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THE TITLE IS WASDA

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ABSTRACT

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Chapter 1

Introduction

1.1 A brief introduction to transcription

In eukaryotic cells, transcription of protein-coding genes is carried out by the protein complex RNA polymerase II (Pol II), and broadly occurs in three sequential stages of transcription initiation, elongation, and termination (Shandilya and Roberts, 2012). During each of these stages, the Pol II complex is associated with distinct sets of factors which modulate the activity of Pol II and carry out co-transcriptional processes such as RNA capping, RNA splicing, histone modification, RNA cleavage, and RNA polyadenylation. Given how fundamental transcription is to gene expression, it is unsurprising that every stage of transcription is highly regulated.

To get a rough idea of just how tightly transcription is regulated, it is useful to consider a back-of-the-envelope calculation of the specificity of transcription initiation in the human genome. That is, what proportion of the human genome at which transcription could initiate does transcription initiation actually occur?

The number of positions at which transcription could theoretically initiate is simply the size of the genome: The human genome is approximately three billion base pairs in length (BNID 111378, Weber et al. (2009)), and since each base pair can be transcribed from each of its two strands, there are 6×10^9 available positions.

The number of positions at which transcription *does* initiate can be estimated from the number of genes transcribed by Pol II and the number of positions that Pol II initiates from for each gene. At last count, the human genome contains about twenty thousand protein-coding genes (Consortium et al., 2012). To be conservative in our estimate with regards to specificity, we will assume that all twenty thousand genes are expressed. We also know that protein-coding genes are only a subset of the genes transcribed by Pol II: Pol II also transcribes multiple classes of non-coding genes, including enhancers and long non-coding RNAs (Kaikkonen and Adelman, 2018). Compared to protein-coding genes, the number of non-coding genes is less certain. If we assume that there are five non-coding genes for each coding gene, this brings our estimate of the number of genes transcribed by Pol II to 1.2×10^5 genes.

As you will see from yeast transcription start site data in later chapters, transcription initiation for a single gene generally occurs at multiple nucleotides, generating multiple major transcript isoforms per gene. Assuming that there are five major transcription start sites (TSSs) per gene, the proportion of the human genome at which transcription initiation occurs is

$$\frac{(1.2 \times 10^5 \text{ genes}) \left(5 \frac{\text{TSSs}}{\text{gene}} \right)}{(6 \times 10^9 \text{ possible TSSs})} = 1 \times 10^{-4}.$$

Our rough estimate says that, when presented with ten thousand positions to choose from, RNA polymerase starts transcription from only one!¹

Many factors are known to contribute to this remarkable specificity. Most notably, transcription initiation requires the presence of specific DNA sequence motifs, which

¹A similar conclusion is reached by examining ENCODE CAGE-seq data: At the time of writing, ENCODE reports roughly 150,000 TSS peaks across 30 cell types/cell lines. Assuming the signal is concentrated at 5 nucleotides per peak, then $\frac{(1.5 \times 10^5 \text{ peaks})(5 \frac{\text{nt}}{\text{peak}})}{6 \times 10^9 \text{ nt}} = \frac{1}{8000}$.

increase the probability of Pol II binding to DNA together with necessary initiation factors (Haberle and Stark, 2018). That factors known to associate with Pol II during transcription initiation control transcription initiation is unsurprising. A less obvious fact is that some transcription *elongation* factors, including histone chaperones and histone modification enzymes, also play a role in restricting where transcription initiation is allowed to occur (Cheung et al., 2008; Hennig and Fischer, 2013; Kaplan et al., 2003). Evidence suggests that these elongation factors are likely required to maintain normal chromatin structure over transcribed regions, and that the disruption of normal chromatin structure allows Pol II to initiate transcription in regions which are normally inaccessible (). Chapters 2 and 3 of this dissertation describe our studies of **Spt6** and **Spt5**, two of the transcription elongation factors involved in this process. One phenotype observed when these factors are disrupted is **intragenic transcription**, transcription appearing to arise from within protein-coding sequences. In chapter 4, I describe our efforts to understand how intragenic transcription might play a role in the cellular response to various stress conditions. The remainder of this introduction provides a brief overview of the considerations taken into account in order to make the data analyses behind this dissertation as transparent and reproducible as possible.

1.2 Reproducible data analysis for genomics

My role in the projects in this dissertation is a mix of **data scientist** and **data engineer**: I build pipelines for processing (usually genomic) datasets, taking raw data through processing, statistical analysis, and data visualization. This mostly entails surveying available tools, selecting the tools most suitable for the task, and coding solutions to problems when existing tools are insufficient.

The analysis of complex datasets like those in genomics presents challenges to

achieving transparency and reproducibility when reporting methods and results. In building the data analysis pipelines behind the results of this dissertation, I have tried to meet these challenges by following best practices that would be standards for publication in an ideal world. All of my data analyses are open source (github.com/winston-lab), and are designed to be reproducible by others: For all publications, an self-contained archive is uploaded which includes everything needed to go from raw data to the figures and results of the publication (e.g. <https://doi.org/10.5281/zenodo.1409826>). This level of accessibility is greatly facilitated by building data analyses using Snakemake (Köster and Rahmann, 2012), one of several available frameworks for workflow management (Di Tommaso et al., 2017; Voss et al., 2017). Snakemake's scalable execution and its ability to specify dependencies in virtual environments allow workflows to truly be reproducible: data analyses can be re-run on personal computers, computing clusters, or cloud environments, and the exact versions of the software used when initially running the data analysis will automagically be deployed.

Open sharing of data and code like this is essential to the scientific process. When analysis pipelines routinely consist of tens of steps with tens of parameters each, seeing the data and code is the only way for those interested to know exactly how the data were handled. Altogether, this allows for more informed evaluation of results from the literature, as well as the possibility of finding and correcting errors in analysis.

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Chapter 2

Genomics of transcription elongation factor Spt6

2.1 Collaborators

Steve Doris optimized TSS-seq and ChIP-nexus protocols
generated TSS-seq and ChIP-nexus libraries

Olga Viktorovskaya generated MNase-seq libraries

Magdalena Murawska generated NET-seq libraries

Dan Spatt various experiments for publication

2.2 Introduction to Spt6 and intragenic transcription

The conserved transcription elongation factor Spt6 interacts directly with RNA polymerase II (Close et al., 2011; Diebold et al., 2010b; Liu et al., 2011; Sdano et al., 2017; Sun et al., 2010; Yoh et al., 2007), histones (Bortvin and Winston, 1996; McCullough et al., 2015), and another elongation factor called Spn1/lws1 (Diebold et al., 2010a; Li et al., 2018; McDonald et al., 2010). The classification of Spt6 as a transcription elongation factor is based on its association with elongating Pol II (Andrulis et al., 2000; Ivanovska et al., 2011; Kaplan et al., 2000; Mayer et al., 2010), and its ability to enhance elongation both *in vitro* (Endoh et al., 2004) and *in vivo* (Ardehali et al., 2009), though Spt6 has also been shown to regulate initiation in a small number

of cases (Adkins and Tyler, 2006; Ivanovska et al., 2011). Evidence suggests that as Spt6 travels with elongating Pol II, it acts as a histone chaperone, reassembling nucleosomes after their displacement from DNA due to transcription (Duina, 2011). Consistent with its histone chaperone function, Spt6 influences chromatin structure (Bortvin and Winston, 1996; DeGennaro et al., 2013; Ivanovska et al., 2011; Jeronimo et al., 2015; Kaplan et al., 2003; Perales et al., 2013; van Bakel et al., 2013); Spt6 is also required for some histone modifications, including H3K36 methylation (Carrozza et al., 2005; Chu et al., 2006; Yoh et al., 2008; Youdell et al., 2008), and, in some organisms, H3K4 and H3K27 methylation (Begum et al., 2012; Chen et al., 2012; DeGennaro et al., 2013; Wang et al., 2017, 2013).

Studies in the yeasts *Saccharomyces cerevisiae* and *Schizosaccharomyces pombe* have previously examined the requirement for Spt6 in normal transcription (Cheung et al., 2008; DeGennaro et al., 2013; Kaplan et al., 2003; Pathak et al., 2018; Uwimana et al., 2017; van Bakel et al., 2013). As Spt6 is essential for viability in *S. cerevisiae*, many of these studies use the same temperature-

sensitive *spt6* mutant used in this project, ***spt6-1004***, which encodes an in-frame deletion of a helix-hairpin-helix domain within Spt6 (Kaplan et al., 2003). When *spt6-1004* cells are shifted from 30 °C to 37 °C for 80 minutes, bulk Spt6 protein levels are depleted to about 20% of wild-type levels (Figure 2.1). A notable phenotype of the *spt6-1004* mutant is the appearance of **intragenic transcripts**, transcripts which

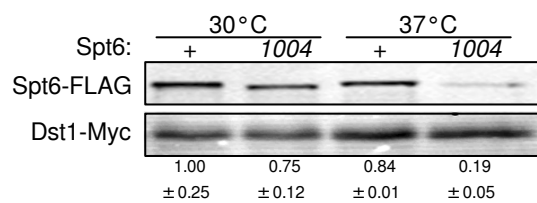


Figure 2.1: Western blot for Spt6 in wild-type and *spt6-1004* cells, at 30 °C and after 80 minutes at 37 °C. Spt6 and Dst1 from a spike-in were detected using α -FLAG and α -Myc antibodies, respectively. The mean \pm standard deviation of three blots are shown below each lane.

appear to arise from within protein-coding sequences, in both sense and antisense orientations relative to the coding gene (Figure 2.2) (Cheung et al., 2008; DeGennaro et al., 2013; Kaplan et al., 2003; Uwimana et al., 2017).

Previous genome-wide measurements

of transcript levels in *spt6-1004* relied on tiled microarrays (Cheung et al., 2008) and RNA sequencing (Uwimana et al.,

2017). Studying intragenic transcription is difficult with these methods, since the signal for an intragenic transcript in the same orientation as the gene it overlaps

is convoluted with the signal from the full-length ‘genic’ transcript (Figures 2.2, 2.3) (Cheung et al., 2008; Lickwar et al., 2009). Therefore, these methods can only discover intragenic transcripts which are highly expressed relative to the corresponding genic transcript, and are likely to find many false positives. Additionally, these methods are assays of steady-state RNA levels, which makes them unable to distinguish whether the intragenic transcripts observed in *spt6-1004* result from: A) new intragenic transcription initiation in the mutant, B) reduced decay of intragenic transcripts which are rapidly degraded in wild-type, or C) processing of full-length protein-coding RNAs.

To address these challenges to studying intragenic transcription, we applied two genomic assays to *spt6-1004*: transcription start-site sequencing (**TSS-seq**), and **ChIP-nexus of TFIIIB**, a component of the RNA polymerase II pre-initiation complex (PIC). TSS-seq sequences the 5' end of capped and polyadenylated RNAs (Arribere and Gilbert, 2013; Malabat et al., 2015), allowing separation of intragenic from genic

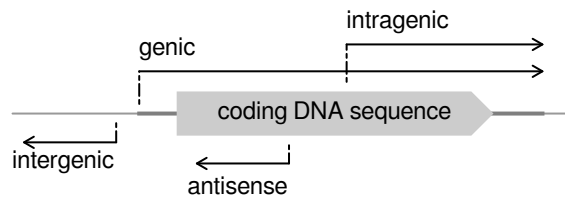
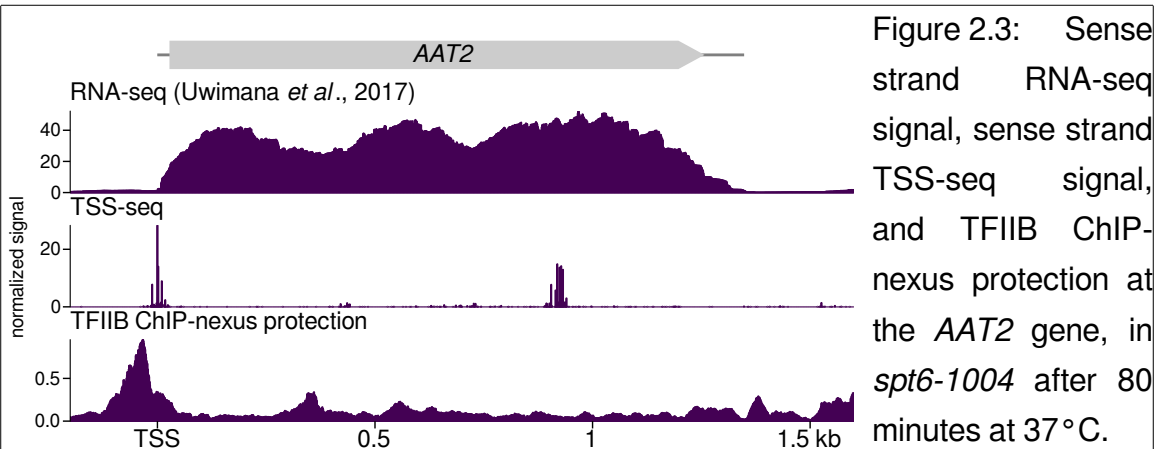


Figure 2.2: Diagram of transcript orientation with respect to coding DNA sequences, for the categories of transcripts referred to in this document.



RNA signals and identification of intragenic transcript starts with single-nucleotide resolution (Figure 2.3). ChIP-nexus is a high-resolution chromatin immunoprecipitation technique, in which the immunoprecipitated DNA is exonuclease digested up to the bases crosslinked with the protein of interest before sequencing (He et al., 2015). When applied to the PIC component TFIIB, ChIP-nexus reports where transcription initiation is occurring, thus allowing us to determine if intragenic transcripts in *spt6-1004* result from new transcription initiation.

2.3 TSS-seq and TFIIB ChIP-nexus results for *spt6-1004*

To study the relationship between Spt6 and transcription, TSS-seq and TFIIB ChIP-nexus libraries were prepared from wild-type and *spt6-1004* cells, both shifted from 30°C to 37°C for 80 minutes. In wild-type cells, TSS-seq and TFIIB ChIP-nexus recapitulate their expected distributions over the genome: Most TSS signal is restricted to annotated genic TSSs, while most TFIIB signal is localized just upstream of the TSS (Figures 2.4, 2.5). In *spt6-1004*, the signal for both assays infiltrates gene bodies, reflecting widespread intragenic expression of capped and polyadenylated transcripts, and suggesting that new transcription initiation contributes to the intra-

genic transcription phenotype. Notably, sense strand TSS-seq signal in *spt6-1004* tends to occur towards the 3' end of genes, while antisense strand TSS-seq signal tends to occur towards the 5' end of genes.

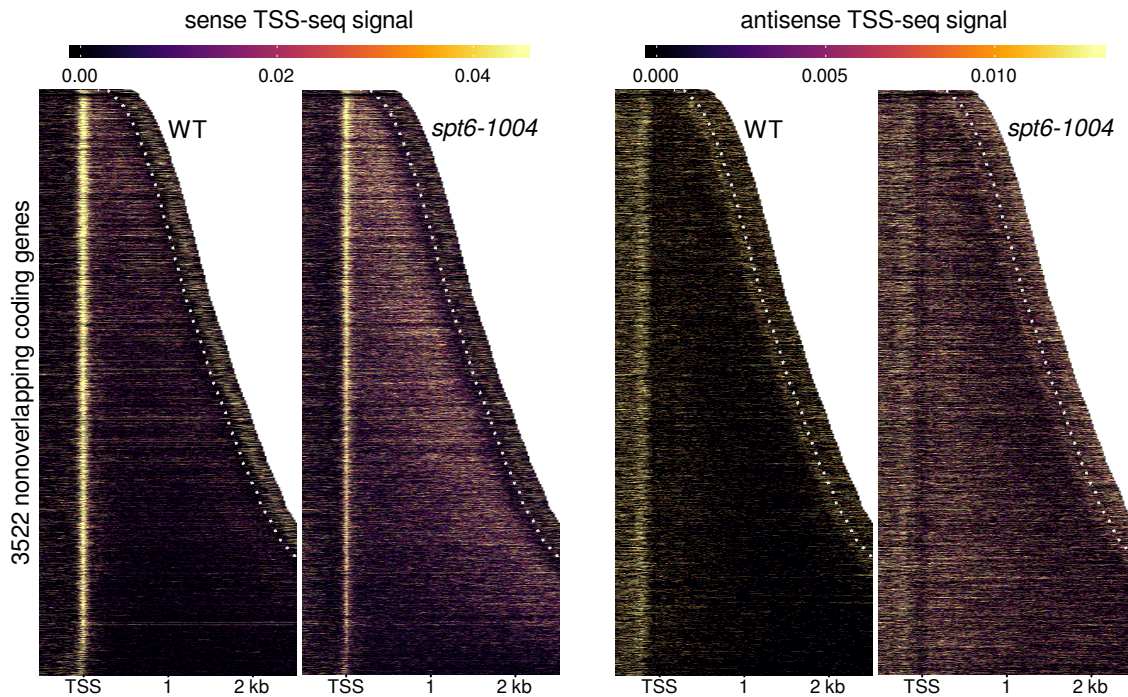


Figure 2.4: Heatmaps of sense and antisense TSS-seq signal from wild-type and *spt6-1004* cells, over 3522 non-overlapping genes aligned by wild-type genic TSS and sorted by annotated transcript length. Data are shown for each gene up to 300 nucleotides 3' of the cleavage and polyadenylation site (CPS), indicated by the white dotted line. Values are the mean of spike-in normalized coverage in non-overlapping 20 nucleotide bins, averaged over two replicates. Values above the 92nd percentile are set to the 92nd percentile for visualization.

The TSS-seq data were quantified by peak calling and differential expression analysis, and classified into genomic categories based on their position relative to coding genes. As suggested by the heatmap visualization (Figure 2.4), we detect significant induction of over 4000 intragenic and antisense TSSs in *spt6-1004* (Fig-

ure 2.6). Compared to previous studies identifying *spt6-1004* intragenic transcription by tiled microarray and RNA-seq (Cheung et al., 2008; Uwimana et al., 2017), we identify intragenic transcription at over 1000 additional genes (Figure 2.7), with the additional information of exact start sites for all identified TSSs.

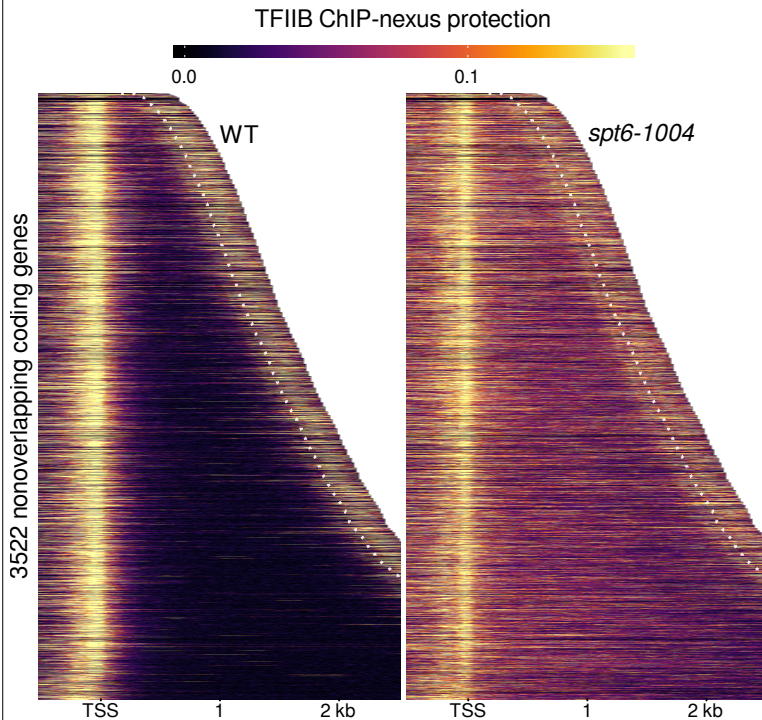


Figure 2.5: Heatmaps of TFIIB binding measured by ChIP-nexus, over the same regions shown in Figure 2.4. Values are the mean of library-size normalized coverage in non-overlapping 20 bp bins, averaged over two replicates. Values above the 85th percentile are set to the 85th percentile for visualization.

The TSS-seq data also revealed an unexpected downregulation of most genic TSSs: In this experiment, we detected a significant downregulation to levels below 67% of wild-type levels at 75% (3579/4792) of genic TSSs (Figure 2.6). As a result of intragenic/antisense induction and genic repression, expression levels in *spt6-1004* of all classes of transcripts become similar to one another (Figure 2.8).

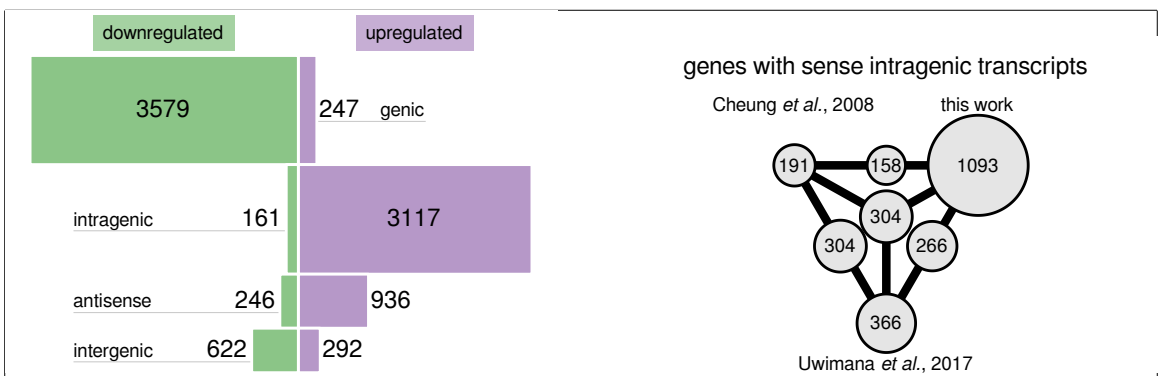


Figure 2.6: Bar plots of the number of TSS-seq peaks differentially expressed in *spt6-1004* versus wild-type, both after 80 minutes at 37°C. The height of each bar is proportional to the total number of peaks in the category, including those not found to be significantly differentially expressed.

The changes in transcript levels in *spt6-1004* observed by TSS-seq correspond with substantial differences in the pattern of TFIIB binding on the genome. While TFIIB in wild-type binds in discrete peaks within promoter regions, TFIIB in *spt6-1004* binds much more promiscuously, with many loci having TFIIB signal spread over broad regions of the genome (Figure 2.9). This difference in binding pattern makes peak calling ineffective for quantifying TFIIB signal in this

genes with sense intragenic transcripts

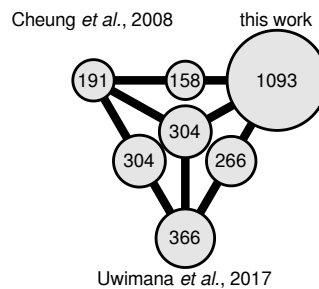


Figure 2.7: Set diagram of the number of genes reported to have *spt6-1004*-induced intragenic transcripts using tiled arrays (Cheung et al., 2008), RNA-seq (Uwimana et al., 2017), and TSS-seq (this work).

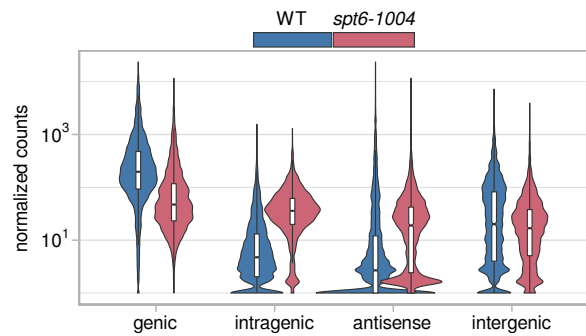


Figure 2.8: Violin plots of expression level distributions for genomic classes of TSS-seq peaks in wild-type and *spt6-1004*, both after 80 minutes at 37°C. Normalized counts are the mean of spike-in size factor normalized counts from two replicates.

case: ChIP-seq peak callers generally use different algorithms for calling ‘narrow’ peaks (e.g. for sequence-specific transcription factors) and ‘broad’ peaks (e.g. for histone modifications), meaning that a single algorithm is unable to call a unified set of peaks that is meaningful for differential binding analyses between wild-type and *spt6-1004*. Therefore, to see if changes in transcript levels in *spt6-1004* correspond to changes in transcription initiation, we compared the change in TSS-seq signal at TSS-seq peaks in *spt6-1004* to the change in TFIIB ChIP-nexus signal in the window extending 200 bp upstream of the TSS-seq peak. Changes in TSS-seq signal in *spt6-1004* are associated with a change in TFIIB signal of the same sign at over 82% of TSSs of any genomic class (Figure 2.10), indicating that the increase in intragenic transcript levels and decrease in genic transcript levels observed in *spt6-1004* are in large part explained by changes in transcription initiation.

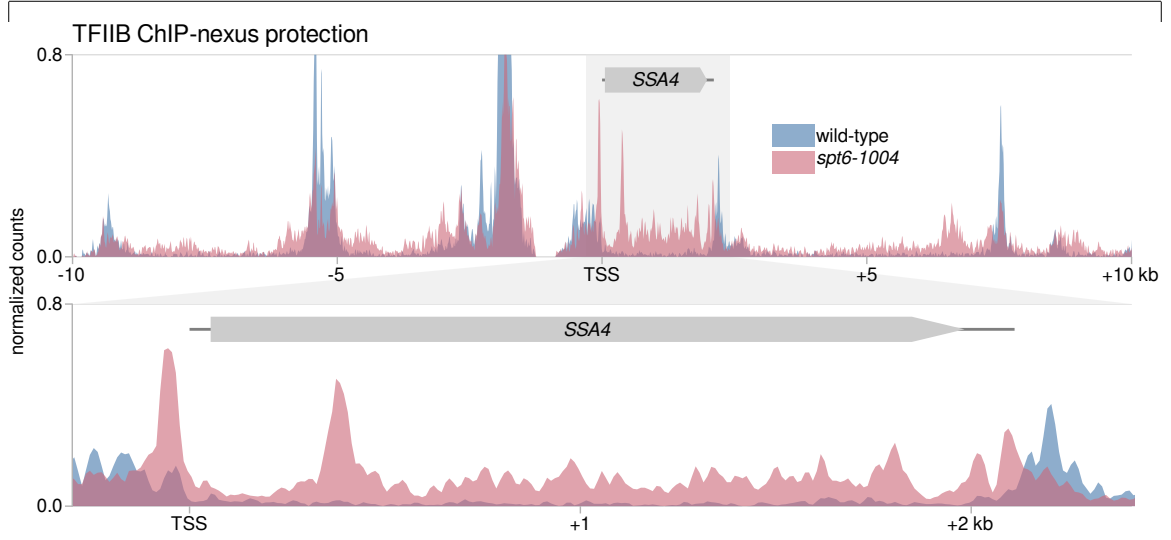


Figure 2.9:

- top) TFIIB ChIP-nexus protection in wild-type and *spt6-1004*, over 20 kb of chromosome II flanking the *SSA4* gene.
- bottom) Expanded view of TFIIB protection over the *SSA4* gene.

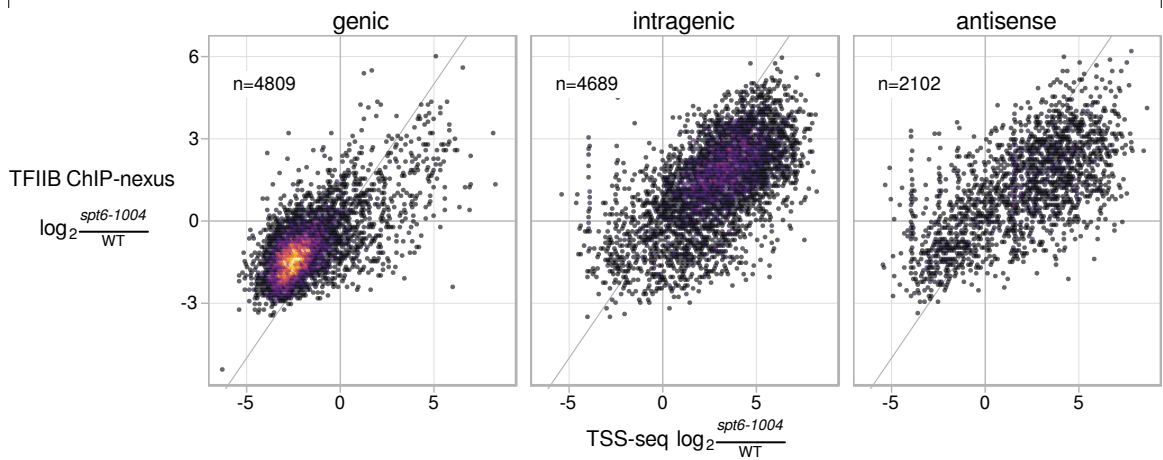


Figure 2.10: Scatterplots of fold-change in *spt6-1004* over wild-type, comparing TSS-seq and TFIIB ChIP-nexus. Each dot represents a TSS-seq peak paired with the window extending 200 bp upstream of the TSS-seq peak summit for quantification of TFIIB ChIP-nexus signal. Fold-changes are regularized fold-change estimates from DESeq2, with size factors determined from the *S. pombe* spike-in (TSS-seq), or *S. cerevisiae* counts (ChIP-nexus).

2.4 MNase-seq results from *spt6-1004*

Because a primary function of Spt6 is to act as histone chaperone that reassembles nucleosomes in the wake of transcription (Duina, 2011), it is reasonable to expect that the transcriptional changes seen in *spt6-1004* would be associated with changes in chromatin structure. The requirement for Spt6 in maintaining normal chromatin structure has been demonstrated in previous studies (Bortvin and Winston, 1996; Ivanovska et al., 2011; Jeronimo et al., 2015; Kaplan et al., 2003; Perales et al., 2013; van Bakel et al., 2013). To re-examine this requirement in higher resolution, we assayed nucleosome protection genome-wide using micrococcal nuclease digestion of chromatin followed by sequencing (MNase-seq).

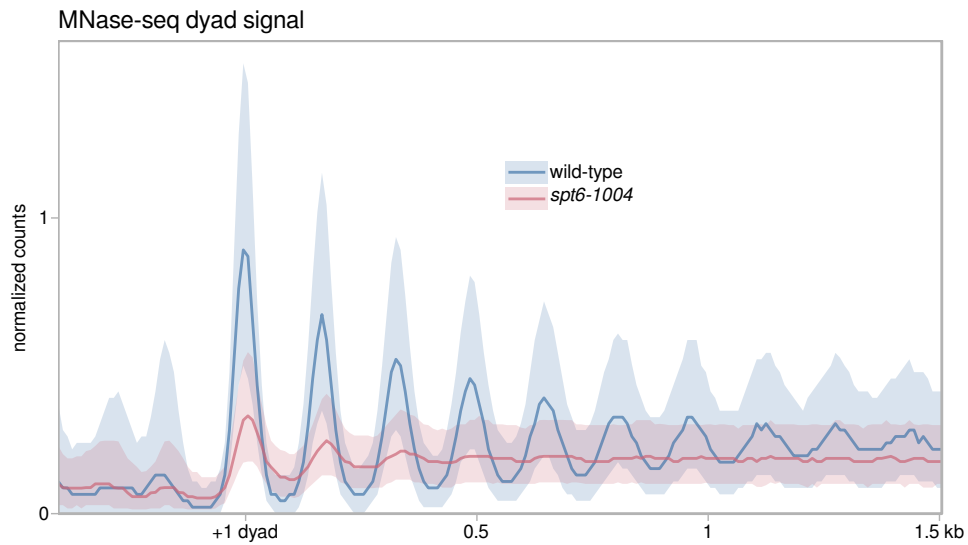


Figure 2.11: Average MNase-seq dyad signal in wild-type and *spt6-1004*, over 3522 non-overlapping genes aligned by wild-type +1 nucleosome dyad. The solid line and shading are the median and inter-quartile range of the mean spike-in normalized coverage over two replicates (*spt6-1004*) or one experiment (wild-type), in non-overlapping 20 bp bins.

In wild-type, the MNase-seq data recapitulate the expected signature over genes, with a nucleosome-depleted region upstream of a strongly positioned ‘+1’ nucleosome, and a regularly phased array of nucleosomes over the gene body (Figure 2.11). In *spt6-1004*, nucleosome signal is severely reduced at canonical nucleosome positions and spreads into inter-nucleosome regions. Changes in aggregate nucleosome sig-

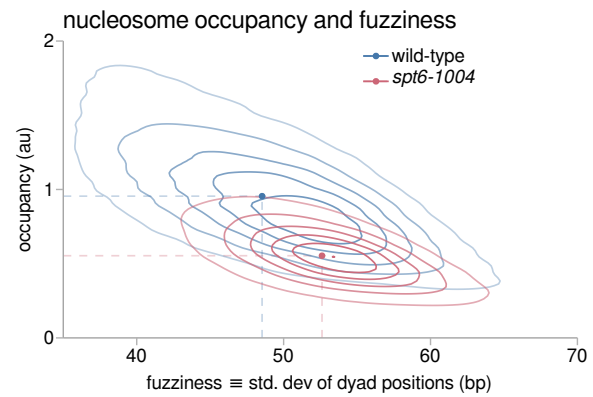


Figure 2.12: Contour plot of the distribution of nucleosome occupancy and fuzziness in wild-type and *spt6-1004*. Dashed lines indicate median values.

nal such as those observed in Figure 2.11 are the combination of changes to nucleosome occupancy (the number of reads assigned to a nucleosome), fuzziness (the standard deviation of read positions for a nucleosome), and position (the coordinate with the maximum reads for a nucleosome) (Chen et al., 2013). Using DANPOS2 (Chen et al., 2013), we called nucleosome positions and quantified these metrics for wild-type and *spt6-1004*. Wild-type nucleosomes span a relatively wide range of occupancy and fuzziness space, with highly occupied nucleosomes tending to be less fuzzy (i.e., more well-positioned) (Figure 2.12). In *spt6-1004*, the population of nucleosomes is much more homogeneous: nucleosome occupancy is decreased globally, and nucleosome fuzziness is restricted to the high end of the wild-type distribution.

Previous studies observed two trends: 1) In wild-type cells, nucleosome positioning is weaker over highly transcribed genes than over moderately transcribed genes (Shivaswamy et al., 2008), and 2) In *spt6-1004* cells, the decrease in nucleosome occupancy is greater for highly transcribed genes (Ivanovska et al., 2011). To re-

examine these trends, we looked at the MNase-seq data in the context of NET-seq data, which reports the position of actively transcribing RNAPII and reflects a gene's level of transcription (Figure 2.13) (Churchman and Weissman, 2012). The data support the first trend: in wild-type, genes with the strongest NET-seq signal have weak patterning of MNase-seq signal. However, we find no obvious relationship between transcription level and the nucleosome occupancy changes observed in *spt6-1004* (Figure 2.13): Genes with the greatest transcription do tend to have lower MNase-seq signal in *spt6-1004*, but this is expected since these genes also have lower MNase-seq signal in wild-type. The discrepancy with prior work might be explained by the greater resolution and breadth of MNase-seq versus MNase and microarray of chromosome III (Ivanovska et al., 2011).

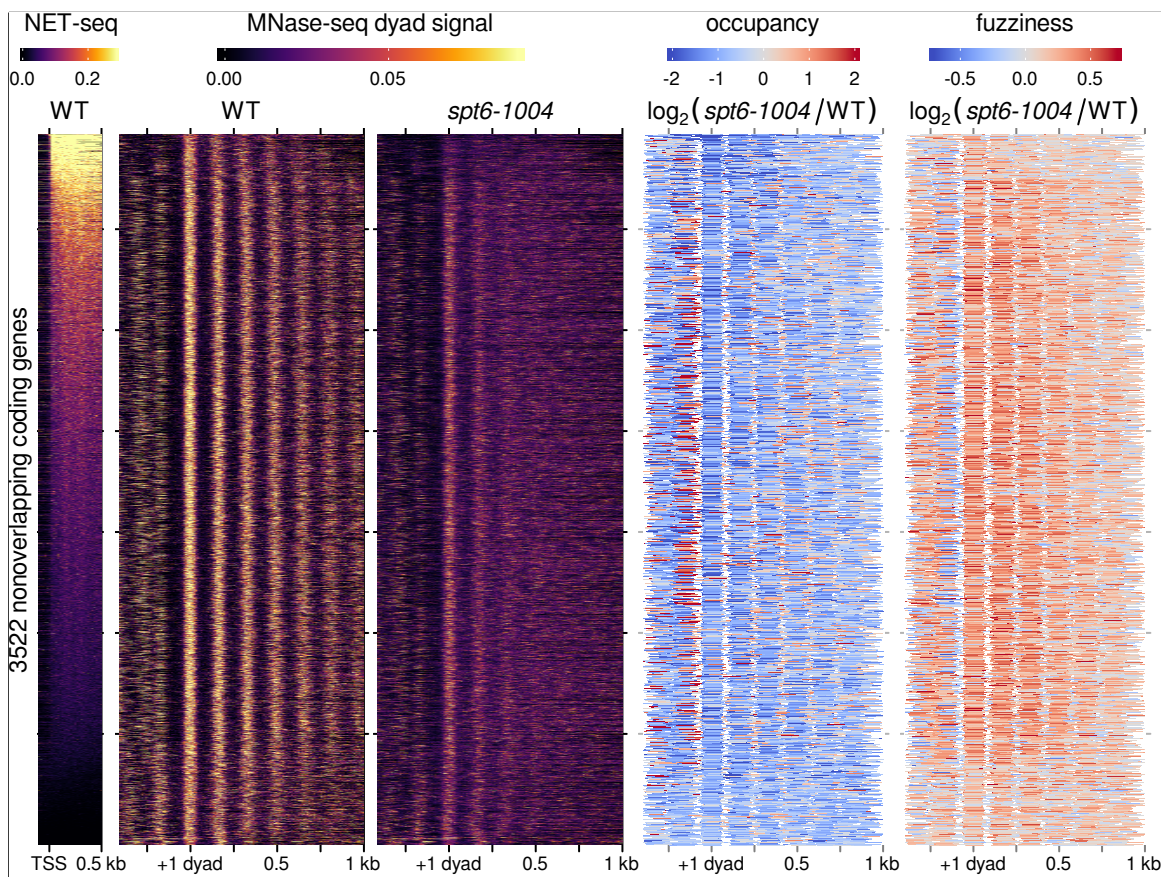


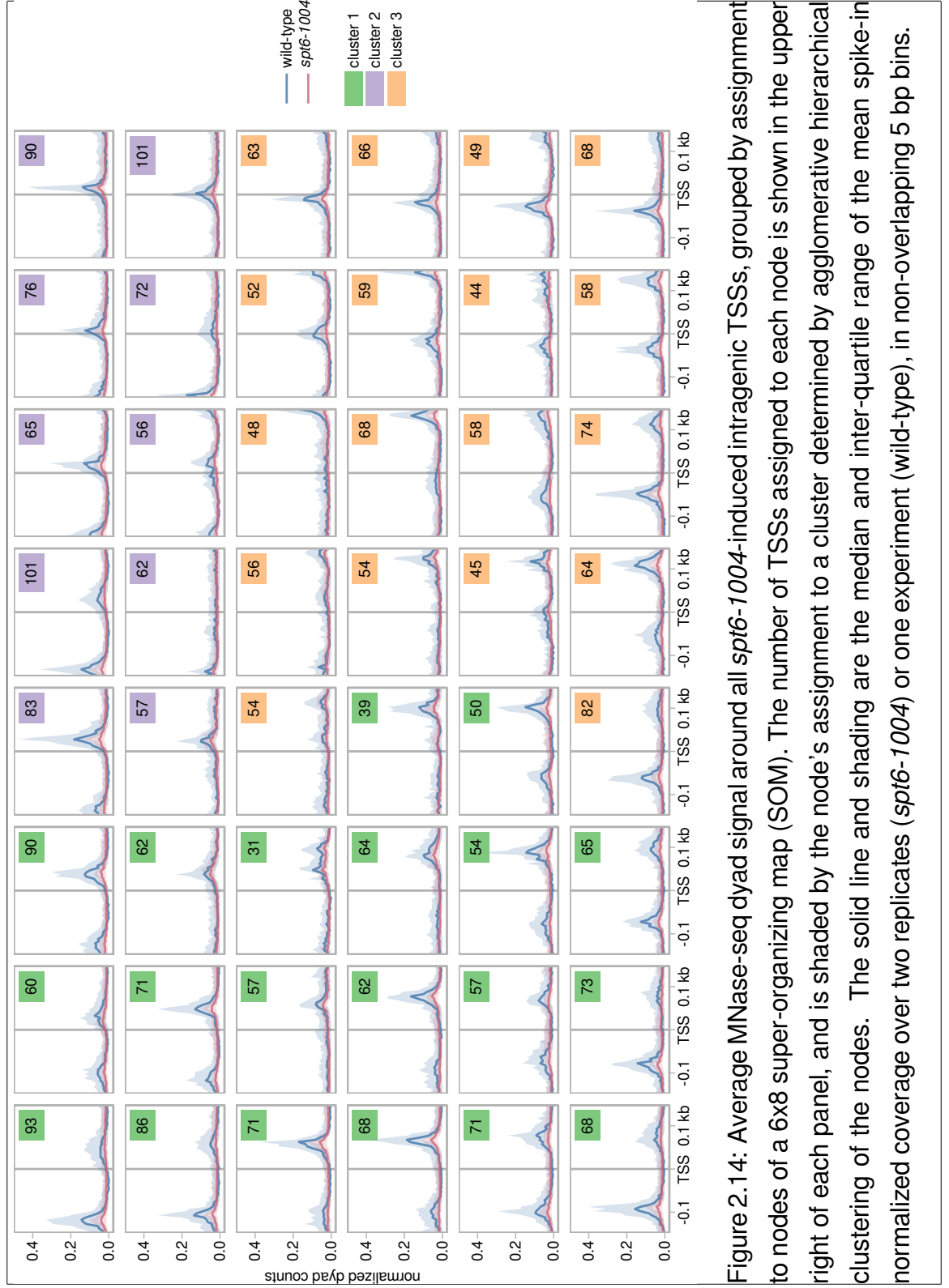
Figure 2.13:

- left) Heatmap of sense strand NET-seq signal for 3522 non-overlapping genes, aligned by genic TSS and sorted by total sense strand NET-seq signal in the window extending 500 nt downstream from the genic TSS. Values are the mean of library-size normalized coverage in non-overlapping 20 nt bins, averaged over two replicates.
- middle) Heatmaps of MNase-seq dyad signal in wild-type and *spt6-1004* for the same genes, aligned by wild-type +1 nucleosome dyad and arranged by sense NET-seq signal as in the leftmost panel. Values are the mean of spike-in normalized coverage in non-overlapping 20 bp bins, averaged over two replicates (*spt6-1004*) or one experiment (wild-type).
- right) Heatmaps of fold-change in nucleosome occupancy and fuzziness for the same genes, aligned by wild-type +1 nucleosome dyad and arranged by sense NET-seq signal as in the leftmost panel.

2.4.1 Clustering of MNase-seq profiles at *spt6-1004*-induced intragenic TSSs

The aggregate MNase-seq dyad signal around all *spt6-1004* intragenic TSSs is aperiodic (Figure 2.15, top left panel), which occurs as a result of destructive interference from offset nucleosome phasing patterns. To discover these phasing patterns, we used the wild-type and *spt6-1004* MNase-seq data flanking intragenic TSSs to train a self-organizing map to assign TSSs with similar MNase-seq patterns to nearby nodes in a rectangular grid (Figure 2.14). This allowed us to see that, although there is considerable diversity in the nucleosome pattern surrounding intragenic TSSs, most intragenic TSSs occur in areas between the positions of nucleosome dyads. By hierarchically clustering the nodes of the self-organizing map, we further grouped intragenic TSSs into three major clusters differing primarily by the phasing of the nucleosome array relative to the TSS, as shown in Figure 2.15. In all three clusters, nucleosomes are disrupted to similar levels in *spt6-1004*.

Because GC-poor DNA sequences are nucleosome disfavoring and are known to occur in promoter regions (Iyer and Struhl, 1995; Kaplan et al., 2008; Tillo and Hughes, 2009; Zhang et al., 2009), we also examined the GC content surrounding the three clusters of intragenic TSSs. For all three clusters, the GC content of the DNA drops just upstream of the TSS to a slightly lesser degree than for genic TSSs (Figure 2.15).



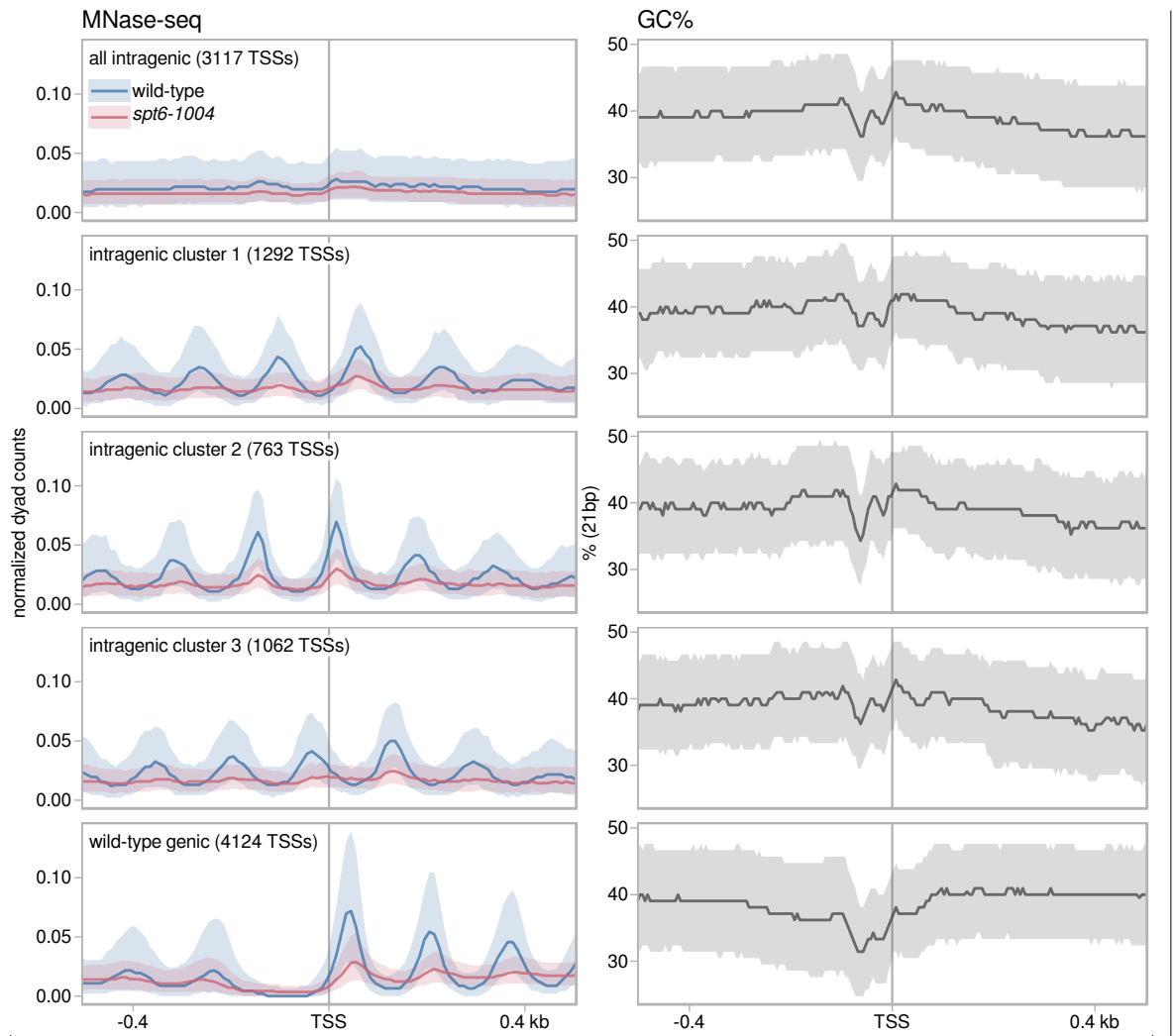


Figure 2.15:

- left column) Average MNase-seq dyad signal for *spt6-1004* intragenic TSSs, both aggregated and grouped into three clusters by the wild-type and *spt6-1004* MNase-seq dyad signal flanking the TSS, as well as all genic TSSs detected in wild-type. Values are the mean of spike-in normalized dyad coverage in non-overlapping 10 bp bins, averaged over two replicates (*spt6-1004*) or one experiment (wild-type). The solid line and shading are the median and inter-quartile range.
- right column) Average GC content of the DNA sequence in a 21 bp window, as above.

2.5 Other features of *spt6-1004* intragenic promoters

MNase-seq indicates that nucleosomes are lost across the entire genome in *spt6-1004*. However, TSSs observed in *spt6-1004* occur in specific locations, suggesting that loss of nucleosomes is necessary but not sufficient for intragenic transcription, and that additional features such as the drop in GC content at intragenic TSSs (Figure 2.15) may be required. The resolution with which we were able to identify intragenic TSSs allowed us to closely examine sequence features that might contribute to intragenic transcription.

2.5.1 Information content and sequence preference

To examine the DNA sequence preference of TSSs in *spt6-1004*, we aligned the sequences of all TSS-seq reads overlapping TSS-seq peaks of each class, and calculated the information content and sequence distribution for each class. Intragenic TSSs have a sequence preference almost identical to the previously observed sequence preference of genic TSSs (Figure 2.16) (Malabat et al., 2015), suggesting that RNA polymerase initiates transcription similarly at genic and intragenic TSSs.

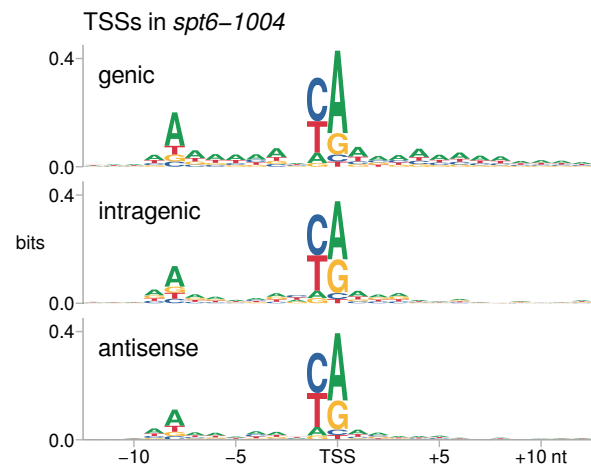


Figure 2.16: Sequence logos depicting information content and sequence preference of TSS-seq reads overlapping genic and intragenic TSS-seq peaks in *spt6-1004*.

2.5.2 Enrichment of the TATA box

A characteristic feature of canonical genic promoters is the presence of a TATA box or TATA-like DNA element which allows for the recruitment of Pol II and general transcription factors via binding of the TFIID complex, which includes TATA-binding protein (Rhee and Pugh, 2012). To examine whether the presence of TATA elements might contribute to *spt6-1004* intragenic transcription, we looked for exact matches to the TATA consensus sequence TATAWAWR in the window extending 200 nucleotides upstream of *spt6-1004* TSSs, finding matches at 13.7% of regions upstream of intragenic TSSs and 24.7% for antisense TSSs, versus 24.4% for all genic TSSs and 8.9% for random locations in the genome. Moreover, the TATA elements found near intragenic and antisense TSSs are highly concentrated in the region 50 to 100 nucleotides upstream of the TSS, where TATA elements are most often found for genic TSSs (Figure 2.17). This further supports the model that *spt6-1004* intragenic promoters are sequences similar to canonical genic promoters, which become accessible for transcription initiation when the normal chromatin state is disturbed.

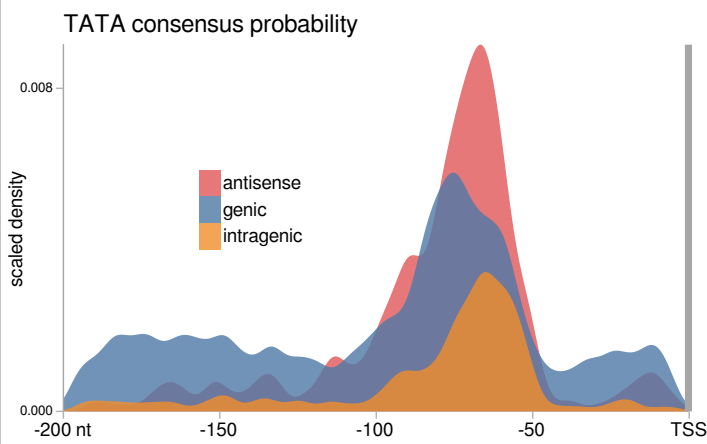


Figure 2.17: Scaled density of occurrences of exact matches to the motif TATAWAWR upstream of TSSs. For each category, a Gaussian kernel density estimate of the positions of motif occurrences is scaled by the number of motif occurrences per region.

2.5.3 Sequence motifs discovered

To discover additional sequence features of *spt6-1004* intragenic promoters, we performed *de novo* motif discovery using MEME-ChIP (Machanick and Bailey, 2011) for the regions -100 to +30 nucleotides relative to TSS summits. The most enriched motif found by MEME at both intragenic and antisense *spt6-1004* TSSs is, with respect to sense genic transcription, a GA-rich motif with 3-nucleotide periodicity 2.18. This motif occurs at only a small subset of intragenic TSSs, but is highly unlikely to occur by chance (compare the expected to observed number of occurrences in Figure 2.18). The motif is not enriched at genic TSSs upregulated in *spt6-1004*, and is not an obvious match to a DNA-binding factor in the databases searched (de Boer and Hughes, 2011; MacIsaac et al., 2006; Newburger and Bulyk, 2008; Ozonov et al., 2012; Teixeira et al., 2017; Weirauch et al., 2014; Zhu and Zhang, 1999). If this motif is directly related to intragenic transcription, we speculate that it might create a DNA structure favorable for transcription initiation.

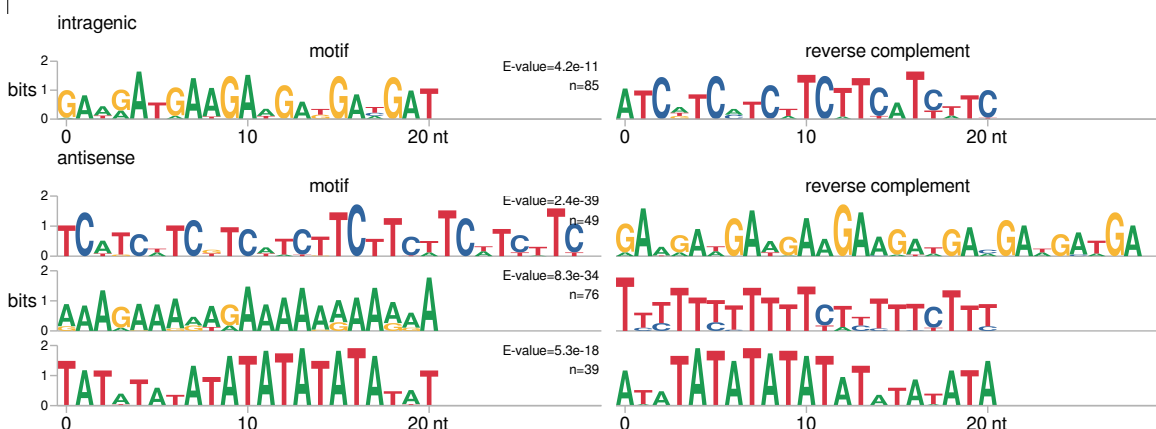


Figure 2.18: Sequence logos of motifs discovered by MEME (Bailey et al., 2015) in the window -100 to +30 bp relative to *spt6-1004* intragenic and antisense TSSs. The number of motif occurrences and the E-value, indicating the expected number of motif occurrences if the input sequences were scrambled, are shown for each motif.

2.6 Discussion

In this work, we integrated multiple quantitative genomic approaches to study the conserved transcription elongation factor Spt6. Our TSS-seq and TFIIB ChIP-nexus results reveal the full extent of intragenic and antisense transcript expression in *spt6-1004*, and show that these transcripts are largely explained by new RNA Pol II transcription initiation. Our MNase-seq results show that this new transcription initiation happens in the context of a global depletion and disordering of nucleosomes from chromatin. We speculate that this dramatic decrease in nucleosome protection of the genome leads to intragenic transcription by allowing initiation factors to access normally inaccessible promoter-like sequences within coding sequences. This model is supported by the similarities we observe between genic and intragenic promoters in DNA GC content, initiation motif, and TATA element frequency. This may also explain the unexpected decrease in transcription initiation we see at almost all genic promoters in *spt6-1004*: Assuming that the pool of transcription initiation factors in the cell is limiting, then making thousands of additional binding sites available to the initiation machinery would decrease the frequency at which the initiation machinery finds its correct targets at genic promoters.

2.7 Methods

2.7.1 Yeast strain construction and growth conditions

All yeast strains were constructed by standard yeast transformation or crosses. The *spt6-1004* and wild-type strains were grown as previously described (Cheung et al., 2008): Cells were grown in YPD at 30 °C to a concentration of approximately 1×10^7 cells/ml ($\text{OD}_{600} = 0.6$), at which point an equal volume of YPD medium pre-warmed to 44 °C was added, and the cultures were shifted to 37 °C for 80 minutes.

2.7.2 Western blotting

The protocols for western blotting and quantification are described in Doris et al. (2018).

2.7.3 Sequencing library preparation (TSS-seq, ChIP-nexus, MNase-seq, NET-seq)

All library preparation methods are detailed in Doris et al. (2018).

2.7.4 Genome builds

The genome build used for *S. cerevisiae* was R64-2-1 (Engel et al., 2014), and the genome build used for *S. pombe* was ASM294v2 (Wood et al., 2002).

2.7.5 TSS-seq data analysis

An up-to-date version of the Snakemake (Köster and Rahmann, 2012) workflow used to process TSS-seq libraries is maintained at github.com/winston-lab/tss-seq. At the time of writing, removal of adapter sequences and random hexamer sequences from the 3' end of the read and 3' quality trimming were performed using cutadapt (Martin,

2011). The random hexamer molecular barcode on the 5' end of the read was then removed and processed using a custom Python script (adapted from Mayer et al. (2015)). Reads were aligned to the combined *S. cerevisiae* and *S. pombe* reference genomes using Tophat2 (Kim et al., 2013) without a reference transcriptome, and uniquely mapping reads were selected using SAMtools (Subgroup et al., 2009). Reads mapping to the same location as another read with the same molecular barcode were identified as PCR duplicates and removed using a custom Python script (adapted from Mayer et al. (2015)). Coverage of the 5'-most base, corresponding to the TSS, was extracted using bedtools genomecov (Quinlan and Hall, 2010) and normalized to the total number of uniquely mapping, non-duplicate *S. pombe* reads. Quality statistics of raw, cleaned, non-aligning, and uniquely aligning non-duplicate reads were assessed using FastQC (Andrews, 2010).

The pipeline additionally performs TSS-seq peak calling, differential expression, classification of peaks into genomic categories, sequence logo visualization, motif enrichment analysis, *de novo* motif discovery, gene ontology analysis (Young et al., 2010), and data visualization with the option to separate data into clusters of similar signal.

2.7.5.1 Reannotation of *S. cerevisiae* TSSs using TSS-seq data

TSS-seq coverage from two replicates of a wild-type *S. cerevisiae* strain grown at 30°C in YPD was averaged and used to adjust the 5' ends of an annotation of major transcript isoforms based on TIF-seq data (Pelechano et al., 2013). The 5' end of the original annotation was changed to the position of maximum TSS-seq signal in a window \pm 250 nt of the original 5' end if the maximum TSS-seq signal was greater than the 95th percentile of all non-zero TSS-seq signal.

2.7.5.2 TSS-seq peak calling

TSS-seq data representing transcription from a single promoter tends to occur as a cluster of signal distributed over a range of positions, rather than a single nucleotide (Arribere and Gilbert, 2013; Malabat et al., 2015). It is reasonable to consider such a cluster of TSS-seq signal as a single entity, because the signals within the cluster are usually highly correlated to one another across different conditions. Therefore, to identify TSSs from TSS-seq data and quantify them for downstream analyses such as differential expression, it is necessary to annotate these groups of TSS-seq signal by using the data to perform peak-calling.

At the time of writing, TSS-seq peak calling for a given experimental group was performed by 1-D watershed segmentation of the data for each sample in the group, followed by filtering for reproducibility within the group by the Irreproducible Discovery Rate (IDR) method (Li et al., 2011). First, a smoothed version of the TSS-seq coverage is generated for each sample using an adaptive two-stage kernel density estimation with a discretized Gaussian kernel (Silverman, 1986). For a given nucleotide, the adaptive kernel bandwidth, σ_{adaptive} , is given by

$$\sigma_{\text{adaptive}} = \sigma_{\text{pilot}} \left(\frac{\rho_{\text{pilot}}}{g} \right)^{-\alpha},$$

where σ_{pilot} is the standard, fixed bandwidth of a Gaussian kernel used to calculate the pilot signal density ρ_{pilot} at that nucleotide, g is the geometric mean of ρ_{pilot} over the whole genome, and α is a parameter in $[0, 1]$ that determines the degree to which the pilot density ρ_{pilot} affects σ_{adaptive} . The adaptive kernel adjusts the kernel bandwidth to be smaller in regions of high signal density and larger in regions of lower signal density, allowing the smoother to better accommodate both ‘sharp’ TSSs where the

signal is distributed over a relatively small window, as well as ‘broad’ TSSs where the signal is more dispersed. For all analyses in this document, adaptive smoothing was performed with $\sigma_{\text{pilot}} = 10$ and $\alpha = 0.2$.

Following smoothing, an initial set of peaks is formed by assigning all nonzero signal in the original, unsmoothed coverage to the nearest local maximum of the smoothed coverage, and taking the minimum and maximum genomic coordinates of the original coverage as the peak boundaries for each local maximum of the smoothed coverage. Peaks are then trimmed to the smallest genomic interval that includes 95% of the original coverage, and the probability of the peak being generated by noise is estimated by a Poisson model where λ , the expected coverage, is the maximum of the expected coverage over the chromosome and the expected coverage in the 2 kb window upstream of the peak (à la the ChIP-seq peak caller MACS2 (Zhang et al., 2008)). Finally, peaks are ranked by their significance under the Poisson model, and a final list of peaks for the group is generated using the IDR method (IDR = 0.1) (Li et al., 2011). In brief, IDR compares ranked lists of regions in order to set a cutoff, beyond which the regions are no longer consistent between replicates.

The python script used for 1-D watershed segmentation of TSS-seq data is available as part of the TSS-seq pipeline, and the IDR implementation used in the pipeline is also available on GitHub.

2.7.5.3 TSS differential expression analysis

For TSS-seq differential expression analysis, TSS-seq peak-calling was performed as described above for both *S. cerevisiae* and the *S. pombe* spike-in. The read counts for each peak in each condition were used as the input to differential expression analysis by DESeq2 (Love et al., 2014), with the alternative hypothesis

$|\log_2(\text{fold-change})| > 1.5$ and a false discovery rate of 0.1. To normalize by spike-in, the size factors of the *S. pombe* spike-in counts were used as the size factors for *S. cerevisiae*, although we note that due to the median of ratios normalization used in DESeq2, the major TSS-seq results of this work are still observed when *S. cerevisiae* size factors are used.

2.7.5.4 Classification of TSS-seq peaks into genomic categories

TSS-seq peaks were assigned to genomic categories based on their position relative to the transcript annotation described above and an annotation of all verified open reading frames (ORFs) and blocked reading frames in *S. cerevisiae* (Engel et al., 2014). First, ‘genic’ regions were defined as follows: If a gene was present in both the transcript and ORF annotations, the genic region was defined as the interval (annotated TSS-30 nt, start codon). If gene was present in the transcript annotation but not the ORF annotation, the genic region was defined as the interval (annotated TSS - 30 nt, annotated TSS + 30 nt). If a gene was present only in the ORF annotation, the genic region was defined as the interval (start codon - 30 nt, start codon). For the purposes of peak classification, regions were considered overlapping if they had at least one base of overlap. TSS-seq peaks were classified as genic if they overlapped a genic region on the same strand. Peaks were classified as intragenic if they were not classified as a genic peak, and their summit position overlapped an open or closed reading frame on the same strand. Peaks were classified as antisense if their summit position overlapped a transcript on the opposite strand. Finally, peaks were classified as intergenic if they did not overlap a transcript, reading frame, or genic region on either strand.

2.7.5.5 TSS information content and sequence composition

TSS-seq alignments were pooled for all replicates in a condition, and the DNA sequence flanking the position of every read overlapping TSS-seq peaks of a particular genomic category was extracted using SAMtools (Subgroup et al., 2009) and bedtools (Quinlan and Hall, 2010). The information content and sequence composition was quantified using WebLogo (Crooks et al., 2004), with the zeroth-order Markov model of the *S. cerevisiae* genomic sequence as the background composition. Sequence logos were plotting using helper functions from ggseqlogo (Wagih, 2017).

2.7.5.6 Enrichment of the TATA box

An up-to-date version of the Snakemake (Köster and Rahmann, 2012) workflow used to test the enrichment of motifs is maintained at github.com/winston-lab/motif-enrichment. To test for enrichment of consensus TATA boxes, FIMO (Grant et al., 2011) was used to search the *S.cerevisiae* genome for matches to the query motif TATAWAWR (where the ambiguous bases are equiprobable) at a p-value threshold of 6×10^{-4} . Regions extending 200 nucleotides upstream of TSS summits were merged if they were overlapping, and were considered to contain a consensus TATA box if the entire motif was overlapping the region on the same strand. The frequency of motif occurrences in the regions of interest was compared to the frequency of occurrences in the regions upstream of 6000 randomly chosen locations in the genome, using Fisher's exact test.

2.7.5.7 *De novo* motif discovery

De novo motif discovery for the regions around TSSs was performed by running MEME-ChIP (Machanick and Bailey, 2011) on the DNA sequence -100 to +30 nu-

cleotides from the TSS summits of the genomic classes of TSSs significantly upregulated in *spt6-1004* versus wild-type.

2.7.6 ChIP-nexus data analysis

An up-to-date version of the Snakemake (Köster and Rahmann, 2012) workflow used to process ChIP-nexus libraries is maintained at github.com/winston-lab/chip-nexus. At the time of writing, filtering for reads containing the constant region of the adapter on the 5' end of the read, 3' adapter removal, and 3' quality trimming were performed using cutadapt (Martin, 2011). The random pentamer molecular barcode on the 5' end of the read was then removed and processed using a custom Python script modified from Mayer et al. (2015). Reads were aligned to the combined *S. cerevisiae* and *S. pombe* genomes using Bowtie2 (Langmead and Salzberg, 2012), and uniquely mapping reads were selected using SAMtools (Subgroup et al., 2009). Reads mapping to the same location as another read with the same molecular barcode were identified as PCR duplicates and removed using a custom Python script modified from Mayer et al. (2015). Coverage of the 5'-most base, corresponding to the point of crosslinking, was extracted using bedtools genomecov (Quinlan and Hall, 2010). The median fragment size estimated by MACS2 (Zhang et al., 2008) over all samples was used to generate coverage of factor protection and fragment midpoints, by extending reads to the fragment size, or by shifting reads by half the fragment size, respectively. Coverage was normalized to the total number of reads uniquely mapping to *S. cerevisiae* (the *S. pombe* spike-in was not used for normalization due to the low number of reads mapping to *S. pombe*). Quality statistics of raw, cleaned, non-aligning, and uniquely aligning non-duplicate reads were assessed using FastQC Andrews (2010).

The ChIP-nexus pipeline additionally performs peak calling, differential occupancy

analysis, and data visualization with the option to separate data into clusters of similar signal.

An second Snakemake workflow for TFIIB-specific analyses is maintained at github.com/winston-lab/chip-nexus-tfib, and performs classification of TFIIB peaks into genomic categories, motif enrichment analysis, and gene ontology analysis.

2.7.6.1 ChIP-nexus peak calling

A number of tools have been created specifically for peak-calling using data from high-resolution ChIP techniques such as ChIP-nexus and ChIP-exo (Hansen et al., 2016; Wang et al., 2014). When applied to our TFIIB ChIP-nexus data, these tools tended to split what appeared to be a single TFIIB binding event into multiple peaks. This may be because TFIIB crosslinks to DNA at multiple points (Rhee and Pugh, 2012), suggesting that while these tools may work well for factors that bind symmetrically to DNA with a single crosslinking point on either side, there is room for improvement for factors with more complex crosslinking patterns.

The ChIP-nexus pipeline currently performs peak calling for a condition using the standard ChIP-seq peak caller MACS2 (Zhang et al., 2008), followed by filtering for reproducibility by the Irreproducible Discovery Rate (IDR) method (IDR = 0.1 for all analyses in this chapter) (Li et al., 2011).

2.7.6.2 TFIIB ChIP-nexus differential occupancy analysis

For TFIIB ChIP-nexus differential binding analysis, TFIIB peaks were called by MACS2 and IDR filtering as described above. A non-redundant list of peaks called in the condition and control groups being compared was generated using bedtools multiinter (Quinlan and Hall, 2010), and the counts of fragment midpoints for each peak in

each sample were used as the input to differential binding analysis by DESeq2 ([Love et al., 2014](#)), with the alternative hypothesis $|\log_2(\text{fold-change})| > 1.5$ and a false discovery rate of 0.1. For estimation of change in TFIIB binding upstream of TSS-seq peaks, TFIIB fragment midpoint counts in the window extending 200 bp upstream of the TSS-seq peak summit were used as the input to DESeq2. *S. cerevisiae* counts were used for size factor calculation.

2.7.6.3 Classification of TFIIB ChIP-nexus peaks into genomic categories

As for TSS-seq peaks, TFIIB ChIP-nexus peaks were assigned to genomic categories based on their position relative to the transcript annotation described above, an annotation of all verified open reading frames (ORFs) and blocked reading frames ([Engel et al., 2014](#)), and [an annotation of ‘genic’ regions derived from the transcript and ORF annotations](#). TFIIB ChIP-nexus peaks were classified as genic if they overlapped a genic region. Peaks were classified as intragenic if they were not classified as a genic peak, and the entire peak overlapped an open or closed reading frame. Finally, peaks were classified as intergenic if they did not overlap a transcript, reading frame, or genic region.

2.7.7 Comparison of TSS-seq to TFIIB ChIP-nexus

An up-to-date version of the Snakemake ([Köster and Rahmann, 2012](#)) workflow used to compare TSS-seq data to TFIIB ChIP-nexus data is maintained at github.com/winston-lab/tss-seq-vs-tfiib-nexus. The pipeline matches and compares peaks from the two assays, and also performs the TFIIB differential occupancy analysis over windows upstream of TSS-seq peaks shown in section 2.3. This differential occupancy analysis was carried out in the same manner as the [differential occupancy](#)

analysis described above, except TFIIB counts were taken over the window extending 200 bp upstream of TSS-seq peak summits.

2.7.8 MNase-seq data analysis

An up-to-date version of the Snakemake (Köster and Rahmann, 2012) workflow used to demultiplex paired-end MNase-seq libraries is maintained at github.com/winston-lab/demultiplex-paired-end. At the time of writing, demultiplexing was performed using fastq-multx (Aronesty, 2013), allowing one mismatch to the barcode, followed by filtering for and removal of the barcode on read 2 using cutadapt (Martin, 2011).

An up-to-date version of the Snakemake (Köster and Rahmann, 2012) workflow used to process MNase-seq libraries is maintained at github.com/winston-lab/mnase-seq. At the time of writing, 3' quality trimming was performed using cutadapt (Martin, 2011). Reads were aligned to the combined *S. cerevisiae* and *S. pombe* genome using Bowtie 1 (Langmead et al., 2009), and correctly paired alignments were selected using SAMtools (Subgroup et al., 2009). Coverage of nucleosome protection and nucleosome dyads were extracted using bedtools (Quinlan and Hall, 2010) and custom shell scripts to get the entire fragment or the midpoint of the fragment, respectively. Smoothed nucleosome dyad coverage was generated by smoothing dyad coverage with a Gaussian kernel of 20 bp bandwidth. Coverage was normalized to the total number of correctly paired *S. pombe* fragments. Quality statistics of raw, cleaned, non-aligning, and correctly paired reads were assessed using FastQC (Andrews, 2010).

The MNase-seq pipeline additionally performs [quantification of nucleosome properties](#), and data visualization with the option to separate data into clusters of similar signal.

2.7.8.1 Quantification of nucleosome properties

Quantification of nucleosome occupancy, fuzziness, and position shifts were calculated using DANPOS2 (Chen et al., 2013), with spike-in normalization by scaling the total counts in condition group libraries by

$$\frac{\text{mean observed percent spike-in in condition libraries}}{\text{mean observed percent spike-in in control libraries}}.$$

2.7.8.2 Clustering of MNase-seq signal at *spt6-1004* intragenic TSSs

The Snakemake (Köster and Rahmann, 2012) workflow for clustering MNase-seq data by self/super-organizing map and hierarchical clustering is maintained at github.com/winston-lab/cluster-mnase-seq. To cluster *spt6-1004* intragenic TSSs based on surrounding MNase-seq signal, spike-in normalized MNase-seq dyad signal in the window ± 150 bp of the TSS-seq peak summit of all intragenic TSS-seq peaks significantly upregulated in *spt6-1004* was binned by taking the mean signal in non-overlapping 5 bp bins, and then averaged taking the mean of two replicates (*spt6-1004*) or one experiment (wild-type). The data were then standardized over each TSS, and the wild-type and *spt6-1004* data were used as equally weighted input layers to a super-organizing map (Wehrens and Buydens, 2007) trained on the input data to assign similar MNase-seq observations in 60-dimensional input space to similar nodes in a 2-dimensional (6×8) rectangular grid. The 48 ‘code vectors’ representing the typical MNase-seq pattern for each node (visualized in Figure 2.14) were then clustered by agglomerative hierarchical clustering using sum of squares distance and Ward linkage. The resulting dendrogram was cut to produce the three clusters of MNase-seq signal shown in Figures 2.14 and 2.15.

2.7.9 NET-seq data analysis

An up-to-date version of the Snakemake (Köster and Rahmann, 2012) workflow used to process NET-seq libraries is maintained at github.com/winston-lab/net-and-rna-seq.

At the time of writing, removal of adapter sequences from the 3' end of the read and 3' quality trimming were performed with cutadapt (Martin, 2011). Reads were aligned to the *S. cerevisiae* genome using Tophat2 (Kim et al., 2013) without a reference transcriptome, and uniquely mapping reads were selected using SAMtools (Subgroup et al., 2009). Coverage of the 5'-most base of the read, corresponding to the 3'-most base of the nascent RNA and the active site of elongating RNA polymerase, was extracted using bedtools genomecov (Quinlan and Hall, 2010) and normalized to the total number of uniquely mapped reads. Quality statistics of raw, cleaned, non-aligning, and uniquely aligning reads were assessed using FastQC (Andrews, 2010).

The NET-seq pipeline additionally performs *ab initio* transcript annotation (Pertea et al., 2015), differential expression analysis, and data visualization with the option to split data into clusters of similar signal. For libraries with unique molecular barcodes and/or spike-ins, the pipeline also handles PCR duplicate removal and spike-in normalization, respectively.

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Chapter 3

Genomics of transcription elongation factor Spt5

3.1 Collaborators

Ameet Shetty generated TSS-seq, MNase-seq, NET-seq, RNA-seq, and ChIP-seq libraries

3.2 Introduction to Spt5 and prior work

Relevant information about Spt5 is summarized as follows (Shetty et al., 2017):

- Spt5 is the only transcription factor known to be conserved in all three domains of life (Hartzog and Fu, 2013; Werner, 2012).
- Spt5 co-localizes with elongating RNA Pol II (Mayer et al., 2010; Rahl et al., 2010).
- Spt5 binds over the Pol II clamp domain, likely stabilizing the elongation complex (Hirtreiter et al., 2010; Klein et al., 2011; Martinez-Rucobo et al., 2011).
- Spt5 physically recruits factors to the elongating transcription complex, in a manner dependent on the modification status of its C-terminal region (CTR) (Hartzog and Fu, 2013):
 - in its unphosphorylated state, the CTR aids in recruiting the mRNA capping enzyme (Doamekpor et al., 2014, 2015; Schneider et al., 2010; Wen

and Shatkin, 1999)

- in its phosphorylated state, the CTR recruits the Paf1 complex, which is important for Pol II elongation (Liu et al., 2009; Mbognign et al., 2013; Wier et al., 2013; Zhou et al., 2009)
- Spt5 helps to recruit mRNA 3' end processing factors (Mayer et al., 2012; Stadelmayer et al., 2014; Yamamoto et al., 2014).
- Spt5 helps to recruit the Rpd3S histone deacetylase complex (Drouin et al., 2010).

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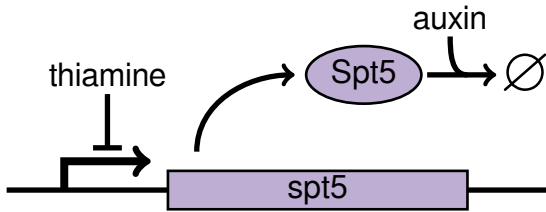


Figure 3.1: Diagram of the dual-shutoff system used to deplete Spt5 from *S. pombe*. Spt5 is expressed from a thiamine-repressible promoter, and tagged with an auxin-inducible degron tag for specific degradation upon addition of auxin.

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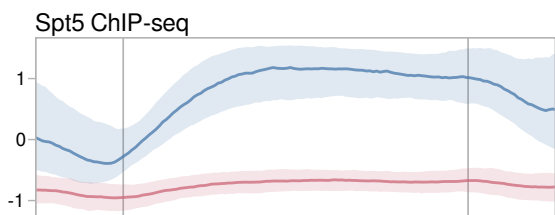
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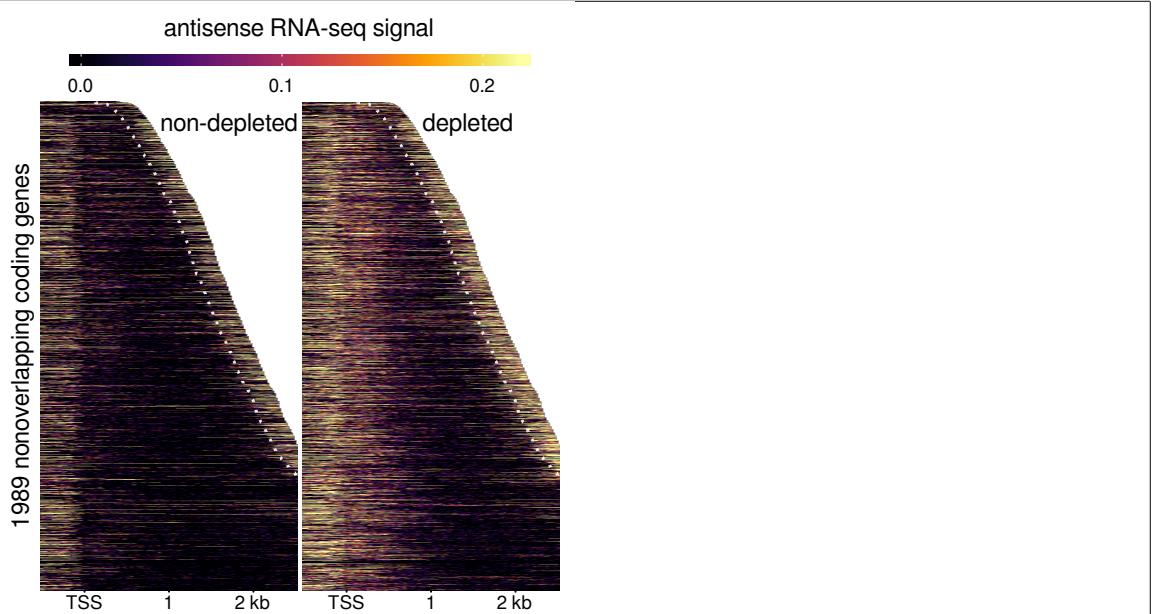
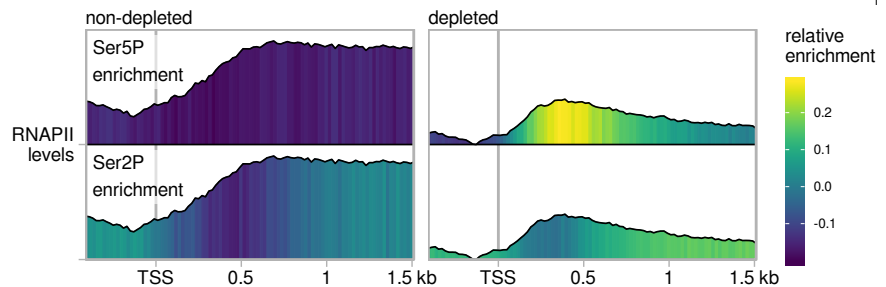


Figure 3.4: Caption wsdasdr zzzz.

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3.3 An aside on spike-in normalization for ChIP-seq

3.4 TSS-seq results from Spt5 depletion

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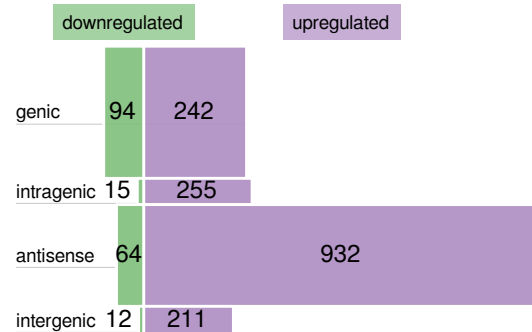


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3.5 MNase-seq results from Spt5 depletion

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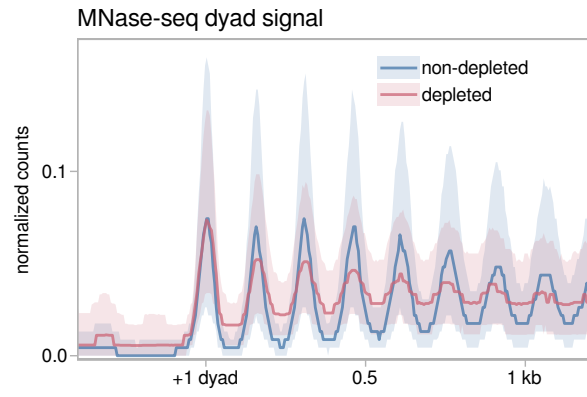


Figure 3.7: Caption wsdasdr zzzz.

Figure 3.8: Caption wsdasdr zzzz.

Figure 3.9: Caption wsdasdr zzzz.

3.5.1 MNase-seq profile at Spt5-depletion-induced antisense TSSs

3.6 Sequence motifs enriched at antisense TSSs

3.7 Discussion

3.8 Methods

3.8.1 A note on spike-in normalization for ChIP-seq experiments with input samples

In the course of determining how to do spike-in normalization for ChIP-seq libraries, I discovered the following error in a published spike-in normalization method. Throughout the following explanation, I use ‘experimental’ and ‘spike-in’ to refer to the two genomes present in the experiment, e.g., experimental signal and spike-in signal.

The goal when including spike-ins in a ChIP-seq experiment is to be able to normalize the experimental signal, such that the normalized signal is proportional to the absolute abundance of the factor being immunoprecipitated. A straightforward method to accomplish this normalization is to linearly scale the experimental signal of a library by a normalization factor, which we will call α . To calculate α for each library, we can use the fact that a normalized ‘spike-in signal’ should be the same for all libraries, since the biological state of the spike-in cells is the same in all libraries. The key to correctly determining α is defining exactly what this spike-in signal is.

The measurement we begin with for determination of the spike-in signal of a library is the number of reads in the library which map uniquely to the spike-in genome

(R_{spike}) . This value will vary based on two factors: the sequencing depth of the library, and the proportion of cells which were spike-in cells (ϕ):

R_{spike} \equiv the number of reads in the library mapping uniquely to the spike-in genome;

ϕ \equiv the proportion of spike-in cells in the sample.

However, the derivation of α is more easily understood in terms of absolute cell numbers rather than ϕ :

C_{exp} \equiv the number of experimental cells used to prepare a library;

C_{spike} \equiv the number of spike-in cells used to prepare a library.

We can express the **number of spike-in reads per spike-in cell** by simply taking the fraction $\frac{R_{\text{spike}}}{C_{\text{spike}}}$. We know that the biological state of a spike-in cell is the same regardless of which sample it belongs to, so we *could* set $\frac{R_{\text{spike}}}{C_{\text{spike}}}$ equal to all samples in order to calculate α . However, this would not account for differences in ϕ between samples: Two libraries representing the same condition and sequenced to the same depth should have equivalent values of $\frac{R_{\text{spike}}}{C_{\text{spike}}}$, which does not hold true if they differed in the proportion of spike-in added.

The metric for ‘spike-in signal’ that leads to the correct expression for α is the **number of spike-in reads per spike-in cell per experimental cell**:

$$\begin{aligned} & \frac{\frac{R_{\text{spike}}}{C_{\text{spike}}}}{C_{\text{exp}}} \\ &= \frac{R_{\text{spike}} C_{\text{exp}}}{C_{\text{spike}}}. \end{aligned}$$

From here, it's simple to calculate α by setting this value to be equal for all samples. Since the actual value of the spike-in signal doesn't matter as long as it is equal for all libraries, we can arbitrarily set it to 1 for convenience:

$$\alpha \frac{R_{\text{spike}} C_{\text{exp}}}{C_{\text{spike}}} = 1$$

$$\alpha = \frac{C_{\text{spike}}}{R_{\text{spike}} C_{\text{exp}}}.$$

Notice that only the ratio of spike-in to experimental cells is needed to calculate α , and not the absolute number of spike-in and experimental cells. We can rewrite this expression in terms of ϕ , the proportion of the sample that was spike-in cells:

$$\phi = \frac{C_{\text{spike}}}{C_{\text{spike}} + C_{\text{exp}}}$$

$$C_{\text{spike}} = \phi (C_{\text{spike}} + C_{\text{exp}})$$

$$C_{\text{spike}} (1 - \phi) = \phi C_{\text{exp}}$$

$$\frac{C_{\text{spike}}}{C_{\text{exp}}} = \frac{\phi}{1 - \phi} \quad \alpha = \frac{C_{\text{spike}}}{R_{\text{spike}} C_{\text{exp}}}$$

$$\alpha = \frac{\phi}{R_{\text{spike}} (1 - \phi)}.$$

This form for α differs from the one presented in ? with no derivation:

$$\alpha = \frac{\phi}{R_{\text{spike}} (1 - \phi)} \quad \alpha_{\text{orlando}} = \frac{\phi}{R_{\text{spike}}}.$$

Working through a few examples with both versions of α will reveal that α_{orlando} leads to incorrect normalization when ϕ is not equivalent for all samples.

In the first example, we will vary sequencing depth between two libraries, keeping everything else constant. Consider a single ChIP library prep in which 20% of the cells were spike-in cells (i.e., $\phi = 0.2$). The library is then unevenly split into two

aliquots and sequenced. One library has four times the reads of the other library.

$$R_{\text{spike}_1} = 1$$

$$R_{\text{spike}_2} = 4$$

$$R_{\text{exp}_1} = 4$$

$$R_{\text{exp}_2} = 16$$

$$\begin{aligned}\alpha_1 &= \frac{\phi}{R_{\text{spike}_1}(1 - \phi)} & \alpha_2 &= \frac{\phi}{R_{\text{spike}_2}(1 - \phi)} & \alpha_{\text{orlando}_1} &= \frac{\phi}{R_{\text{spike}_1}} & \alpha_{\text{orlando}_2} &= \frac{\phi}{R_{\text{spike}_2}} \\ \alpha_1 &= \frac{0.2}{1(0.8)} & \alpha_2 &= \frac{0.2}{4(0.8)} & \alpha_{\text{orlando}_1} &= \frac{0.2}{1} & \alpha_{\text{orlando}_2} &= \frac{0.2}{4} \\ \alpha_1 &= \frac{4}{16} & \alpha_2 &= \frac{1}{16} & \alpha_{\text{orlando}_1} &= \frac{4}{20} & \alpha_{\text{orlando}_2} &= \frac{1}{20}.\end{aligned}$$

The total levels of spike-in normalized experimental signal can be found for each library by multiplying α by R_{exp} , for our version of α ,

$$\text{signal}_1 = \alpha_1 R_{\text{exp}_1}$$

$$\text{signal}_2 = \alpha_2 R_{\text{exp}_2}$$

$$\text{signal}_1 = \frac{4}{16}(4)$$

$$\text{signal}_2 = \frac{1}{16}(16)$$

$$\text{signal}_1 = 1$$

$$\text{signal}_2 = 1$$

and for α_{orlando} :

$$\text{signal}_{\text{orlando}_1} = \alpha_{\text{orlando}_1} R_{\text{exp}_1}$$

$$\text{signal}_{\text{orlando}_2} = \alpha_{\text{orlando}_2} R_{\text{exp}_2}$$

$$\text{signal}_{\text{orlando}_1} = \frac{4}{20}(4)$$

$$\text{signal}_{\text{orlando}_2} = \frac{1}{20}(16)$$

$$\text{signal}_{\text{orlando}_1} = 0.8$$

$$\text{signal}_{\text{orlando}_2} = 0.8$$

Only the relative abundances within normalization methods matter, so in this case both calculations correctly normalized for library size and say that the normalized signal in the two libraries are the same.

Now let's consider two libraries from two different conditions with $\phi = 0.1$. In condition 2, there is a known global decrease in experimental signal expected. This time, we will skip the algebra:

$$R_{\text{spike}_1} = 1$$

$$R_{\text{spike}_2} = 4$$

$$R_{\text{exp}_1} = 9$$

$$R_{\text{exp}_2} = 6$$

$$\alpha_1 = \frac{4}{36} \quad \alpha_2 = \frac{1}{36} \quad \alpha_{\text{orlando}_1} = \frac{4}{40} \quad \alpha_{\text{orlando}_2} = \frac{1}{40}$$

$$\text{signal}_1 = 1 \quad \text{signal}_2 = 1/6 \quad \text{signal}_{\text{orlando}_1} = 0.9 \quad \text{signal}_{\text{orlando}_2} = 0.15$$

Both methods correctly detect that experimental signal levels in library 2 are 1/6th that of library 1.

Finally, let's consider two libraries from the same condition which were spiked in with different amounts of spike-in cells. Both libraries are sequenced to the same depth. Since the libraries are from the same condition, we expect their total experimental signal to be the same after normalization, even though they had different

amounts of spike-in added.

$$\phi_1 = 0.2$$

$$\phi_2 = 0.4$$

$$R_{\text{spike}_1} = 2$$

$$R_{\text{spike}_2} = 4$$

$$R_{\text{exp}_1} = 8$$

$$R_{\text{exp}_2} = 6$$

$$\begin{aligned}\alpha_1 &= \frac{\phi_1}{R_{\text{spike}_1}(1 - \phi_1)} & \alpha_2 &= \frac{\phi_2}{R_{\text{spike}_2}(1 - \phi_2)} & \alpha_{\text{orlando}_1} &= \frac{\phi_1}{R_{\text{spike}_1}} & \alpha_{\text{orlando}_2} &= \frac{\phi_2}{R_{\text{spike}_2}} \\ \alpha_1 &= \frac{0.2}{2(0.8)} & \alpha_2 &= \frac{0.4}{4(0.6)} & \alpha_{\text{orlando}_1} &= \frac{0.2}{2} & \alpha_{\text{orlando}_2} &= \frac{0.4}{4} \\ \alpha_1 &= \frac{3}{24} & \alpha_2 &= \frac{4}{24} & \alpha_{\text{orlando}_1} &= \frac{1}{10} & \alpha_{\text{orlando}_2} &= \frac{1}{10}\end{aligned}$$

$$\text{signal}_1 = \alpha_1 R_{\text{exp}_1}$$

$$\text{signal}_2 = \alpha_2 R_{\text{exp}_2}$$

$$\text{signal}_1 = \frac{3}{24} (8)$$

$$\text{signal}_2 = \frac{4}{24} (6)$$

$$\text{signal}_1 = 1$$

$$\text{signal}_2 = 1$$

$$\text{signal}_{\text{orlando}_1} = \alpha_{\text{orlando}_1} R_{\text{exp}_1}$$

$$\text{signal}_{\text{orlando}_2} = \alpha_{\text{orlando}_2} R_{\text{exp}_2}$$

$$\text{signal}_{\text{orlando}_1} = \frac{1}{10} (8)$$

$$\text{signal}_{\text{orlando}_2} = \frac{1}{10} (6)$$

$$\text{signal}_{\text{orlando}_1} = 0.8$$

$$\text{signal}_{\text{orlando}_2} = 0.6$$

Here, our method correctly normalizes the two samples to the same total experimental signal while using the Orlando α results in an apparent decrease in signal in library

2. This is because the Orlando α fails to account for the fact that when you add more spike-in to a sample, you necessarily decrease the proportion of the sample that is experimental. In most experiments with spike-ins, this isn't really a problem because we assume that ϕ is the same for all samples. However, with ChIP-seq experiments that include input samples, if we assume that the experimental and spike-in input sample read counts are proportional to the amounts of experimental and spike-in cells mixed, we can plug these values in for values of ϕ to get a more reliable estimation of experimental signal levels. In this case, it becomes important to use the correct equation for α .

So, putting everything together, here's how I use the spike-in to normalize an IP ChIP-seq library paired with an input ChIP-seq library.

As stated above, we assume that the experimental and spike-in read counts in the input sample are proportional to the numbers of experimental and spike-in cells used to prepare the library:

$$R_{\text{input}_{\text{exp}}} \propto C_{\text{exp}},$$

$$R_{\text{input}_{\text{spike}}} \propto C_{\text{spike}}$$

Therefore, we can plug these values in for C for both the input and IP libraries (using the form of α without ϕ):

$$\alpha_{\text{input}} = \frac{C_{\text{input}_{\text{spike}}}}{R_{\text{input}_{\text{spike}}} C_{\text{input}_{\text{exp}}}}$$

$$\alpha_{\text{input}} \propto \frac{R_{\text{input}_{\text{spike}}}}{R_{\text{input}_{\text{spike}}} R_{\text{input}_{\text{exp}}}}$$

$$\alpha_{\text{input}} \propto \frac{1}{R_{\text{input}_{\text{exp}}}}$$

$$\alpha_{\text{IP}} = \frac{C_{\text{input}_{\text{spike}}}}{R_{\text{IP}_{\text{spike}}} C_{\text{input}_{\text{exp}}}}$$

$$\alpha_{\text{IP}} \propto \frac{R_{\text{input}_{\text{spike}}}}{R_{\text{IP}_{\text{spike}}} R_{\text{input}_{\text{exp}}}}$$

Notice how α_{input} reduces down to normalizing by the experimental library size, with no dependence at all on the spike-in. This makes sense because the input always represents the same state, regardless of how much spike-in is added to it. The function of the spike-in in the input is only to allow us to estimate abundances in the corresponding IP library. Rewriting α_{IP} in the form

$$\alpha_{\text{IP}} \propto \frac{1}{R_{\text{IP}_{\text{spike}}} \frac{R_{\text{input}_{\text{exp}}}}{R_{\text{input}_{\text{spike}}}}}$$

shows that α_{IP} will basically scale the experimental IP signal to the same scale as the experimental input signal, using the spike-in as a link between the two samples. This makes it natural to subtract the normalized input signal from the normalized IP signal: since they are on the same scale, the resulting coverage can be interpreted as reporting how much more IP signal was detected than was expected based on the input.

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Chapter 4

Stress-responsive intragenic transcription

4.1 Collaborators

Steve Doris generated TSS-seq and ChIP-nexus libraries

Dan Spatt polyribosome fractionation, fitness competitions,
and other experiments

James Warner fitness competitions and other experiments

4.2 Possible functions for intragenic transcription in wild-type cells

4.3 Discovery of stress-induced intragenic promoters by TFIIB ChIP-nexus and TSS-seq

4.4 Chromatin landscape of oxidative-stress-induced promoters.

4.5 Polysome enrichment of oxidative-stress-induced intragenic transcripts

4.6 TSS-seq analysis of oxidative stress in *Saccharomyces sensu stricto* species

4.7 Functions of intragenic DSK2 expression in oxidative stress

4.8 Discussion

4.9 Methods

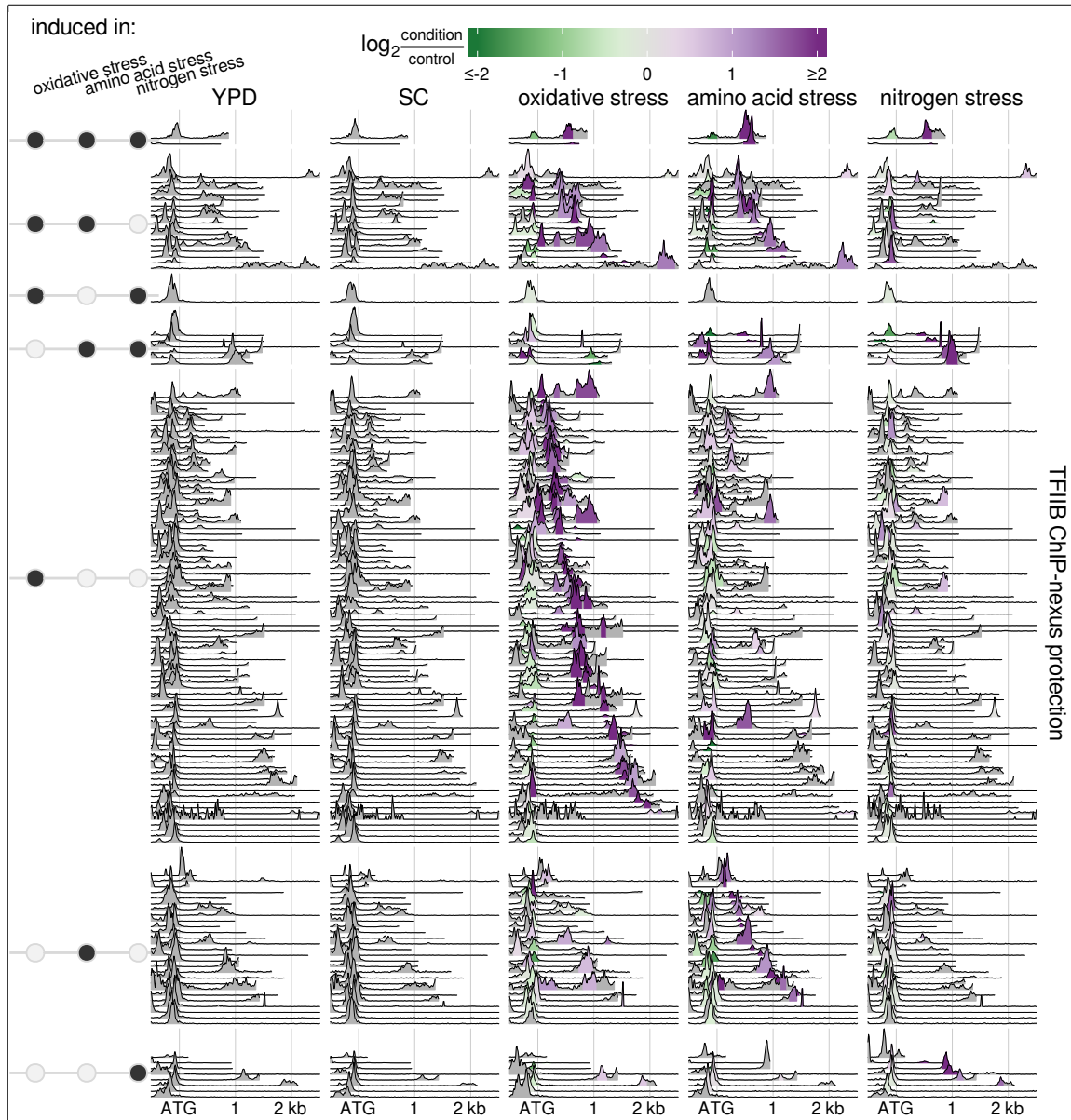


Figure 4.1: Relative TFIIB ChIP-nexus protection over all genes with an intragenic TFIIB peak significantly induced in one or more of the stress conditions tested, as depicted in the left panel. Genes are aligned by start codon, and are sorted within each group by the distance from the start codon to the summit of the induced intragenic TFIIB peak. Data are shown for each gene up to the stop codon of the gene. Regions where TFIIB peaks are called are shaded in the stress conditions according to the fold-change of the peak relative to the corresponding control condition.

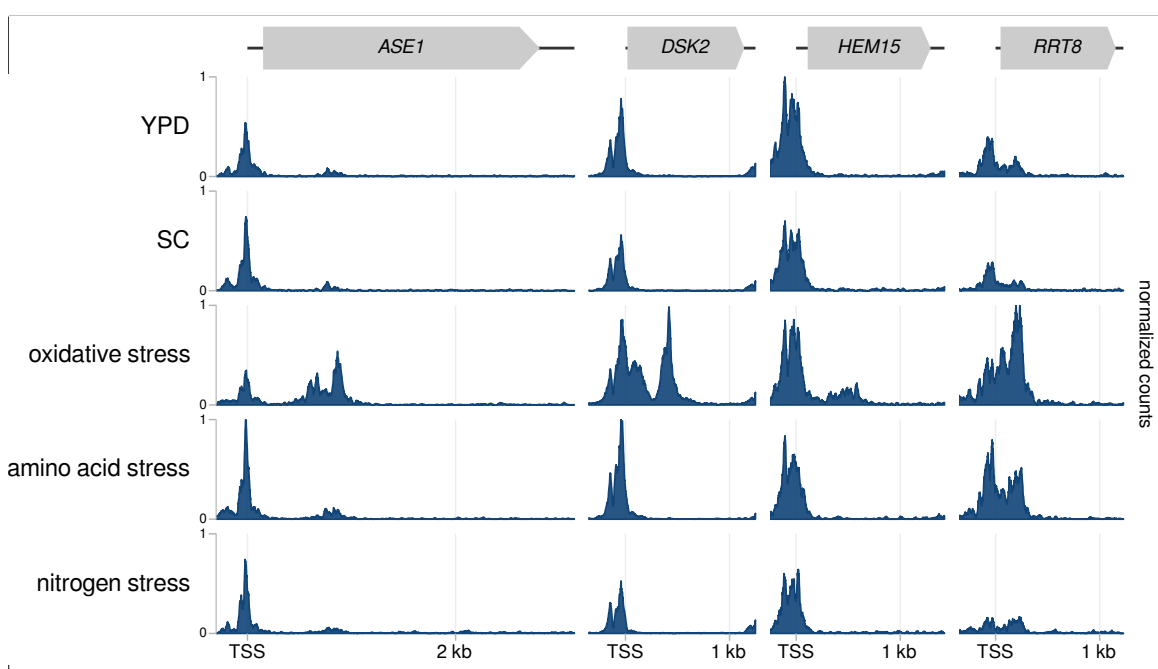


Figure 4.2: Caption asdfkj asldkfjlkj.

4.10 Bibliography

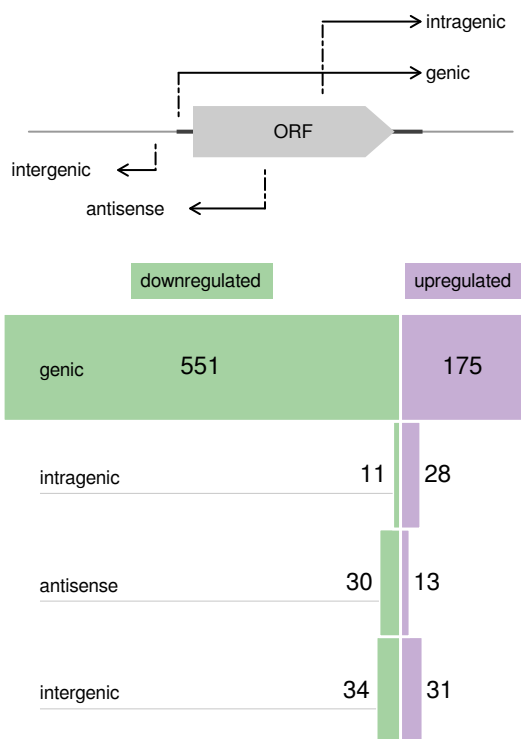


Figure 4.3: Caption dsafklj asldkfjlkj.

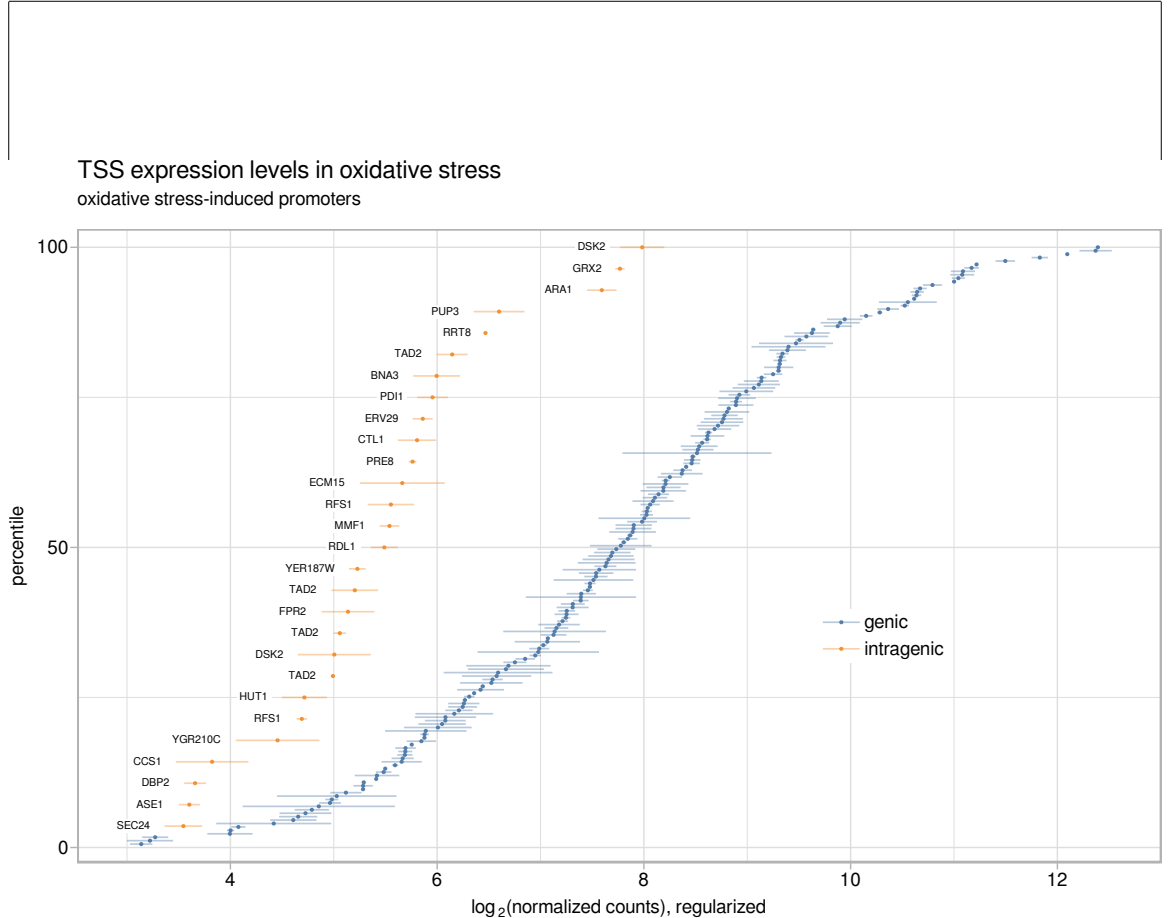


Figure 4.4: Caption dsafklj zzzz.

Figure 4.5: Caption dsafklj .

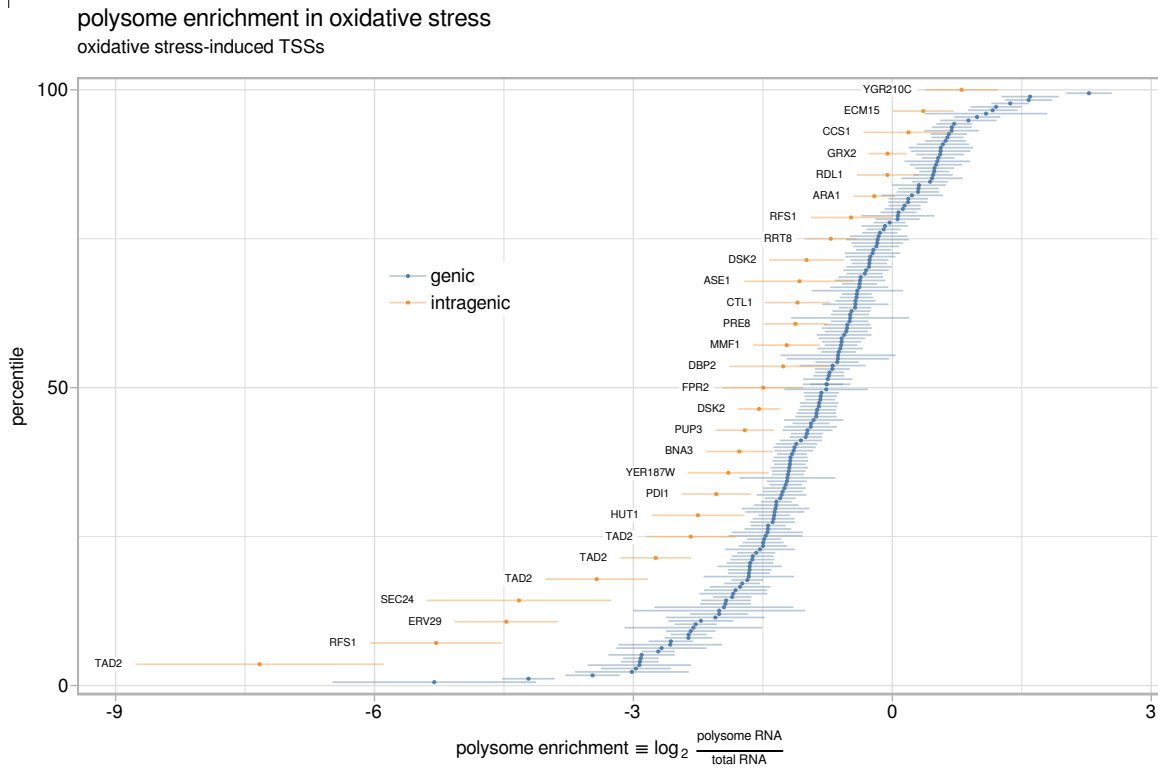


Figure 4.6: Caption wsdasdr zzzz.

Figure 4.7: Caption dsafklj .

Figure 4.8: Caption dsafklj .

Figure 4.9: Caption dsafklj .

Figure 4.10: Caption dsafklj .

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Vita

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