Genome-wide binding analysis of 195 DNA Binding Proteins reveals “reservoir” promoters and human specific SVA-repeat family regulation

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Abstract

A key aspect in defining cell state is the complex choreography of DNA binding events in a given cell type, which in turn establishes a cell-specific gene-expression program. In the past two decades since the sequencing of the human genome there has been a deluge of genome-wide experiments which have measured gene-expression and DNA binding events across numerous cell-types and tissues. Here we re-analyze ENCODE data in a highly reproducible manner by utilizing standardized analysis pipelines, containerization, and literate programming with Rmarkdown. Our approach validated many findings from previous independent studies, underscoring the importance of ENCODE’s goals in providing these reproducible data resources. This approach also revealed several new findings: (i) 1,362 promoters, termed ‘reservoirs,’ have up to 111 different DNA binding-proteins localized on one promoter yet do not have any expression of steady-state RNA (ii) The human specific SVA repeat element may have been co-opted for enhancer regulation. Collectively, this study performed by the students of a CU Boulder computational biology class (BCHM 5631 – Spring 2020) demonstrates the value of reproducible findings and how resources like ENCODE that prioritize data standards can foster new findings with existing data in a didactic environment.

Introduction

In the postgenomic era[1,2] there have been efforts to adapt classical biochemical protocols studying a few DNA regions to genome-wide events. One of the first of these genome-wide assays was Chromatin Immunoprecipitation (ChIP) followed by hybridization of co-precipitate DNA fragments to microarrays (or ChIP-CHIP) representing many thousands of DNA locations...
This application was first demonstrated in yeast and quickly adapted to many species[3–7]. With the advent of massively parallel sequencing technologies, bound DNA from the biochemical ChIP could be sequenced (ChIP-seq) to unbiasedly detect binding events (reviewed[8,9]). This rapid change in platforms for ChIP analyses resulted in many data sets that differed greatly in their results (ChIP-ChIP versus ChIP-seq)[10,11]. Only three years after sequencing of the human genome it became clear that uniform experimental and data standards were essential to limit a deluge of irreproducible results.

To this end, the field turned to the publicly available ENCODE consortium as the largest and most standardized repository of ChIP-seq data sets[12–15]. The goal was to develop standardized experimental and computational pipelines. Over the past 17 years since its inception, many thousands of ChIP-seq experiments have been performed. Often these large consortium studies analyze these data sets across cell types and tissues[13,16–19]. In contrast, fewer studies have investigated dozens of DNA binding proteins (DBPs) in one cell type.

Observing how hundreds of DBPs are bound relative to each other in the same cellular context provides a unique perspective. This allows a promoter-centric approach across hundreds of possible DNA binding events. Thus, we can address the underlying properties of combinatorial binding at promoters and, in turn, how this relates to promoter activity. Moreover, this approach allows us to systematically investigate numerous DBPs for possible enrichment in noncoding regions such as repetitive element class and families. Overall, this strategy is limited in cellular diversity, but rich in relative information of binding events at a given promoter.

By investigating these properties for 195 DBPs in K562 cells, we were able to reproduce known findings from independent data sets. For example, the number of binding events at a promoter
correlates with RNA expression output (both nascent and mature transcripts)[17,18]. We also
made several new observations. Specifically, we identify 1,362 promoters that do not produce a
mature transcript despite having up to 111 DBP binding events. We termed these promoters
“reservoirs” because these promoters serve as ‘reservoirs’ for DBPs. Importantly, reservoirs are
distinct from super-enhancers and highly overrepresented by long noncoding RNA (lncRNA)
promoters. We also observed that the human specific SVA repeat is one of the few repeat
families that had specific DBP enrichment, with a total of three DBPs specific to SVA repeats.

Looking further we found that SVA repeats reside adjacent to or within enhancers and are often
transcribed; suggesting they may have been co-opted in late primates as enhancer elements.

Overall, we demonstrate the utility of implementing data-science and reproducibility standards to
gain new insights combinations of genome-wide DNA binding events. We further note that the
design of this study was intended for didactic purposes and carried out by students in a
classroom setting.

Results

We first set out to survey the encode portal for the largest number of ChIP-seq experiments that
satisfied the following criterion: (i) target was considered a DNA binding protein (DBP), the
experiment used validated antibodies, sequencing was performed with 100bp paired end reads
and were in the same cell setting. We found the maximum number of samples that meet these
requirements were performed in K562 cells. Specifically, there are 1,076 FASTQ files comprised
of 195 DBPs meeting these criteria in K562. Rather than analyzing the peaks already called by
ENCODE for these experiments we chose to re-analyze the raw data using a community-
curated pipeline developed by “nf-core”[20]. This approach meets the highest data
reproducibility standards by using a container for all software and producing extensive
documentation at every stage of analysis within the nf-core/chipseq (v1.1.0) pipeline (Fig 1A).

**Fig 1. Framework of ChIP-seq analyses and peak calling across replicates.** (A) Schematic
of data quality requirements (2 or more replicates, 100bp reads, validated antibody) resulting in
1,076 FASTQ files representing 195 unique DNA binding proteins. FASTQs were processed
using the nf-core/chipseq pipeline (QC and peak calling). All FASTQ files passed nf-core quality
control metrics. (B) Browser view of raw data, individual replicate peak calls and our consensus
peaks. All scales are from 0 to 1 representing minimum and maximum reads in that window
using UCSC auto-scale. Peaks from individual replicates are in gray and consensus peaks
called are represented by black boxes.

The nf-core pipeline consists of documented analyses and quality control metrics that results in
significant windows or peaks of DNA binding events for each replicate[20]. After the
standardized pipeline gave us peak calls, we used this data to support our analysis and
exploration of the data. Our approach was to use R and Rmarkdown to document the analyses.
Compiling the 11 Rmarkdown files provided in the GitHub repository
(https://github.com/boulderrinnlab/CLASS_2020) will reproduce all the results and figures of this
study.

After calling significant peaks (MACS broad peak) for each replicate ChIP experiment for each
of the 195 DBPs, we wanted to develop consensus peaks across replicates. Briefly, we filtered
to peaks on canonical chromosomes and required that peaks overlap by at least 1nt in all
replicates for a given DBP. Peaks that overlap in all replicates are then merged by the union of
peak widths (Fig 1B-C). We observed five DBPs that did not have any peaks overlapping across replicates perhaps suggesting that these are promiscuous antibodies, or these proteins have heterogeneous binding across K562 cell populations (MCM2, MCM5, MCM7, NR3C1, TRIM25).

We next plotted the distribution of the number of consensus peaks for each DBP and found that many DBPs had very few peaks. In order to capture the majority of DBPs and still provide a reasonable number of peaks for analyses (e.g., permutation analyses), we chose a cutoff of 250 peaks (15% percentile, Supplemental Fig 1A). This results in 161 proteins to carry forward in the analysis and in losing the following proteins: ARNT BCLAF1 COPS2 CSDE1 DNMT1 eGFP-ETS2 FOXA1 KAT8 KDM4B MCM2 MCM5 MCM7 NCOA1 NCOA2 NCOA4 NR0B1 NR3C1 NUFIP1 PYGO2 THRA TRIM25 TRIP13 XRCC3 YBX1 YBX3 ZBTB8A ZC3H8 ZNF318 ZNF830.

Promoter centric binding properties of 161 DNA Binding Proteins

We next plotted the relationship between the number of consensus peaks observed for each DBP and how many promoters overlapped (36,814 IncRNA and mRNA promoters). We observe a linear relationship (slope = 0.31 for mRNA and IncRNA promoters) between the number of peaks and or size of peaks and the number of overlaps with promoter regions (Fig 2A). Somewhat surprising was this trend was even more pronounced when comparing overlaps within gene-bodies rather than promoter regions (Fig 2B). This suggests we could have an observation bias at promoters where promoter binding simply increases with the number of peaks observed for a given DBP and not due to preferential binding at promoters.
Fig 2. Promoter binding properties of 161 DBPs. (A) Schematic of promoter overlap strategy. Number of overlapping promoters (y-axis) per number of peaks for each DBP (x-axis). (B) Same as in (A) but for overlapping gene bodies instead of promoters. (C) Binary clustering of 161 DBPs based on promoter binding profiles (consensus peaks). Zoom out of specific regions.

To detect preferential binding at promoters, we took a permutation-based approach for each DBP’s peak-profile across the genome. Briefly, we took the consensus peaks for each DBP and randomly placed them across the genome, while controlling for (i) number of peaks, (ii) width of peaks and (iii) number of peaks on each chromosome. We then performed a Fisher exact test of the observed binding at promoters versus expected binding in the empirically derived null distributions. We observed that nearly all DBPs exhibit significant overlap with promoters versus the rest of the genome, despite being involved in many different DNA regulatory processes (Supplemental Fig 1B).

To more closely examine the results of our consensus peak strategy we performed manual inspection of samples with two or more replicates (Fig 1B-C). We find that our peaks are consistent with what would be expected of highly reproducible binding events. We see that most Pol2 and ATF3 peaks show good agreement between replicates. Interestingly in this example ATF3 is not localized to the promoter but in an upstream region that could be a newfound enhancer or upstream regulatory element. Overall, these analyses are consistent with our consensus overlap strategy representing expected and newfound features in peak size profiles.
Global analysis of similarities in binding profiles

To determine if there were underlying similarities and differences of 161 DBPs that passed our conservative filtering, we first performed hierarchical clustering (Fig 2C) on binary vectors representing binding events on 36,814 IncRNA and mRNA promoters defined in GENCODE 32 where 1 = bound, 0 = not bound for each promoter and DBP. As a quality control check, we looked for clustering of known factors. The binary vector profiles validated that POLR2A, POLR2B, and SUPT5H form a distinct cluster. Known family members, such as ATF3 and ATF2 co-cluster together as well, along with the eGFP-ATF3 control. This indicates that these DBPs had similar binding profiles with or without the eGFP tag. However, 11 cases of eGFP-tagged samples clustered together, despite having widely different functions. This may suggest that in some rare cases the tag can alter the binding profiles in a manner that is more consistent with the tag than DBP function.

As an unbiased approach to find underlying properties in DBP binding profiles, we also performed UMAP[21] dimensionality reduction for the global binding profile of each DBP (Fig 3A). Briefly, UMAP uses algebraic topology to reduce the data dimensionality. We further clustered this reduced representation using density-based clustering (HDBscan[22]). We observed a total of seven clusters. Similar to binary clustering, we identify a clear cluster of POLR2A, POLR2B, and SUPT5H and other basal transcriptional associated factors (TAF) as would be expected. This is another example of high reproducibility as POL2 has three different antibodies with 2 replicates each that are all highly concordant with thousands of peaks each.

Fig 3. Binding properties of DBPs and expression output of promoters. (A) UMAP dimensionality reduction to identify DBPs with similar promoter binding profiles. (B) Four
discrete clusters of binding patterns around promoter TSSs with 3Kb up- and downstream. Line is the average profile of all peaks in each cluster. (C) The number of peaks per DBP versus number of mRNA or IncRNA promoter overlaps. X-axis is the number of DBPs overlapping either IncRNA (red) or mRNA (black) promoters. (D) Chi-squared test for enrichment of DBPs between IncRNAs and mRNAs. The x-axis is the log2(observed over expected) and y-axis is the P-value.

Next we compared specific features of the DBP with their position in the reduced space by mapping metadata (e.g., type of DNA binding domain) onto the UMAP points (Fig 2A, Supplemental Fig 2A-D). We found no clear association with (i) type of DNA binding domain, or (ii) annotation as a transcription factor, (iii) RNA-seq expression of bound genes, or other properties. Collectively, these results recapitulate known biological functions of DBPs while including potential new factors across these different promoter regulatory functions.

**Promoter binding specificity of 161 DNA binding proteins**

We next wanted to assess the underlying promoter features associated with each DBP. Specifically, we wanted to determine where each DBP is bound relative to the TSS of 36,814 IncRNA and mRNA promoters. To this end, we generated 'binding profile plots' by calculating the read counts across all promoters centered at the TSS with 3kb flanking up- and downstream (Fig 3B, Supplemental Fig 2E-F). We next clustered the 161 DBPs based on their promoter profile plot. We split the dendrogram into clusters by ‘cut-height’ (h = 65). We observed 4 distinct clusters with at least two DBPs. The first distinction is that about half exhibit a narrow peak profile (71) and half with a broader peak profile (74). In both cases these profiles
peak near the TSS. Interestingly, 6 genes (eGFP-PTRF, ZBTB33, SMARCA5, HDGF, eGFP-ZNF512, eGFP-ZNF740) have the inverse pattern: depletion of binding at the TSS with strong enrichment at flanking regions (Supplemental Fig 2E-F).

Previous studies have identified several differences in binding features at coding (mRNA) compared to noncoding (lncRNA) promoters. Here we wanted to independently test this across 161 DBPs to determine if there was an enrichment or depletion at mRNA versus lncRNA promoters. We counted the number of lncRNA and mRNA promoter overlaps separately and observed the same linear trend of more peaks resulting in more binding events for both lncRNAs and mRNAs. However, the slope for mRNA is 0.19 (R = 0.75, P < 1e-10) and lncRNA is 0.088 (R = .87, P < 1e-10) suggesting a two-fold reduction on an average lncRNA promoter (Fig 3C).

We then performed permutation analysis (above) separately for lncRNAs and mRNAs to determine if the observed overlap is greater than expected by chance (Supplemental Fig 3C). Similar to our previous observation, nearly all DBPs were significantly (Fisher-exact P < 0.05) enriched at both lncRNA and mRNA promoters yet with a smaller magnitude of enrichment of binding events on lncRNA promoters (similarly to previously reported[17]). We observed two DBPs that were significantly depleted: BRCA1 on mRNA promoters, and ZNF507 on both lncRNA and mRNA promoters. Four DBPs showed neither enrichment or depletion at lncRNA or mRNA promoters. In total 155 of the 161 tested DBPs were enriched at lncRNA and mRNA promoters more than expected by chance (Supplemental Fig 2G).

Our previous permutation test above demonstrated that most DBPs bind both lncRNA and mRNA promoters more than expected by chance. But this approach does not account for DBPs that may prefer lncRNA or mRNA promoters. Thus, we hypothesized that some DBPs may have
a bias in binding for mRNA relative to lncRNA promoters and vice-versa. To test this, we performed a Chi-squared test to compare the number of binding events for each DBP at lncRNA versus mRNA promoters. Interestingly, although most DBPs are enriched on mRNA promoters, there were a few with a relative bias toward lncRNA promoters (P < 0.05): BRCA1, eGFP-ZNF507, EWSR1, eGFP-TSC22D4 (Fig 3D). Interestingly, BRCA1 prefers to bind outside of promoters, yet if it does bind a promoter BRCA1 prefers lncRNA over mRNA promoters.

**Repeat family and class binding preferences for 161 DNA binding proteins**

In order to determine if DBPs are enriched or depleted in TE classes and families we performed a permutation enrichment analysis. As above, we randomly shuffled peaks around the genome and calculated the number of overlaps with repeat family and classes from RepeatMasker Open-3.0 occurring by chance (Fig 4A).

**Fig 4. Many DBPs are enriched or depleted on repeat families and classes.** (A) Heat map of Z-scores of observed overlaps of each DBP versus the overlap distribution of 1,000 random permutations of each DBPs profile genome wide. Red indicates depletion and blue enrichment (negative versus positive Z-scores respectively). The observed and permuted Z-scores are for overlaps with repeat classes. (B) The same permutation analysis as in (A), but for observed versus permuted overlaps with repeat families. Red indicates depletion and blue enrichment.
We observe that some classes, such as Simple Repeats and tRNAs, were enriched for most DBPs, while others, such as the LINEs and Satellites, were depleted for most DBPs (Fig 4A). The LINE class was depleted of all DBPs with the exception of five DBPs with zinc finger-like motifs (ZNF507, ZNF316, ZNF184, ZNF24 and ZNF512). Additionally, the LTR class was depleted for most DBPs, but enriched for a subset of 23 DBPs (Fig 4B).

Overall, we found that most small TE families were not significantly enriched or depleted for specific DBPs. However, a subset of 23 DBPs were enriched in the ERV1 family, but depleted in the L1 family. These 23 DBPs are the same that were enriched in the LTR class. This is consistent with ERV1 family TEs being a part of the LTR class. Similarly, the MIR family shows a similar enrichment pattern to the tRNA family (Fig 4B). Thus, using this approach we can provide a map of which DBPs are specifically bound to which repeat family.

We did observe a subset of 6 DBPs (NUFIP1, ZC3H8, PHF21A, ARHGAP35, NCOA4, PYGO2) enriched in snRNAs, but no other TE family. Each of these DBPs, except NCOA4, contains a zinc-finger-like DNA binding domain, and a few (NUFIP1 and ZC3H8) are known to be a part of the snRNA biogenesis pathway, perhaps suggesting some form of feedback. The L1 family is depleted for almost every DBP, but is highly enriched for ZNF507, an interaction which has been previously described in an undergraduate thesis and confirmed genome-wide here (https://web.wpi.edu/Pubs/E-project/Available/E-project-042618-111020/unrestricted/MQP.pdf).
The human specific SVA repeat family has enhancer like features

Although most families are not enriched for specific DBPs, the human specific SVA repeat family is specifically enriched for three DBPs: ZBTB33, CBFA2T2, and CBFA2T3. Interestingly, all three of these DBPs are known transcriptional repressors. The SVA family is the youngest family of TEs, is enriched in gene-rich areas of the genome[23–25], and can cause human disease[24]. Based on these interesting features we further explored the binding of these factors on the SVA repeat.

We first retrieved histone modification ChIP data for K562 cells from ENCODE and visualized the coverage centered on the 5,882 SVA repeats with 5kb up- and down-stream. We find that Lysine 4 mono-methylation (K4me1) is the only histone modification enriched on SVA elements – all others were depleted (Supplemental Fig 3A). Moreover, the enrichment of K4me1 is on the 5’ end of the SVA element suggesting it could be an insulator for enhancers or part of the enhancer element. This pattern is so sharp we were concerned about mapability to the SVA element – despite observing the 5’ enrichment of K4me1. We reasoned that ZBTB33, CBFA2T2 and CBFA3T3 should be enriched across the SVA element. We performed the same analysis above for the these 3 DBPs and find there is strong mapping to these SVA regions, which suggests a low potential for the histone mark depletion to be an artifact of low mappability (Supplemental Fig 3B). We next looked at the expression level of SVA elements relative to other repeat family members. Interestingly, we observed that SVA elements have more transcription (Supplemental Fig 3C) than LTR family members that are known to function as promoters[26]. Together, these results demonstrate that the SVA region has enriched and fully mappable coverage of K4me1, ZBTB33, CBFA2T2, CBFA3T3 and are expressed.
Of the 5,882 SVA elements genome-wide, 255 SVAs were found to contain consensus peaks for all three enriched DBPs: ZBTB33, CBFA2T2, CBFA3T3. We took the same approach above for this subset of bound SVA elements. We see even stronger enrichment of K4me1 (Fig 5A) and also coverage by ZBTB33, CBFA2T2 and CBFA3T3 (Fig 5B). Interestingly, the shape and position of ZBTB33 is distinctly different than that of CBFA2T2/3T3 (Fig 5B). It suggests that ZBT33 binds on the 5’ region near K4me1 and CBFA2T2/3T3 have overlapping positions on the 3’ end of SVA elements. Closer examination of nascent, steady state RNA-sequencing (see below) and K4me1 ChIP shows a very interesting pattern of the SVA elements being transcribed and or producing bi-directional RNAs in K4me1 enriched (Fig 5C-D). This is very similar to what has been seen for enhancer regions genome wide[27,28]. Thus, the SVA transposon may have evolved (neutrally or positively) to ‘co-opt’ binding of DBPs adjacent or within enhancer regions.

**Fig 5. Human SVA repeats are enriched for DBPs and enhancer properties.** (A) Heatmap of histone modification reads centered on SVA and 5Kb up- and down-stream for the 255 SVA elements containing ZBTB33 and CBFA2T2/3T3 peaks. Here red indicates enrichment, while blue indicates depletion. Above is the average profile line of enrichment within and outside SVA elements. (B) Same as (A) but coverage of ZBTB33 and CBFA2T2/3T3. The K4me1 plot is same as in (A) for direct comparison. (C) Browser examples in the same format as Fig 1.
Promoter binding of 161 DNA binding proteins versus promoter expression output

Here we set out to investigate how binding events at individual promoters relate to the concomitant expression of the gene-product at that promoter. To this end, we analyzed ENCODE K562 total RNA sequencing data from two replicates. We calculated the average read coverage across replicates and quantified by transcripts per million reads (TPM); while considering the variance between replicates in further analyses. We first asked if the number of binding events at a promoter correlated with expression. We observed a positive correlation ($R = 0.6, P < 2.2\text{e}-16$) between the number of DBPs bound at a promoter and expression output of the promoter (Supplemental Fig 4A).

We next wanted to determine if this trend is similar for mRNA and lncRNA promoters separately. Indeed, we see that both lncRNA and mRNA promoters have a positive correlation to binding events and expression output (Fig 6A). We observed that lncRNAs have lower expression in general than mRNA as previously determined[29–32]. Yet despite these expression differences, both exhibit a positive relationship between number of binding events and promoter activity. This is consistent with observations in a previous study using a different yet overlapping subset of 73 DBP ChIP datasets[32].

Fig 6. Reservoir promoters are comprised of ghosts and zombies. (A) Number of DBPs bound to a promoter (x-axis) versus $\log_{10}$ (TPM) of transcription as measured by total RNA-seq. (B) Box plot comparing mRNA (black) and lncRNA (red) expression as a function of off, low, medium, and high expressed transcripts. Y-axis is the number of DBPs and X-axis each...
category. (C) Y-axis is the mean expression level in windows of 5 genes excluding the center
gene, with a step (slide) of 1 gene. X-axis is by category of windows containing a reservoir, non-
reservoir or super-enhancer. Y-axis is mean expression in each 5 gene window. (D) Density plot
of number of DBPs bound at a promoter at expressed (grey) versus non-expressed promoters
(red), separated by lncRNA and mRNA promoter types. (E) Nascent TPM expression (y-axis)
compared to number of DBPs bound at a promoter. (F) Density plots of DBPs ghost reservoirs
(those without nascent expression, PRO-seq TPM < 0.001) vs those with detectable nascent
expression (zombie reservoirs).

Although we saw a linear trend with binding events and expression output above, we wanted to
refine this analysis to a binned approach. Specifically, we binned lncRNA and mRNA promoters
by expression output of: Off: < 0.001 TPM, Low: (0.001,0.137] TPM, Medium: (0.137,3] TPM,
and High: >3 TPM. Interestingly, at ‘low’ and ‘off’ expressed promoters there is no difference in
binding event distributions between lncRNA and mRNA promoters (Fig 6B). Thus, they both
have similar numbers of binding events -- and can have dozens of DBPs bound -- despite
having little to no expression output. In contrast, mRNA promoters show significant increases in
binding events, compared to lncRNA promoters, at medium and high expressed promoters.
Thus, in the middle to high ranges of expression is where we begin to see the differences
between mRNA and lncRNA promoters. Collectively, these results identify over a thousand
promoters that resemble the DBP content of highly-expressed promoters yet do not have any
detectable expression by RNA-seq.
Promoters with numerous binding events but lack gene-expression output

Based on the observation of over a thousand promoters that have numerous DBPs bound, but do not produce a transcript identified by RNA-seq, we wanted to further characterize the global properties of this subset of promoters. First, we made density plots of the number of binding events at promoters. We observed a bimodal distribution of binding events where the cutoff between the distributions is around seven binding events at a promoter (54% percentile). Based on these two distinct distributions, we focus our analysis on those promoters with more than seven binding events (Supplemental 4B) and further required that the RNA-seq output was less than 0.001 TPM. This resulted in 1,362 promoters which had a relatively high number of binding events but lack of RNA-seq output from these promoters. Interestingly, 981 of the 1,362 are comprised of IncRNA promoters (Supplemental Fig 4C). This is a significant over-representation of IncRNAs in these high-binding non-expressed promoters over what would be expected by chance (hypergeometric p-value = 1.1 x 10^{-88}).

There are two trivial explanations that could explain these high binding low expression promoters: (i) these are simply super enhancer\cite{33–35} annotations (as they share similar properties of many binding events) and or (ii) the promoter is regulating a neighboring gene.

Our first concern is that super-enhancers (SE) share the similar property of many binding events, we wanted to determine how many of these regions were super enhancers. For super-enhancer annotations we used the SE-DB\cite{36} that is comprised of 331,601 super-enhancers from 542 tissues and cells, including K562. We first retrieved the SE annotations in K562 with the hg19 reference genome alignments. We then lifted over these annotations from hg19 (732
annotated SEs) to hg38. We found 714 annotations have one match to the genome and took a conservative approach of not including the 18 SEs with multi-mapping in the genome (often too many chromosomes). Of these 714 regions, 35 overlapped with the 1,363 reservoirs (P = .991 Hypergeometric). Thus, reservoirs are distinct from SE annotations and are enriched with repressor complexes unlike SEs.

Another concern is that these promoters we identified could regulate a neighboring gene; this would be most obvious for bidirectional promoters. Thus, we first defined a set of promoter types: (i) bidirectional, if another promoter on opposite strand overlaps within 1,000 bp upstream of the TSS on the opposite strand (147 / 11%); (ii) multiple nearby promoters, if there is more than one promoter on either strand within 1,000 bp (91 / 7%); (iii) nearby on same strand if there is another promoter upstream within 1,000 bp (113 / 8%); (iv) none (1,011 / 74%), if there are no promoters within 1,000 bp (Supplemental Fig 4D). Collectively, very few reservoirs had shared promoters of any type i-iii (26%), thus this cannot likely account for the lack of transcription at the observed or neighboring promoter (since there are so few). Nonetheless we calculated the TPM of promoter(s) neighboring reservoirs. We observed that 68% of these shared promoters did have a neighboring gene expressed (subcategories in Supplemental Fig 4E) for a total 240 (15%) of reservoirs that could affect neighboring gene expression. Thus, neighboring promoters of any orientation cannot account for the general lack of expression observed at reservoirs (Chi-squared p-value 2e-22).

Although bidirectional expression cannot explain why these promoters seem inert, we wanted to look more globally at the transcription environment of these promoter regions and their 5 neighboring genes. Specifically, we used a “sliding-window” approach to calculate the median TPM expression value for windows of 5 genes. Each window is centered on one gene and the mean of the neighboring four genes is calculated excluding the center gene. We first plotted the
distribution of windows where the center gene is a reservoir compared to those with non-
reservoir center genes. We also removed the 35 reservoirs that were annotated as super-
enhancers. We observed that the Wilcoxon test statistic (Fig 6C) between means was
significant (P < 9e-06), however the means were very similar (mean = 7.2 for reservoir, mean =
8.4 for non-reservoir). To be sure this is not an artifact of our permutation analysis we performed
the same analysis for windows of genes centered on super-enhancers versus non super-
enhancers. Indeed, we see that super-enhancers reside in regions of significantly higher
transcriptional activity (p < 2.5e-12) with a large fold change (4.5x) in mean expression (mean
super-enhancer = 37 TPM, mean = non super-enhancer 8.4 TPM) (Fig 6C).

Collectively, these results identify a subset of promoters that appear to be a ‘holding place’ for
DNA binding events. Thus, we will refer to these promoters as ‘reservoirs’ since they: (i) are
distinct from super-enhancer annotations; (ii) are located in more transcriptionally silenced
neighborhoods; (iii) share the property of many DNA binding properties as those promoters that
are highly expressed and (iv) have no expression output as measured by RNA sequencing.

DNA binding properties of reservoir promoters
To understand if reservoir promoters are enriched for certain DBPs, we compared the density of
DNA binding events at lncRNA and mRNA reservoir and non-reservoir promoters which had
greater than seven binding events. We observed a shift toward fewer binding events for both
lncRNA and mRNA reservoirs (Fig 6D). However, it’s notable that there are still reservoirs along
the whole range of DBP binding. Although reservoirs have fewer binding events in general, we
wanted to determine if there was enrichment of certain DBPs on reservoirs. Using a Chi-
squared test to compare the number of bound promoters for reservoirs versus non-reservoirs
we observed that 31 DBPs were depleted on reservoirs and only one gene enriched (P < 0.001
and > 2-fold depletion/enrichment, Supplemental Fig 4F). This is in contrast to the lncRNA and mRNA comparisons above where we saw global depletion of all DBPs on lncRNA promoters. Thus far, reservoirs are deviant from all trends observed for the other ~33,000 promoters tested above.

We wanted to further globally characterize reservoir promoters using UMAP dimensionality reduction as in Fig 2A. Unlike with all promoters we only observe two distinct clusters across reservoirs (Supplemental Fig 4G). However, gene-ontology analysis revealed that both clusters are strongly enriched for similar processes such as regulation of transcription (P < 1e-20).

Perhaps as expected, Pol2 and associated transcriptional machinery are some of the most significantly depleted from reservoirs; consistent with their lack of expression. Despite a global depletion of Pol II at reservoirs, we were surprised that over a quarter of reservoirs (417) had Pol II binding events, suggestive of ‘paused’ transcription. While only one DBP (eGFP-TSC22D4) reached the fold-change threshold, two more were found to be significant (P < 0.001) with small enrichments. All three are associated with repressive activity. TSC22D4 and CBFA2T2 are both known repressors while EHMT2 facilitates transcription repression through methylation of H3K9. Collectively, these findings show that reservoir promoters are distinct from super enhancers, bound by many DBPs and yet are not transcribed.

Nascent Expression and chromatin properties of reservoir promoter

Since reservoirs don’t have mature transcriptional products despite many promoter binding events, we next examined if reservoirs have “nascent” transcription detected via PRO-seq
These approaches are so precise they can identify specific DBP binding sites through PRO-seq nascent RNA read out [37,38]. Thus, we hypothesized that reservoir promoters would exhibit nascent transcription owing to so many DNA binding events. This could also be similar to more well established “paused” promoters as reviewed[39].

To determine the nascent transcription properties of reservoirs, we obtained two replicate pro-seq data sets that measure the amount of nascent transcription at a promoter. We used “Rsubread”[40] to calculate TPM values of nascent transcription across the same 6 Kb promoter window defined for DBP binding. We first plotted the relationship of nascent sequence at reservoirs versus non-reservoirs (Supplemental Fig 5A). Although statistically different (P < 3e-9) the distributions are fairly similar for reservoirs (mean = 0.41) and non-reservoirs (mean = 0.51) with a fold change of only 1.25. Thus, consistent with lack of RNA-seq expression, reservoirs also have slightly lower nascent expression than non-reservoirs (Supplemental Fig 5A). Next, we compared the relationship between the number of DBPs bound and nascent expression levels (Fig 6C). Similar to what was observed for RNA sequencing and previous studies(17,32) (Fig 3C), nascent transcription also has a significant (R = .3, P < 2e-16) positive correlation with the number of DBPs bound at that promoter (Fig 6C).

Interestingly, we observed a subset of reservoirs that have many DNA binding events but do not have nascent transcriptional activity. Specifically, we found 355 (25%) promoters with more than 7 and as many as 60 binding events that have neither nascent nor mature expression (PRO-seq TPM < 0.001, Fig 6F). We refer to these reservoirs without nascent or mature transcription as ‘ghosts’, as there is no presence of transcriptional activity. We also found 964 promoters with more than seven binding events that had no mature expression but did have nascent expression. These are referred to as ‘zombies,’ as there is some presence of activity.
We next investigated if the chromatin environment discriminates between ghost and zombie promoters. We therefore retrieved ENCODE ChIP data from K562 for a euchromatic and heterochromatic histone modification; Histone 3 Lysine 27 acetylation (H3K27ac) versus Histone 3 Lysine 27 trimethylation (H3K27me3) respectively. To this end, we downloaded peak files called in two independent replicates for each histone modification from ENCODE analysis pipelines. To validate our re-analysis of these ChIP-seq experiments we first determined if K27ac correlates and K27me3 anticorrelates with global nascent transcription as would be expected. Indeed, we see that those promoters containing K27ac have increased nascent expression (P < 2e-16, fold change = 4) (Supplemental Fig 5B). Similarly, we checked the trend for K27me3 status (Supplemental Fig 5C). As expected, we see that promoters containing K27me3 have lower nascent expression (P < 2e-16, Fold change = 0.3, Supplemental 5C).

Having validated that our analysis of PRO-seq faithfully represents known biological processes (e.g., K27ac enriched with higher expression) we wanted to zoom in only on reservoirs. We first compared K27ac status versus nascent transcription levels on reservoirs. As was seen with all promoters we see a significant difference in nascent expression between K27ac containing reservoirs and those without that mark (P < 0.0006, fold change = 1.65, Supplemental Fig 5D). Similarly, K27me3 status on reservoirs is negatively associated with nascent expression levels (P < 0.0002, fold change = 0.55, Supplemental Fig 5E). However, chromatin environment doesn’t fully explain the presence of zombie promoters, as there are promoters with and without nascent expression in each category of chromatin state.

To understand the difference between ghosts and zombies, we compared DBP binding events, the distribution of nascent transcription, and histone marks. We did not observe a significant difference in distribution of DBPs between ghosts and zombies (P = 0.064, fold change = 1.04, Fig 6F, Supplemental Fig 5F). Thus, unlike all other cases tested, the number of DNA binding
events cannot account for the difference in those that do and don’t have nascent expression. Collectively, these findings demonstrate that more than 60 DBPs bound to the same promoter do not exhibit nascent nor transcript production and are ‘ghosted by Pol II’. All properties identified above can be found in S2 Table.

Discussion

A fundamental question in biology is to understand when and where DBPs localize on a given promoter and in turn how these combinations affect expression output. Thanks to heroic efforts by ENCODE and other genome consortium efforts we now have standardized DNA binding profiles for hundreds of DBPs[12–15,31]. Moreover, these datasets go through several quality control measures before being released by ENCODE (see ENCODE portal). Thus, these important resources provide two opportunities: one for data-reproducibility standard advancements based on such well documented data; and a second to re-analyze these data-sets to find novel insights into the genome-wide localization of DNA binding proteins.

This study found a vast majority of ENCODE data to be highly reproducible -- both with known biology and in data quality. However, we do note that it may be recommended to be sure replicates have reproducible peak profiles as we observed a few ChIP-seq experiments that did not have any overlapping replicate peaks. This led us to identify 5 (2%) experiments that did not have any reproducible peaks. However, a majority of the experiments (98%) have peaks that overlap in all replicates as applied in this study. Moreover, taking into account the number of observations (promoters) it is needed to be sure there are sufficient replicable peaks called for each DBP. We found 30 more samples that had fewer than 250 peaks between replicates (14th quartile). Considering the number of observations (promoters) it is also important to be sure
there are sufficient peak numbers for permutation analysis and statistical comparisons. Finally, we noted that many of the proteins tagged with “eGFP” had similar binding profiles based on the tag and not DBP function (Fig 2C). We did not see differences in number or sizes of peaks compared to antibody-based ChIP. Yet it is surprising that 15 different DBPs all cluster together based on the “eGFP” tag despite diverse biological roles and all having similar consensus peak profiles.

These large and standardized data-sets also provide a unique opportunity to search for novel insights into the relationship of DBPs and expression output. Thus, we can compare 161 DBPs from the perspective of a promoter to determine how many bind and how this influences promoter output. Consistent with two recent studies using orthogonal datasets and approaches[17,18] we found that the more DBPs at a given promoter the more it tends to be expressed. This was similar for IncRNA and mRNA promoters alike. This analysis similarly validated these studies finding that mRNA promoters are more enriched in general than IncRNAs for DBPs[17,18].

Surprisingly, we observed 1,362 promoters had numerous DBPs (more than seven and up to 111 DBPs on one promoter) bound yet did not have expression output. In fact, these promoters had similar DBP events as the most highly expressed mRNA promoters. We termed these regions reservoirs as they seem to be a holding spot for DBPs. Notably, reservoirs are highly over-represented for IncRNA promoters relative to mRNA promoters (p < 2 e-12). We also determined that reservoirs are not super-enhancers previously defined by having many DBP binding events. Unlike super-enhancers, reservoirs have many different DBPs bound rather than many binding events of cell-specific transcription-factors in a defined region[33–36,41]. Another difference from super-enhancers is the lack of Pol II, although we do find that a quarter...
of reservoirs do have Pol II machinery bound. Perhaps suggesting that they are “paused promoters”[39,42] potentiated with up to 111 DBP binding events.

Further investigation into reservoirs revealed that almost half produced “nascent” transcription as measured by PRO-seq. This is consistent with the above hypothesis of paused promoters. What is more surprising is that half of the reservoirs also did not produce nascent transcripts within 6Kb of the TSS (ghosts). The distribution of number of DBPs was not different between poised and ghost promoters. Nor could we find enrichment of specific DBPs that separate these categories. Another possibility is that ghosts are positioned in a three-dimensional space with “DBP” hubs[43,44]. Finally, it could be that the large number of binding events at these promoters causes a ‘liquid phase state transition’ owing to so many proteins in a confined space.

Our permutation-based approach to determine if a DBP prefers a genomic feature allowed us to extend beyond promoters into the noncoding genome. Specifically, we were interested in determining if certain DBPs were specific to repetitive elements, such as transposons, across the genome. Comparing random permutation versus observed overlaps revealed something somewhat surprising: that repeat classes and families such as ‘simple-repeats’ and tRNA repeats were strongly enriched for all DBPs tested. In contrast, Line and Satellite repeats were strongly depleted for all DBPs. Thus, some repeat sequences ‘repel’ DNA binding and some ‘recruit’ DBPs without discretion.

In some cases, we did observe some interesting biases for DBPs and repeat elements. One example is the human specific repeat family ‘SVA’ as one of the newest evolving repeats in humans compared to primates. Specifically, three genes had a strong bias of binding SVA elements -- all three of which are known transcriptional repressors. Recently studies have
identified that primate specific transposons can be co-opted to generate promoters of newly evolving enhancers and even lncRNAs[26,45–47]. Thus, unlike many existing examples of co-option in the case of SVA, it could have selective pressure for binding motifs of the observed repressors and hitherto to unknown repressor motifs – or hitherto unknown promoter regulatory elements.

Collectively, this exercise in data-science, reproducibility and scale in a singular cellular context has been informative to understand relativistic promoter binding events across 161 DBPs. This has led us to understand new features of the coordination of this binding with respect to promoter expression output. Perhaps most importantly, 15 graduate students learned data-sciences and reproducibility measures that not only provide new insight into reservoir promoters but also a logical framework for future objective teaching exercises of genomic data-science.

All markdown files needed to reproduce the results and figures of this manuscript can be found here: https://github.com/boulderrinnlab/CLASS_2020.

**Materials and Methods**

**Data, Code and Markdown**

Accessions and sample information for the DBPs included in this study can be found in S1 table. All data and analyses are publicly available on our GitHub: https://github.com/boulderrinnlab/CLASS_2020.

All analyses, code, and compiled markdown are available in S1 File.
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Supporting Information

S1 Fig. (A) Distribution of number of consensus peaks observed for each DBP with cutoff at 15th percentile shown as red line. (B) Permutation analysis of DBP significance of overlapping a promoter versus 1,000 random samplings of the same peak profiles for each DBP genome wide. Showing enrichment and depletion status for DBPs (Fisher Exact P < 0.01).

S2 Fig. UMAP dimensionality reduction based on DBP binding profiles and overlaid with: (A) DNA binding domain annotations. (B) enrichment score on reservoir promoters (C) TF annotation status (D) Median RNA-seq expression level of bound promoters. (E) Examples promoter binding profile. Grey line indicates 95% confidence interval and black line is the mean value. (F) Heatmap of each promoter binding profile for individual DBPs centered at TSS. Red indicates degree of binding. Cluster of binding profiles for each DBP. The four clusters are separated by white space. (G) Enrichment for each DBP at IncRNA and mRNA promoters.
versus 1,000 random samplings of the same profiles for each DBP across the genome. Blue indicates Z-score of observed versus permuted distribution.

S3 Fig. Heatmaps as in Fig 5 for all SVA elements in the human genome. (A) Histone modifications (B) DBPs enriched at SVAs. (C) Expression of SVA elements relative to other LTR containing endogenous retroviruses (ERVs).

S4 Fig. (A) X-axis, number of DBPs bound per promoter for all promoters. Y-axis is the log_{10}(TPM) expression of resulting transcript as measured by RNA-seq. (B) Cumulative distribution of binding events on promoters. Red line indicates approximately the 50th percentile of binding events occurring at 7 DBPs bound per promoter. (C) Stacked box plots of lncRNA (red) and mRNA (black) promoters in reservoirs versus non-reservoirs. (D) Stacked box plots of promoter types in reservoir (right) versus non reservoir (left) (E) Bar plot of the 25% of reservoir promoters that have other promoters nearby. True equals a neighboring gene promoter is expressed, False is not expressed. (F) X-axis is Chi-squared test value as \log_{2}(observed/expected), Y-axis is the log10 of Chi-squared P-value. (G) UMAP reduction using only DBP binding to only reservoir promoters.

S5 Fig. (A) Density plot of nascent expression at reservoirs versus non-reservoirs. (B) Box plot of nascent expression without (left) and with (right) H3K27ac modifications. (C) Same as (B) for K27me3. (D-E) Same as (B) for reservoir versus non-reservoir promoters. (F) Boxplot of DBP distribution at ghosts versus non-ghosts.

S1 Table. Sample information for DNA binding proteins in study.

S2 Table. Promoter-level summary of DBP properties examined. Each observation (row) is a promoter and each column a variable investigated in this study.

S1 File. All scripts used to analyze the data and produce figures.
Figure 1

(A) K562 Cells + Validated Antibody + > 1 replicate + 100 bp PE → 1,076 FastQ Files

(B) POLR2A_1, POLR2A_2, POLR2A_3, POLR2A_4, Rep_1, Rep_2, Rep_3, Rep_4 → Consensus → 20 Kb

(C) POLR2B_1, POLR2B_2, SUPT5H_1, SUPT5H_2, Rep_1, Rep_2 → Consensus → 20 Kb

nf-core/chipseq v1.1.0 with Singularity for QC & MACS peak calls → Replicated peaks (188 DBPs) → 161 DBPs > 250 replicated peaks

RNA-seq, PRO-seq, K4me1, K4me3, K27ac
Figure 3
Figure 6