCI)

The procedure described by (Cole et al. 2014) for preparing DNA libraries for Illumina pair-end sequencing was followed with the following modifications. Immediately prior to harvesting, cells were fixed with 1% formaldehyde (20 min at room temperature) and glycine was added to 0.3 M for 5 min. Cells were collected by centrifugation, washed twice with ice-cold Tris-buffered saline (pH 7.5), and cell pellets were stored at -80°C. Each frozen cell pellet from 100 mL culture was suspended in 0.4 mL FA buffer (50 mM Heps-KOH [pH 7.5], 140 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.1% sodium deoxycholate, 1 mM PMSF, 1 µg/mL leupeptin, 1 µg/mL pepstatin A and 10 µg/mL aprotinin). 0.6 mL glass beads were added, and cells were vortexed for 45 min at 4°C. The beads were removed, FA buffer added to 1 mL, and the cell lysate was spun at 14,000 rpm for 10 min at 4°C. The pellet was resuspended in 1 mL FA buffer and spun again. The final pellet was resuspended in 0.5 mL FA buffer and sonicated using a Diagenode Bioruptor 300 at 3°C with 40 cycles of 30 s at high power and 30 s at rest, with a 15 min cooling period after 20 cycles. Debris was removed by spinning for 20 min at 4°C. The supernatant was divided into 100 µL aliquots and stored at -80°C. A 100 µL aliquot of supernatant was adjusted to 200 µL by adding 100 µL FA buffer, 4 µL of 1M Tris (pH8.0), 2 µL of 0.5 M EDTA and 10 µL of 10% SDS and incubated at 65°C overnight to reverse the cross-links. DNA was purified by treating with proteinase K (add 100 µg proteinase K) for 4 h at 55°C, extracting twice with an equal volume of chloroform, and precipitating with 0.7 volume of isopropanol and washed with 1 mL of 70% ethanol. Purified DNA was quantified using the NanoDrop ND-1000. For immunoprecipitation, anti-H3 antibody (1 µg, Abcam ab1791) or anti-Rpb3 antibody (1.5 µL, NeoClone, W0012) was first incubated in 200 µL FA buffer for 3 h with 50 µL of Dynabeads anti-Rabbit IgG (Invitrogen, Cat # 11204D) or 50 µL of Dynabeads Pan Mouse IgG (Invitrogen, Cat # 110.42), respectively, that had been pre-washed twice with 500 µL PBS/BSA buffer. The resulting anti-H3 or anti-Rpb3 conjugated Dynabeads were washed twice with 500 µL PBS/BSA, and mixed with chromatin extract containing 2.5 µg DNA and FA buffer at a final volume of 200 µL, and rotated at 4°C for 3 h. The beads were washed twice with 1 mL FA buffer containing 0.25% SDS, once with 1 mL FA buffer, twice with 1 mL Wash buffer II (50 mM Heps-KOH [pH7.5], 500 mM NaCl, 1 mM EDTA, 1% Triton X-100 and 0.1% sodium deoxycholate), twice with 1 mL Wash buffer III (10 mM Tris-Cl [pH8.0], 250 mM LiCl, 1 mM EDTA, 0.5% NP-40 and 0.5% sodium deoxycholate), and once with 1 mL TE. Bound chromatin was eluted with 100 µL elution buffer (50 mM Tris-Cl [pH8.0], 10 mM EDTA and 1% SDS) at 65°C for 15 min and the eluate collected. Elution was repeated with 150 µL elution wash buffer (10 mM Tris-Cl [pH8.0], 1 mM EDTA and 0.67% SDS) for 10 min at 65°C. The eluates were pooled and incubated at 65°C overnight to reverse the cross-links, followed by proteinase K digestion (0.4 mg/mL, 4 h, 55°C). The digest was cooled, mixed with one-fifth volume of 5M potassium acetate, and extracted twice with an equal volume of chloroform. DNA was precipitated with 0.7 volume isopropanol in the presence of 15 µg glycoBlue (Ambion/Invitrogen), washed once with 70% ethanol and dissolved in 45 µL TE (10 mM Tris-Cl [pH 8.0], 0.1 mM sodium EDTA). Immunoprecipitated purified DNA was repaired as described (Cole et al. 2012), purified using a Qiagen PCR column and eluted with 50 µL TE. A 5’-phosphate and a 3’-dA were added, followed by ligation to Illumina paired-end adaptor (cat. #1005711) (Cole et al. 2012). The adaptor-ligated immunoprecipitated...
DNA (5 µL for anti-H3 or 7.5 µL for anti-Rpb3) was subjected to PCR (22 cycles for H3 and 25 cycles for Rpb3) in 50 µL with Illumina paired-end primers (InPE1.0 and InPE2.0) and an index primer at the concentrations recommended by the vendor. DNA was purified using a Qiagen PCR column. To remove primer-dimers and larger DNA fragments, 200-450 bp fragments were resolved by gel electrophoresis using a precast 2% agarose gel and E-Gel system (Invitrogen) and purified using Zymo clean Gel DNA Recovery Kit, and the final concentration was determined with a Qubit 2.0 Fluorometer (Invitrogen). DNA fragment size was verified by agarose gel electrophoresis prior to sequencing.

Supplemental Results

Comparison of conventional H3 ChIP and H3 ChIP-seq results for ARG1, HIS4, ARG4 and CPA2

Overall, there is good agreement between the measurements of H3 occupancy by ChIP-seq (Figs. S13-S15 summarized in Fig. S18) and conventional ChIP (Fig. S3B) for the 4 canonical target genes ARG1, HIS4, ARG4 and CPA2. Both approaches implicated only Ydj1 and Snf2 at ARG1 and all three co-factors at HIS4. Whereas conventional ChIP revealed roles for all three co-factors at ARG4 (Figs. 1E & S3B), only Gcn5 and Ydj1 were implicated at this gene by ChIP-seq measurements (Fig. S18). However, snf2Δ produced a defect in H3 eviction at ARG4 in ChIP-seq assays (Fig. S13A) with a p-value of 0.09 (Fig. S18 row 9), just above the threshold of p<0.05 used to construct the Venn diagram in Fig. 4C. Given the high precision of conventional ChIP measurements for ARG4 (Fig. S3B), we elected to place ARG4 among the group of genes employing all three co-factors for H3 eviction (Fig. 4C). Whereas conventional ChIP and ChIP-seq measurements both implicated Gcn5 and Snf2 at CPA2, the role of Ydj1 was clearly established only by conventional ChIP analysis. ChIP-seq measurements for CPA2 in Fig. S13D are suggestive of an additive effect of snf2Δ and ydj1Δ, consistent with Ydj1 functioning at CPA2; however, because this was not observed for the corresponding gcn5Δ/ydj1Δ ChIP-seq data in Fig. S14D, we refrained from implicating Ydj1 at CPA2 in Fig. 4C.
Supplemental Figure S1. SM-induced depletion of histone H3 from promoters of four canonical Gcn4 target genes requires Gcn4. (A) Sequence coordinates of primers for conventional ChIP analysis of promoter occupancies relative to the ATG start codon. Coordinates of TSS were obtained from [Xu et al., 2009], and coordinates of -1 and +1 nucleosomes determined as described in Fig. S10B. (B-E) Promoter occupancies of H3 measured by conventional ChIP in the indicated strains cultured in SC medium (U) or treated with sulfometuron methyl (SM) for 30 min to induce Gcn4 (I). Cross-linked chromatin was immunoprecipitated with anti-H3 antibodies and DNA from immunoprecipitated (IP) and input samples was subjected to PCR in the presence of [33P]-dATP to amplify radiolabeled fragments of the relevant gene or non-transcribed sequences from chromosome V analyzed as a control, which were resolved by PAGE and quantified by phosphorimaging analysis. Ratios of target gene to ChrV signals in IP samples were normalized to the corresponding ratios for Input samples to yield occupancy values. Mean (+/-SD) values were calculated from 3 biological replicates.
Supplemental Figure S2. Snf2 and Gcn5 promote H3 eviction from promoters of Gcn4 target genes only under inducing conditions. Conventional ChIP analysis of H3 in the indicated strains under inducing (I) or uninducing (U) conditions, conducted as in Fig. 1. Mean (±SD) values were calculated from 3 biological replicates.
### Supplemental Figure S3.

Summary of H3 EDI values and Gcn4 occupancies at canonical Gcn4 target genes in different mutants. (A-B) H3 and Gcn4 occupancies were determined by conventional ChIP analysis, as in Fig. 1A-D, and EDI values were calculated as (H3\(_{\text{mutant}}\) - H3\(_{\text{WT}}\))/\((\text{H3}_{\text{mutant}}^{\text{gcn4\_D}}) - \text{H3}_{\text{WT}}\))

| Mutant | H3 EDI | Gcn4 occupancy (%WT) |
|--------|--------|----------------------|
|        | ARG1   | ARG4   | HIS4 | CPA2 | ARG1 | ARG4 | HIS4 | CPA2 |
| 1      | gcn4\_A |        |      |      |      |      |      |      |      |
| 2      | gal1T\_A/med15\_A | -0.1 | -0.02 | -0.12 | -0.11 | 1.2 | 1.2 | 1.0 | 1.1 |
| 3      | snf2\_A | -0.16 | 0.15 | 0.83 | -0.05 | 2.1 | 1.6 | 1.0 | 1.5 |
| 4      | snf2\_A | -0.08 | 0.05 | -0.1 | -0.04 |      |      |      |      |
| 5      | ada1\_A | -0.09 | 0.61 | 0.25 | 0.1 | 1.8 | 1.1 | 1.0 | 0.98 |
| 6      | ada1\_A | -0.31 | 0.39 | 0.11 | 0.05 | 2.2 | 1.4 | 1.4 | 1.3 |
| 7      | adf1\_A | -0.27 | -0.09 | -0.15 | -0.2 | 1.7 | 1.4 | 2.1 | 1.8 |
| 8      | yrt109\_A | -0.11 | -0.06 | 0.06 | -0.18 |      |      |      |      |
| 9      | gcn4\_A | 0.29 | 0.37 |      |      | N/A | 0 | 0.41 | 0.31 |
| 10     | snf2\_A | -0.12 | -0.14 | -0.21 | -0.25 |      |      |      |      |
| 11     | snf2\_A | 0 | 0.14 | 0.92 | -0.02 | 1.4 | 1.2 | 1.1 | 1.2 |
| 12     | ydt1\_A | -0.09 | -0.1 | -0.12 | -0.2 |      |      |      |      |
| 13     | ydt1\_A | -0.15 | -0.02 | -0.02 | -0.03 |      |      |      |      |
| 14     | ydt1\_A | -0.1 | -0.06 | -0.16 | -0.03 |      |      |      |      |
| 15     | ada1\_A | 0.19 | 0.13 | 0.14 | 0.13 | 0.84 | 0.86 | 0.68 | 0.91 |
| 16     | ada1\_A | 0.31 | 0.13 | -0.01 | -0.07 | 0.98 | 1.0 | 1.1 | 1.2 |
| 17     | ada1\_A | 0.14 | 0.13 | 0.11 | 0    |      |      |      |      |

Mean EDI values were determined from the EDIs calculated from 3 or more biological replicates, as shown for selected mutants in Fig. S4A, with all SEMs <20% of the mean values with 3 exceptions: gen5\_D at HIS4 (SEM/mean=0.68), gen5\_D at CPA2 (SEM/mean=0.55), and gen5\_D ydj1\_A at ARG1 (SEM/mean=0.24). EDI ranges are color-coded as indicated. Mean Gcn4 occupancies normalized to those measured in WT cells were determined from replicate experiments as described in Figs. 1C & E, with all SEMs <20% of the mean values shown. N/A, not applicable. H3 and Gcn4 ChIP analyses were not conducted for the spt16-169 mutant because it harbors a mutant HIS4 promoter.
**Supplemental Figure S4.** Summary of conventional ChIP analysis of H3 at four canonical Gcn4 target genes in different mutants. (A) Mean EDI values (± SD) were calculated as in Fig. S3A from 3 or more biological replicates for each gene in 8 different mutants. Asterisks indicating significant differences in Student’s t-tests (p<0.05) are color-coded to indicate the strains being compared; thus, the cyan asterisk above the ARG1 bar for gcn5Δydl1Δ indicates a mean EDI significantly different than that indicated by the ARG1 bar for the ydl1Δ strain (labeled cyan). (B) Measurements of Gcn4 occupancies in the panel of mutants analyzed in (A) (plus the snf2Δydl1Δ/vector strain presented for comparison) with mean (±SD) values calculated from 3 biological replicates.
**Supplemental Figure S5.** Efficient doxycycline-mediated repression of Snf2 expressed from $P_{TEA-SNF2}$ in different mutants. Strains of the indicated genotypes cultured in SC medium were treated (+) or left untreated (-) with 10µg/mL doxycycline for 8h and subjected to Western analysis with antibodies against Snf2 or Gcd6 (analyzed as loading control).
Supplemental Figure S6. Summary of conventional ChIP analysis of H3 and Gcn4 occupancies at four canonical Gcn4 target genes in different mutants. (A) Summary of EDI values and Gcn4 occupancies for the four genes in the indicated mutants, calculated and displayed as in Fig. S3A. (B) Measurements of Gcn4 occupancies for mutants analyzed in (A) with mean (± SD) values calculated from 3 biological replicates.
Supplemental Figure S7. H3 or Rpb3 densities in biological replicate experiments are highly correlated. Scatterplots are profiles of Rpb3 reads averaged over the CDS for each gene (A-B) or H3 reads in the 450 bp spanning nucleotides -389 to +61 (relative to the ATG codon) for each gene (C-D) calculated from ChIP-seq data in replicate experiments designated -1, -2, or -3, with the Pearson’s correlations coefficient indicated. For Rpb3, reads were obtained from sequenced fragments of 50-150 bp for 5783 genes, with 6 genes (A) and 1 gene (B) excluded as their extremely large read numbers would compress most of the data points into a small sector of the plots. H3 reads were obtained from sequenced fragments of 50-300 bp for 5783 genes, with 5 genes excluded from both plots (C-D) due to their extremely large read numbers.
Supplemental Figure S8. SM-induction evokes high-level Rpb3 in CDS and depletion of H3 in promoters and CDS at four canonical Gcn4 target genes. Reads at each coordinate base of Rpb3 (upper panels) and H3 (lower panels) for the indicated chromosomal regions containing ARG1 (A), ARG4 (B), HIS4 (C), and CPA2 (D), determined from the average of ChIP-seq data from 3 biological replicates. Note different scales on the x-axis chosen to display the entire CDS for Rpb3 (upper plots) or zoom in on promoters for H3 occupancies (lower plots). TSS and TTS positions were taken from (Xu et al. 2009). Violet-blue-violet bars in H3 profiles represent annotated -1, NDR and +1 regions as determined in Fig. S10B.
**Supplemental Figure S9.** Identification of genes exhibiting increased Rpb3 occupancies in CDS on SM-induction. Density plot of average Rpb3 occupancies per bp in CDS calculated from the average of ChIP-seq data from 3 biological replicates for SM-induced (Rpb3\textsubscript{I}) or uninduced (Rpb3\textsubscript{U}) WT cells. Individual genes are marked by blue filled circles; the coloring indicates increased density of genes (red is maximum). Dashed lines indicate differences of ≥2-fold. The four Gcn4 targets analyzed by conventional ChIP, ARG1, ARG4, HIS4, and CPA2, were among the genes displaying the largest Rpb3 induction ratios and induced levels of Rpb3 occupancy.
Supplemental Figure S10. Nuc-seq analysis of WT cells identifies consensus positions of the -1 nucleosome, NDR, and +1 nucleosomes and locations of these regions at each gene. Paired-end sequencing of mononucleosome core particles prepared by MNase digestion of nuclei was conducted for SM-induced or uninduced WT cells, averaging results from 2 biological replicates for each condition. (A) TSS plots for all genes showing consensus locations of -1, NDR, and +1 regions. (B) Nucleosome densities for all genes sorted by NDR length and aligned relative to the NDR centers (dotted line), with each row giving results for a single gene. Pink and cyan zones demarcate 147bp surrounding the dyads of the -1 and +1 nucleosomes, respectively, and the intervening NDRs are in lime green.
Supplemental Figure S11. Promoter depletion of H3 in SM-induced cells is correlated with changes in Rpb3 occupancies for 211 induced genes and for all genes. (A) The ratio of relative H3 occupancies per bp in the [-1,NDR,+1] windows defined in Fig. S10B in WT_U versus WT_I cells plotted against the ratio of Rpb3 occupancies per bp in CDS in WT_I versus WT_U cells, for the set of 211 SM-induced genes (defined in RESULTS), calculated from the average of ChIP-seq data from 3 biological replicates. Pearson’s (r) and Spearman’s (ρ) correlation coefficients are indicated. (B) Same as (A) except for all genes.
Supplemental Figure S12. Elimination of Snf2, Ydj1, or Gcn5 elevates H3 occupancies at highly induced genes. Noted box-plots of relative H3 occupancies per bp in the [-1,NDR,+1] window under inducing conditions for the 204 highly induced genes.
Supplemental Figure S13. Continued on next page.
Supplemental Figure S13. Continued on next page.
Supplemental Figure S13. Effects of snf2\(
\Delta\)
 and ydj1\(\Delta\)
 single or double mutations on promoter H3 occupancies of 70 SM-induced exemplar genes. (A-E) Mean (± SD) relative H3 occupancies per bp in the \([-1,NDR,+1]\) regions determined by ChIP-seq analysis for each of three biological replicates for WT under inducing and uninducing conditions, and mutant strains under SM-inducing conditions. Asterisks indicate significant differences in Student’s \(t\)-tests: *, \(p<0.05\); **, \(p<0.01\). Genes are listed in descending order of H3 eviction in WT cells, quantified as WT\(U\)/WT\(I\).
Supplemental Figure S14. Continued on next page.
Supplemental Figure S14. Continued on next page.
**Supplemental Figure S14.** Effects of *gcn5Δ* and *ydl1Δ* single or double mutations on promoter H3 occupancies of 70 SM-induced exemplar genes. (A-E) Mean (± SD) relative H3 occupancies per bp in the [-1,NDR,+1] regions determined by ChIP-seq analysis for each of three biological replicates for WT under both inducing and uninhibiting conditions, and mutant strains under SM-inducing conditions. Asterisks indicate significant differences in Student’s t-tests: *, p<0.05; **, p<0.01. Genes are listed in descending order of H3 eviction in WT cells, quantified as WT_U/WT_I. Results for WT and *ydl1Δ* are re-plotted from Fig. S13 for comparison.
Supplemental Figure S15. Continued on next page.
Supplemental Figure S15. Effects of *gcn5Δ, P_{TET-SNF2}, ydj1Δ, P_{TET-SNF2} gcn5Δ, and P_{TET-SNF2} gcn5Δ ydj1Δ/*cGCN4 single, double or triple mutations on promoter H3 occupancies of 70 SM-induced exemplar genes. (A-E) Mean (± SD) relative H3 occupancies per bp in the [-1,NDR,+1] regions determined by ChIP-seq analysis for each of three biological replicates for WT under both inducing and uninducing conditions, and mutant strains under SM-inducing conditions. Asterisks indicate significant differences in Student’s *t*-tests: *, *p*<0.05; **, *p*<0.01. Genes are listed in descending order of H3 eviction in WT cells, quantified as WT_U/WT_I. Results for WT, *gcn5Δ* and *ydj1Δ* are re-plotted from Fig. S14 for comparison.
Supplemental Figure S16. Caption on next page.
Supplemental Figure S16. Snf2, Gcn5, and Ydj1 cooperate in promoter H3 eviction at many of the 70 exemplar induced genes. (A-B) Cluster analysis of changes in relative H3 occupancies per bp in the [-1,NDR,+1] regions for the 70 exemplar genes in single, double and triple mutants. Relative H3 densities for each gene in the appropriate mutants shown in Fig. S15 were averaged and z-scores calculated for the results for each mutant. Hierarchical clustering analysis of the average relative H3 occupancies in the [-1,NDR,+1] regions was performed as described in Fig. 4A-B. Boxed genes display additive effects of mutations; those with bullets show strong eviction defects only in double mutants.
**Supplemental Figure S17.** Distribution of histone eviction defect indices for the 70 exemplar SM-induced genes in single mutants lacking Snf2, Ydj1, or Gcn5. The proportions of the 70 exemplar genes exhibiting EDI values falling into different bins is plotted for each mutant using the mean EDI values from Supplementary file S5, sheet 2. EDI was calculated as the increase in H3 occupancy in the mutant normalized to the difference in occupancy between uninduced and induced WT cells.
### Supplemental Figure S18.

Continued on next page.
Supplemental Figure S18. Summary of effects of single, double or triple mutations that eliminate or deplete Snf2, Ydj1, and Gcn5 on promoter H3 occupancies of 70 SM-induced exemplar genes. Compilation of p-values from Student’s t-tests conducted on results shown in Figs. S13-S15. Asterisks and muted colors indicate gene/mutant combinations where H3 occupancies are reduced rather than elevated in the mutant, indicating defects in nucleosome replacement versus eviction. Dependence on Snf2 for nucleosome eviction is summarized for each gene in column 8 (“Total Snf2-dep”) using colors indicated at the bottom signifying p-value ranges for the comparison yielding the most significant difference between strains containing or lacking SNF2, tabulated in the preceding 5 columns. Similar procedures were followed to tabulate dependence on Ydj1 (columns 9-13) or Gcn5 (columns 14-17) for nucleosome eviction. Totals for each column are listed at the bottom. Results in columns 8, 13, and 17 for p-values <0.05 (yellow/lime/blue), summarizing dependence on each co-factor, were used to construct the Venn diagram in Fig. 4C.
**Supplemental Figure S19.** Reductions in Pol II occupancy are associated with increased H3 promoter occupancy for 1000 genes with the greatest eviction of promoter nucleosomes in WT induced cells. (A–C) Heat maps for the identically-ordered 1000 genes shown in Fig. 2D, constructed as described there, for the indicated mutants.
**Supplemental Figure S20.** Mean defects in promoter H3 eviction are associated with mean defects in Pol II induction for the group of 70 induced exemplar genes. (A) EDI/TDI values for each gene/mutant combination were calculated as in Fig. 6A-B, and the mean EDI and TDI values were calculated for the group of 70 genes and plotted (±95% CI) for each strain, yielding the indicated Pearson’s coefficient. (B) EDI and TDI values were calculated for each gene/mutant combination as in Figs. 6A-B and seventy correlation analyses were conducted for each of the 70 exemplar genes between the EDI and TDI values determined for the 8 different mutants analyzed in (A), and the Pearson’s coefficient ($r$) was tabulated for all 70 genes. The average $r$ value for the 70 genes is 0.51, with a standard deviation of 0.35.
**Supplemental Figure S21.** Defects in promoter H3 eviction are associated with defects in Pol II induction for SM-induced exemplar genes in mutants with single, double or triple mutations eliminating or depleting Snf2, Ydj1, and Gcn5. (A-F) Scatterplots of EDIs versus TDIs calculated for all 70 exemplar SM-induced genes as in Fig. 6C, for the indicated mutants. In one or more plots, data for RTS1, CIT2, YPS1, LYS20, LYS21, NRG1, or STE2 were omitted (but included in calculations of Pearson’s (r) and Spearman’s (ρ) correlation coefficients) because their large negative TDI values would confine the majority of data points to a small portion of the x-axis, as follows. (A) ydj1Δ: CIT2, RTS3, YPS1; (B) snf2Δ: RTS3; (C) gcn5Δ: LYS20, RTS3; (D) snf2∆ydj1Δ/leGCN4: RTS3, LYS21; (E) gcn5Δ ydj1Δ: RTS3, LYS20; (F) P_TET-SNF2gcn5Δ ydj1Δ/leGCN4: CIT2, TRS3. In (F), NRG1 and STE2 were also omitted owing to their large EDI values.
Supplemental Figure S22. Elimination of Ydj1, Snf2, Gcn5 increases H3 occupancies in the NDRs of most genes without substantially altering Pol II occupancies. (A-C) Heat maps for the identically-ordered 4744 genes shown in Fig. 7A, constructed as described in Fig. 5 for the indicated mutants.
Supplemental Figure S23. Elimination of Snf2, Ydj1, or Gcn5 elevates promoter H3 occupancies genome-wide but reduces Pol II occupancies only at highly expressed genes. (A) Notched box-plots of relative H3 occupancies per bp in the [-1,NDR,+1] window (left) and log2 values of Rpb3 occupancies per bp in CDS (right) under inducing conditions for the first decile (with highest Rpb3 occupancies) of the group of 4744 weakly or non-induced genes. (B) Same as (A) except for the 5th decile of Rpb3 occupancy in WT cells. (C) Same as (A) except for the tenth decile of the group of 4744 genes (with lowest Rpb3 occupancies). Averaged H3 and Rpb3 ChIP-seq data from 3 biological replicates was employed.
Supplemental Figure S24. Elimination of Snf2, Ydj1, or Gcn5 elevates H3 occupancies and reduces Pol II occupancies at the 70 exemplar induced genes. Notched box-plots of H3 occupancies in the [-1,NDR,+1] window (A) and log₂(Rpb3) occupancies (B) under inducing conditions for the 70 exemplar induced genes. Data in (A) was presented in Fig. 3D and is reproduced here for comparison to (B).
## Supplemental Table S1. Yeast strains used in this study

| Name        | Parent | Relevant genotype \(^1\) | Reference                        |
|-------------|--------|---------------------------|----------------------------------|
| BY4741\(^2\) |        | MATa his3Δ1 leu2Δ met15Δ ura3Δ | Research Genetics                |
| 249\(^2\)   | BY4741\(^2\) | gcn4Δ::kanMX4           | Research Genetics                |
| 1586\(^2\)  | BY4741\(^2\) | snf2Δ::kanMX4           | Research Genetics                |
| 3012\(^2\)  | BY4741\(^2\) | ydj1Δ::kanMX4           | Research Genetics                |
| 1742\(^2\)  | BY4741\(^2\) | gal11Δ::kanMX4          | Research Genetics                |
| 1038\(^2\)  | BY4741\(^2\) | ada1Δ::kanMX4           | Research Genetics                |
| 7285\(^2\)  | BY4741\(^2\) | gen5Δ::kanMX4           | Research Genetics                |
| 1310\(^2\)  | BY4741\(^2\) | asf1Δ::kanMX4           | Research Genetics                |
| 1490\(^2\)  | BY4741\(^2\) | rtt109Δ::kanMX4         | Research Genetics                |
| 5119\(^2\)  | BY4741\(^2\) | nap1Δ::kanMX4           | Research Genetics                |
| 5922\(^2\)  | BY4741\(^2\) | yta7Δ::kanMX4           | Research Genetics                |
| 711\(^2\)   | BY4741\(^2\) | hsc82Δ::kanMX4          | Research Genetics                |
| 1512\(^2\)  | BY4741\(^2\) | ssa2Δ::kanMX4           | Research Genetics                |
| 1703\(^2\)  | BY4741\(^2\) | htz1Δ::kanMX4           | Research Genetics                |
| HQY1595     | 7285   | gen5Δ::kanMX4 ydj1Δ::hphMX4 | This work                       |
| HQY1598     | 1586\(^2\) | snf2Δ::kanMX4 ydj1Δ::hphMX4 | This work                       |
| HQY1602     | 771    | hsc82Δ::kanMX4 hsp82Δ::hphMX4 pHQ2066/LEU2 hsp82-T1011 | This work                       |
| HQY1603     | 771    | hsc82Δ::kanMX4 hsp82Δ::hphMX4 pHQ2067/LEU2 hsp82-G170D | This work                       |
| HQY1607     | 1512   | ssa1Δ::hphMX4 ssa2Δ::kanMX4 | This work                       |
| HQY1610     | 1586   | snf2Δ::kanMX4 yta7Δ::hphMX4 | This work                       |
| HQY1611     | 7285   | gen5Δ::kanMX4 yta7Δ::hphMX4 | This work                       |
| HQY1612     | 1586   | snf2Δ::kanMX4 asf1Δ::hphMX4 | This work                       |
| HQY1623     | 7285   | HIS3*:P\(_{TET}\)-SNF2 gen5Δ::kanMX4 | This work                       |
| HQY1625     | 249\(^2\) | HIS3*:P\(_{TET}\)-SNF2 gcn4Δ::kanMX4 | This work                       |
| HQY1627     | BY4741\(^2\) | HIS3*:P\(_{TET}\)-SNF2 | This work                       |
| HQY1657     | HQY1595 | HIS3*:P\(_{TET}\)-SNF2 gen5Δ::kanMX4 ydj1Δ::hphMX4 | This work                       |
| BLY49       | MATa   | sht1-3ts his3-Δ200 ura3-52 lys2-801 ade2-101 | Chai et al. \(^*\) 2005         |
| L577        | MATα   | spt16-197 his4Δ912 lys2Δ128 ura3-52 | Malone et al. \(^*\) 1991        |

\(^1\) HIS3* designates the HIS3 allele from S. kluyveri.

\(^2\) Strains are purchased from Research Genetics.
## Supplemental Table S2. H3-PESCI studies

| Yeast strain (genotype) | Sample ID | No. of aligned paired reads | Pearson correlation<sup>2</sup> | Exp. 1 | Exp. 2 |
|-------------------------|-----------|----------------------------|-------------------------------|--------|--------|
| BY4741 (WT, uninduced)  | AGH0220-1 (Exp. 1) | 22,933,214                 |                               |        |        |
|                         | AGH0220-2 (Exp. 2) | 20,223,628                 | 0.958                         |        |        |
|                         | AGH0220-3 (Exp. 3) | 21,078,138                 | 0.958                         |        |        |
| BY4741 (WT, induced)    | AGH0220-4 (Exp. 1) | 27,288,850                 |                               |        |        |
|                         | AGH0220-5 (Exp. 2) | 22,101,846                 | 0.929                         |        |        |
|                         | AGH0220-6 (Exp. 3) | 22,386,501                 | 0.922                         |        | 0.956  |
| 1586 (snf2Δ)            | AGH0406-10-1 (Exp. 1) | 23,691,663               |                               |        |        |
|                         | AGH0406-10-2 (Exp. 2) | 30,728,286               | 0.912                         |        |        |
|                         | AGH25-1 (Exp. 3)     | 27,312,607                 | 0.806                         |        | 0.791  |
| 3012 (ydj1Δ)            | AGH0406-11-1 (Exp. 1) | 22,351,818               |                               |        |        |
|                         | AGH0406-11-2 (Exp. 2) | 23,409,609               | 0.928                         |        |        |
|                         | AGH25-2 (Exp. 3)     | 26,326,214                 | 0.858                         |        | 0.864  |
| 7285 (gcn5Δ)            | AGH08-5 (Exp. 1)     | 13,992,780                 |                               |        |        |
|                         | AGH09-5 (Exp. 2)     | 10,618,715                 | 0.817                         |        |        |
|                         | AGH25-3 (Exp. 3)     | 26,846,307                 | 0.791                         |        | 0.763  |
| HQY1598 (snf2Δydj1Δ/leGCN4) | AGH13-1 (Exp. 1) | 30,295,240                 |                               |        |        |
|                         | AGH13-2 (Exp. 2)     | 26,707,317                 | 0.921                         |        |        |
|                         | AGH13-3 (Exp. 3)     | 23,557,188                 | 0.878                         |        | 0.854  |
| HQY1595 (gcn5Δydj1Δ)    | AGH08-6 (Exp. 1)     | 14,521,858                 |                               |        |        |
|                         | AGH09-6 (Exp. 2)     | 12,233,963                 | 0.864                         |        |        |
|                         | AGH25-4 (Exp. 3)     | 25,370,469                 | 0.835                         |        | 0.821  |
| HQY1627 (P<sub>TET</sub>-SNF2) | AGH08-4 (Exp. 1) | 14,492,452                 |                               |        |        |
|                         | AGH09-4 (Exp. 2)     | 12,155,964                 | 0.840                         |        |        |
|                         | AGH25-8 (Exp. 3)     | 24,176,184                 | 0.822                         |        | 0.859  |
| HQY1623 (P<sub>TET</sub>-SNF2gcn5Δ) | AGH08-8 (Exp. 1) | 14,462,524                 |                               |        |        |
|                         | AGH09-8 (Exp. 2)     | 9,372,348                  | 0.812                         |        |        |
|                         | AGH25-10 (Exp. 3)    | 22,921,217                 | 0.719                         |        | 0.689  |
| HQY1657 (P<sub>TET</sub>-SNF2gcn5Δydj1Δ/leGCN4) | AGH08-12 (Exp. 1) | 12,230,198                 |                               |        |        |
|                         | AGH09-12 (Exp. 2)    | 11,067,423                 | 0.746                         |        |        |
|                         | AGH27-5 (Exp. 3)     | 25,380,703                 | 0.491                         |        | 0.505  |

1 Using Bowtie 2 to align reads to the SacCer2 version of the genome sequence.
2 Pearson correlations between experiments for the H3 average occupancy in the combined [-1,NDR,+1] regions of 5403 annotated genes.
**Supplemental Table S3.** Rpb3-PESCI studies

| Yeast strain (genotype) | Sample ID | No. of aligned paired reads | Pearson correlation
|-------------------------|-----------|----------------------------|---------------------|
|                         |           | Exp. 1                     | Exp. 2              |
| **BY4741 (WT, uninduced)** |           |                            |                     |
| AGH03-1 (Exp. 1)       |           | 27,714,859                 | 0.997               |
| AGH03-2 (Exp. 2)       |           | 31,174,598                 | 0.995               |
| AGH03-3 (Exp. 3)       |           | 32,460,758                 | 0.995               |
| **BY4741 (WT, induced)** |           |                            |                     |
| AGH03-4 (Exp. 1)       |           | 23,003,834                 | 0.995               |
| AGH03-5 (Exp. 2)       |           | 29,921,812                 | 0.995               |
| AGH03-6 (Exp. 3)       |           | 26,823,064                 | 0.994               |
| **249 (gcn4Δ)**        |           |                            |                     |
| AGH12-1 (Exp. 1)       |           | 16,573,573                 | 0.998               |
| AGH12-2 (Exp. 2)       |           | 15,234,053                 | 0.950               |
| AGH12-3 (Exp. 3)       |           | 12,149,580                 | 0.952               |
| **1586 (snf2Δ)**       |           |                            |                     |
| AGH12-4 (Exp. 1)       |           | 18,787,456                 | 0.997               |
| AGH12-5 (Exp. 2)       |           | 17,401,885                 | 0.952               |
| AGH12-6 (Exp. 3)       |           | 23,806,884                 | 0.957               |
| **3012 (ydj1Δ)**       |           |                            |                     |
| AGH12-7 (Exp. 1)       |           | 20,262,851                 | 0.997               |
| AGH12-8 (Exp. 2)       |           | 20,954,044                 | 0.936               |
| AGH12-9 (Exp. 3)       |           | 25,344,969                 | 0.931               |
| **7285 (gcn5Δ)**       |           |                            |                     |
| AGH12-10 (Exp. 1)      |           | 23,998,691                 | 0.991               |
| AGH12-11 (Exp. 2)      |           | 23,412,718                 | 0.864               |
| AGH12-12 (Exp. 3)      |           | 22,255,960                 | 0.866               |
| **HQY1598 (snf2Δydj1Δ/leGCN4)** | AGH13-4 (Exp. 1) | 23,925,582                 | 0.975               |
|                         | AGH13-5 (Exp. 2) | 26,980,682                 | 0.970               |
|                         | AGH13-6 (Exp. 3) | 24,924,859                 | 0.975               |
| **HQY1595 (gcn5Δydj1Δ)** | AGH14-8 (Exp. 1) | 20,744,649                 | 0.977               |
|                         | AGH14-9 (Exp. 2) | 24,995,164                 | 0.926               |
|                         | AGH14-10 (Exp. 3) | 20,314,069                 | 0.927               |
| **HQY1627 (P_{TET-SNF2})** | AGH15-8 (Exp. 1) | 18,963,524                 | 0.980               |
|                         | AGH15-9 (Exp. 2) | 25,390,805                 | 0.950               |
|                         | AGH15-10 (Exp. 3) | 23,667,405                 | 0.959               |
| **HQY1623 (P_{TET-SNF2gcn5Δ})** | AGH16-8 (Exp. 1) | 20,681,102                 | 0.978               |
|                         | AGH16-9 (Exp. 2) | 25,091,761                 | 0.926               |
|                         | AGH16-10 (Exp. 3) | 21,059,642                 | 0.943               |
| **HQY1657 (P_{TET-SNF2gcn5Δydj1Δ/leGCN4})** | AGH18-12 (Exp. 1) | 24,057,383                 | 0.970               |
|                         | AGH18-13 (Exp. 2) | 20,092,762                 | 0.840               |
|                         | AGH18-14 (Exp. 3) | 28,621,240                 | 0.855               |

1. Using Bowtie 2 to align reads to the SacCer2 version of the genome sequence.
2. Pearson correlations between experiments for the Rpb3 occupancies averaged over the CDS for all 5783 genes with the CDS longer than 170 bp.
### Supplemental Table S4. Nuc-seq studies

| Yeast strain (genotype) | Sample ID          | No. of aligned paired reads¹ | Pearson correlation² r |
|------------------------|--------------------|------------------------------|------------------------|
| BY4741 (WT, uninduced) | AGH01-1 (Exp. 1)   | 8,699,065                    | 0.983                  |
|                        | AGH01-2 (Exp. 2)   | 8,060,225                    |                        |
| BY4741 (WT, induced)   | AGH01-3 (Exp. 1)   | 7,968,815                    | 0.987                  |
|                        | AGH01-4 (Exp. 2)   | 10,220,772                   |                        |

¹ Using Bowtie 2 to align reads to the SacCer2 version of the genome sequence.

² Pearson correlations between experiments for the nucleosome occupancies in the 450 bp spanning nucleotides -389 to +61 (relative to the ATG codon) of each gene for all 5783 genes with the CDS longer than 170 bp.
Additional Supplemental Files

Supplemental File S1

ChIP-seq analysis of H3 and Rpb3 occupancies in uninduced and SM-induced WT cells. Sequence reads from three biological replicates were combined and H3 occupancies at each bp were normalized to the average occupancy per bp on the respective chromosome, to yield relative H3 occupancies. Rpb3 occupancies were normalized to correct for differences in total read depth in different experiments. The relative H3 occupancies per bp in the [-1,NDR,+1] intervals for each gene, identified as in Fig. S10B, are listed for all 5332 genes for which the [-1,NDR,+1] region could be defined (sheet 4), 4744 genes exhibiting <1.5-fold induction of Rpb3 on SM induction (sheet 3), 211 genes selected from a group of 223 exhibiting SM-induction of Rpb3 by ≥2-fold (sheet 2), and 70 exemplar genes from the group of 211 genes displaying an H3_U/H3_I ratio of ≥1.415 (sheet 1). The group of 223 SM-induced genes does not include all genes exhibiting Rpb3_I/Rpb3_U ≥2-fold as it excludes certain genes that appear to exhibit read-through transcription from adjacent, highly induced genes that does not extend across the entire CDS, or that have unusually low reads under one or both conditions. Of the 223 genes that satisfy these criteria, the [-1,NDR,+1] region could not be defined for 12 genes, yielding the group of 211 genes listed in sheet 2. Note that 14 genes in the group of 211 belong to pairs of divergently oriented genes that share a common NDR, and that one of the two genes in the pair (highlighted in yellow) was excluded in certain analyses of SM-induced genes described in RESULTS. In addition, of the 73 genes from the group of 211 genes that display an H3_U/H3_I ratio of ≥1.415 (sheet 2), 6 genes belong to pairs of divergently oriented genes that share a common NDR, such that one of the two genes in the pair (highlighted in yellow) was excluded to produce the 70 exemplar genes (sheet 1). Sheets 1-3 list the relative H3 occupancies per bp in the [-1, NDR, +1] zones under uninducing (H3-U) or inducing (H3-I) conditions, and the Rpb3 occupancies per bp averaged over the CDS under the same two conditions (Rpb3-U, Rpb3-I) for each gene. Sheet 5 summarizes results of microarray analysis of mRNA expression in WT cells treated or untreated with SM for 2h, with 1370 genes exhibiting an SM-induced increase in mRNA expression by ≥1.5 and 1298 genes with a similar fold reduction in expression. Four independent RNA preparations were made from untreated or SM-treated WT cells and used for 4 microarray hybridizations, including two dye swaps, to generate mean values calculated from the 4 replicate experiments (columns F and L) (Saint et al. 2014).

Supplemental File S2

Average H3 occupancies of -1, NDR, and +1 zones in WT and mutant cells. Relative H3 occupancies were calculated from the data combined from 3 biological replicates as described in File S1. Sheets 2-17: Relative H3 occupancy per bp (Avg. H3 Occ.) was calculated for each gene for the [-1,NDR,+1] combined zones (sheets *_3zones), or individual -1 nucleosome regions (sheets *_-1), NDRs (sheets *_NDR), or +1 nucleosome regions (sheets *_+1), for WT uninduced or induced cells and the indicated 8 mutants cultured under inducing conditions in the groups of 70, 204, 4744 or all (5403) genes. Sheet 1: Mean (+ 95% CI) Avg. H3 Occ. values were calculated for the indicated gene sets (70, 204, 4744 or all 5403) in the indicated strains, along with the fold-changes in mean values between mutant and induced WT cells (rel to WT-I). The mean values (+ 95% CI) calculated for the individual -1, NDR, and +1 zones for the 70 exemplar genes are plotted in Fig. 3E.

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Supplemental File S3

Summary of effects of single or double mutations that eliminate Snf2, Ydj1, or Gcn5 on promoter H3 occupancies of 70 induced exemplar genes. (Sheet 1) Compilation of p-values from Student’s t-tests conducted on results shown in Fig. S13. Asterisks and muted colors indicate gene/mutant combinations where H3 occupancies are reduced in the mutant, indicating defects in nucleosome replacement versus eviction. Dependence on Snf2 or Ydj1 for nucleosome eviction is summarized for each gene in columns 8 or 9 using the color code at the bottom signifying p-value ranges for the comparison yielding the most significant difference between strains containing or lacking SNF2 or YDJ1, which are tabulated in the preceding 5 columns. Totals for each column are listed at the bottom. (Sheet 2) Same as sheet 1 but for mutants lacking GCN5 and YDJ1 and compiled from t-tests conducted on results shown in Fig. S14.

Supplemental File S4

ChIP-seq analysis of Rpb3 occupancies averaged over the CDS in SM-induced WT and mutant cells. Rpb3 occupancy data from 3 biological replicates, from reads of 50-300 bp sequenced fragments, were normalized for differences in total read number, averaged, and normalized to CDS length, to yield the mean values listed here for all 5783 genes (sheet 4), 4744 genes exhibiting <1.5-fold induction of Rpb3 on SM induction (sheet 3), 211 genes selected from a group of 223 exhibiting SM-induction of Rpb3 by ≥2-fold (sheet 2), and 70 exemplar genes from the group of 211 genes displaying an H3_U/H3_I ratio of ≥1.415 (sheet 1).

Supplemental File S5

EDI and TDI values for 70 exemplar genes. (Sheet 1) Mean (± SEM) EDI and TDI values calculated for each gene in the P_{TET}-SNF2 gcn5 Δ double mutant by averaging EDI and TDI values calculated from three biological replicates, for the analysis in Fig. 6C. (Sheet 2) EDI and TDI values were calculated for each gene/mutant combination using the average of H3 or Rpb3 ChIP-seq data from three biological replicates for the analyses in Figs. 6A-B, S17, S20A-B, and S21A-F.

Supplemental File S6

Perl and MATLAB scripts used in this study. Archive File S6.zip contains two folders, Perl_scripts and MATLAB_scripts, with the Perl and MATLAB scripts used in this study. Perl_scripts folder contains the scripts necessary to size-select the raw paired-end reads according to the insert sizes, and to compute the number of mapped reads over promoters, coding regions, and downstream regions (see demo.docx for a short tutorial with the necessary steps to process the *.sam files containing the aligned paired-end reads that were generated with Bowtie 2). MATLAB_scripts folder contains the following scripts: Detect_NDRs_and_flanking_nucleosomes detects the locations of the nucleosome depleted regions and the flanking nucleosomes (+1/-1) corresponding to each gene promoter; Compute_H3_averages_TSS generates Excel tables with the genome-wide average H3 occupancy in the window [TSS-1000, TSS +1000]; Compute_H3_averages_3_zones generates Excel tables with the average H3 occupancy in the three zones (-1 nucleosome, NDR, +1 nucleosome) for every gene.
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