Variable reproducibility in genome-scale public data: A case study using ENCODE ChIP sequencing resource

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Abstract

Genome-wide data is accumulating in an unprecedented way in the public domain. Re-mining this data shows great potential to generate novel hypotheses. However this approach is dependent on the quality (technical and biological) of the underlying data. Here we performed a systematic analysis of chromatin immunoprecipitation (ChIP) sequencing data of transcription and epigenetic factors from the encyclopaedia of DNA elements (ENCODE) resource to demonstrate that about one third of conditions with replicates show low concordance between replicate peak lists. This serves as a case study to demonstrate a caveat concerning genome-wide analyses and highlights a need to validate the quality of each sample before performing further associative analyses.

1. Introduction

The ENCODE project has set up a model to generate large data to understand mammalian transcription control by pooling resources across the globe. The easy accessibility of these data, including processed files, has resulted into many researchers world-wide using the ENCODE resource to learn new biology, either by integrating existing data [1] or by integrating with their own new data [2]. An especially valuable part of the ENCODE resource concerns the identification of the binding sites of transcription factors and chromatin modifiers at genome-wide scale using chromatin immuno-precipitation (ChIP) followed by high throughput sequencing (ChIP-seq). This relies on a complex experimental setup involving antibodies, a fragmentation step, chemical or UV cross-link and library preparation from very small amount of DNA [3]. Using these data, regulators have been assigned various roles that are sometimes contradictory [4]. This suggests that despite the high technical quality control of individual samples by the ENCODE consortium [5], serving as a model for many other consortia (BLUEPRINT Epigenetics, Roadmap Epigenomics, FAANG), the post-processing of samples to determine their biological relevance is still incomplete. To investigate this issue, we performed a systematic analysis of the conditions of ChIP sequencing data with replicate experiments in the ENCODE resource.

The ENCODE consortium at UCSC provides 690 ChIP sequencing data sets for transcription factors and transcription regulators, performed using 189 different antibodies in 91 cell lines, sometimes under different cell treatments. It contains 57 conditions where ChIP-seq for the same factor in the same cell line with the same treatment (or absence of treatment) were done multiple times (between 2 and 5, for a total of 135 experiments and corresponding peak lists) and not merged. These experiments were not merged because they differ in one or more of the following: antibody used for the ChIP, laboratory that performed the experiment, or library preparation protocol for the sequencing. It is not mentioned whether these replicates give redundant information, complementary information, or if one of the replicates is of higher quality than the other(s), leaving the end user unarmed to decide which peak list should be used. Notably, these peak lists have also not been merged in other ENCODE data repositories such as encodeproject.org or factorbook.org [6]. Therefore, for each of the 57 conditions with unmerged replicates, we compared individual experiment quality using various approaches and identified 18 conditions...
where more than 50% of a peak list was irreproducible in the other experiment(s). In these cases, downstream analyses of one peak list or another can lead to different results.

2. Materials and methods

ChIP-seq experiments with replicates in the ENCODE TFBS dataset were identified and uniform TFBS peaks files for the corresponding samples were downloaded from the website http://hgdownload.cse.ucsc.edu/goldenPath/hg19/encodeDCC/wgEncodeAwgTfbsUniform/ (Uniform SPP peaks). Common and sample specific peaks were identified using bedtools [7], with a minimal overlap of one base pair. Additionally, for 6 conditions with 2 replicates, alternative peakSeq peak lists were downloaded from http://ftp.ebi.ac.uk/pub/databases/ensembl/encode/integration_data_jan2011/byDataType/peaks/jan2011/peakSeq/optimal/hub/. Bam files for the corresponding experiments were downloaded from https://www.encodeproject.org/, and merged using Samtools [8]. The mean coverages around peaks were computed using Repitools [9]. Statistically significant differentially bound regions were identified using MANorm [10] with a 150 base pair read shift. Peaks were annotated with the closest TSS using bedtools and GENCODE v21 transcripts annotations [11], after converting the annotation table into hg19 coordinates using liftOver (99.83% conversion success). De novo motif analysis was performed using HOMER [12] function findMotifsGenome.pl. Cluster analysis of all regions conversion success). Using the fraction of common peaks in each experiment as metrics to identify a set of common peaks, detected in every replicate, but missed in a given experiment ("dissimilar"), we classified the replicates (2 similar, 2 sensitive and 2 dissimilar) classified the conditions observed between replicate experiments reflect differences in ChIP enrichment signals. In a handful of exceptional cases, mean coverage at undetected peaks was very close to mean coverage at sample specific peaks (experiment 1 for Rad21 ChIP-seq in HepG2, Fig. S53C), or peaks were called in one experiment despite a low read enrichment (experiment 1 for NRSF ChIP-seq in HepG2, Fig. S36C). These exceptions might be due to potential artefacts in peak detection. Mean coverage at common peaks was almost always higher than at sample specific peaks, confirming that intersecting peak lists indeed selects peaks of higher signal to noise ratio. Notable exceptions to this are replicate 2 for CHD1 ChIP-seq in H1-hESC (Fig. S9C), c-Myc ChIP-seq in H1-hESC (Fig. S10C), Pol2 ChIP-seq in K562 (Fig. S50C) and ATF3 ChIP-seq in K562 (Fig. S55C).

Peak lists generated by another peak caller (peakSeq) for 6 replicates (2 similar, 2 sensitive and 2 dissimilar) classified the samples in the same groups (Fig. S1). This emphasizes that differences observed between replicate experiments reflect differences in raw data and they were not introduced by biased peak detection.

3. Results

3.1. A third of ENCODE conditions with replicates are of low concordance, while about a fourth has sensitivity issues

We identified 57 conditions within the ENCODE transcription and epigenetic factors ChIP sequencing data where the same experiment was done multiple times (between 2 and 5) for the same factor in the same cell line with the same treatment (or absence of treatment), and the replicates were provided without being merged. For example, the USF1 ChIP-seq in A549 cells treated with 0.02% of ethanol was performed two times by the HudsonAlpha laboratory with the same antibody but with two different library preparation protocols (the "PCR1x" and the "V0422111" protocols). Two corresponding peak lists are provided by the ENCODE ("wgEncodeAwgTfbsHaibA549Usj1V0422111Et0h02UnPnKx.narrowPeak" and "wgEncodeAwgTfbsHaibA549Usj1Pe1xEtoh02UnPnKx.narrowPeak") and both experiments are in different lanes in the UCSC track (respective experiment ID numbers 68 and 69, http://hgdownload.cse.ucsc.edu/goldenPath/hg19/encodeDCC/wgEncodeRegTfbsClustered/wgEncodeRegTfbsClustered.bed.gz).

We first compared the peak overlap between replicate experiments to identify a set of common peaks, detected in every replicate. Using the fraction of common peaks in each experiment as a measure of replicate consistency, we classified the 57 conditions in three categories (similar, sensitive and dissimilar). If the overlap between all replicate peak lists and the common peak list was 50% or more (i.e. if at least 50% of the peaks were reproducible in every replicate), then the condition was labelled as "similar" (Fig. 1A). Within the non-similar conditions, a special case was made for conditions where one peak list was at least two times bigger than another and more than 70% of the smaller peaks list overlapped with the common peak list. This was labelled as having differences of sensitivity between replicates, and the condition was classified as "sensitive" (Fig. 1B) where the differences across replicates might reflect differences in ChIP efficiency, with one experiment with a lower false positive rate, while the other a lower false negative rate. The remaining conditions were labelled as "dissimilar" (Fig. 1C). Of the 57 conditions with replicates, 26 were classified as "similar", 13 were "sensitive" and 18 were "dissimilar" (Table S1). Five of the 18 "dissimilar" conditions used different antibodies suggesting the differences might reflect undocumented isoform specificity of these antibodies. The complete analysis of overlapping peak lists for each condition is available as Figs. S4–60, and is summarized in Table S1.

In order to check whether the discrepancies between peak lists from different replicate experiments might be due to a bias in the peak detection (or peak calling) procedure instead of representing true ChIP enrichment differences, we obtained the raw read counts for each sample from the ENCODE resource. For each of the 135 experiments (from 57 conditions), we computed mean coverage at "common peaks" (detected in all replicate experiments targeting a given TF in a given cell line under a given treatment), at peaks detected in a given experiment but not in any other replicate ("sample specific peaks"), and at peaks detected in any of the other replicates, but missed in a given experiment ("undetected peaks", Fig. 1, coverage plots). For the majority of 135 experiments, the mean coverage at common and sample specific peaks was higher than at undetected peaks, demonstrating that differences in peaks indeed largely reflect differences in ChIP enrichment signals. In a handful of exceptional cases, mean coverage at undetected peaks was very close to mean coverage at sample specific peaks (experiment 1 for Rad21 ChIP-seq in HepG2, Fig. S53C), or peaks were called in one experiment despite a low read enrichment (experiment 1 for NRSF ChIP-seq in HepG2, Fig. S36C). These exceptions might be due to potential artefacts in peak detection. Mean coverage at common peaks was almost always higher than at sample specific peaks, confirming that intersecting peak lists indeed selects peaks of higher signal to noise ratio. Notable exceptions to this are replicate 2 for CHD1 ChIP-seq in H1-hESC (Fig. S9C), c-Myc ChIP-seq in H1-hESC (Fig. S10C), Pol2 ChIP-seq in K562 (Fig. S50C) and ATF3 ChIP-seq in K562 (Fig. S55C).

The "sensitive" conditions mainly represent conflicts due to a large variation between the numbers of peaks detected across replicates. We therefore systematically investigated criteria responsible for the variability in number of detected peaks. One of the main factors influencing the number of detected peaks is sequencing depth [13]. For each experiment, we thus compared the number of peaks and the number of uniquely aligned reads. Though the number of detected peaks and number of reads frequently showed a positive correlation in general (e.g. Pol2 ChIP-seq in GM12878, Figs. S43D and E), this was not the case for 3 out of 13 (23%) conditions (Fig. S2A): BHLHE40 ChIP-seq in HepG2 (Fig. 1B, number of peaks and millions of reads plots), CEBP ChIP-seq in HepG2 (Figs. S7D and E) and CREST ChIP-seq in K562 (Figs. S13D and E). We also explored if differences in signal over noise ratio (i.e. the level of enrichment obtained after the ChIP) of these ChIP-seq could explain the differences in the number of
detected peaks. The FRIP (Fraction of Reads in Peaks) metric used by the ENCODE consortium [5] is sensitive to the number of detected peaks. We therefore computed peak FPKM (Fragments per Kilo base per Million) for each experiment at common peaks detected in every replicate experiments (Fig. 1C, common peak height plot, Figs. S5–61F). For the 13 “sensitive” conditions, the relationship between the total number of peaks detected in an experiment and the median peak FPKM at common peaks was...
always positive (Fig. S2B). This analysis therefore suggests that the level of ChIP enrichment (i.e. the signal over noise ratio) is a better predictor of number of detected peaks than the sequencing depth in a ChIP-seq experiments. Lower ChIP enrichment therefore cannot practically be compensated for by increasing sequencing depth in order to detect more peaks. This also highlights the importance of normalizing data according to their signal over noise ratio, using appropriate tools, such as MAntp[10] and DiffBind[14], when comparing two peak lists from ChIP-seq of the same factor in different cell types or conditions. We applied MAntp to find statistically different bound regions between replicate experiments for 6 conditions with replicates (Fig. S1). In three (JUND ChIP-seq in HepG2, YY1 ChIP-seq in GM12878 and Pol2 ChIP-seq in GM12891) of six conditions, more than 50% of the peaks show statistically significant differential binding between replicates. For the other three conditions (RAD21 ChIP-seq in K562, CMYC ChIP-seq in Hela-S3 and PAX5 ChIP-seq in GM12878), less than 25% of the peaks show statistically significant differential binding between replicates. Thus overall (but not always), MAntp (or similar methods) can indeed help producing a robust peak list shared by all replicate experiments. However, it should be noted that false positive peaks are likely to display low number of reads in each replicate experiment, and thus are unlikely to be called differentially bound across experiments.

3.3. Data analysis and integration to determine biological validity of a sample

The promoter proximity of binding and its preference for specific sequence motifs are characteristics of many transcription factors and can be used as a proxy to establish biological validity of ChIP-seq samples. We annotated peaks with their distance to the nearest transcription start sites (TSS). This revealed differences between replicates, especially for dissimilar conditions (Fig. 1C, peak distribution plot, and Figs. S5–61G). We also performed de novo motif discovery on all 135 experiments. Out of the 18 conditions with dissimilar peak lists, 6 (33%) showed a discrepancy between the de novo motifs identified in the replicate experiments (Fig. 1C, motif logos, Fig. S3 and Table S1). This was the case for one of the 13 sensitive conditions (8%), and one of the 26 similar conditions (4%). We then systematically investigated the replicates for the dissimilar conditions to determine whether these or any other evidence put higher confidence on one or few replicate(s) over other(s). We first illustrate two cases where additional evidence put higher confidence on one or few replicate(s) over the other.

1. HDAC2 experiments in K562 cell line – histone deacetylase HDAC2 experiments in K562 cell line were generated by the Broad and HudsonAlpha laboratories using different antibodies. When comparing to other ENCODE ChIP-seq experiments, HudsonAlpha HDAC2 ChIP-seq clusters with P300 (as identified by the Sydh laboratory) while Broad HDAC2 clusters with HDAC6 (Fig. S4A). In H1-hESC cell line, HDAC2 (HudsonAlpha antibody) and P300 cluster together as well. The discrepancy between the two HDAC2 experiments is likely to be due to different antibody specificities. Wang et al. [15] identified that a cell-line specific secondary motif that mediates the binding of HDAC2 in K5622 was a GATA motif. Accordingly, the GATA motif is the top motif enriched in the HudsonAlpha sample (P value < 1e–1216), whereas the REST-NRSF motif is the top enriched motif in the Broad sample (P value < 1e–275). As GATA1 is a master regulator of erythropoiesis and K562 is a leukaemia derived cell line, the HudsonAlpha HDAC2 experiment (and antibody) appears more reliable than the Broad HDAC2 in K562.

2. CMYC experiments in H1-hESC cell line – CMYC replicate experiments in H1-hESC cell line were generated by the Stanford and UTA laboratories. Amongst H1-hESC samples, the Stanford CMYC replicate clusters together with MAX and is part of a larger cluster including RNA polymerase II and other promoter associated factors, such as TAF1 and TBP. This is in accordance with the proven role of CMYC in RNA polymerase II elongation [16]. The UTA CMYC sample in contrast does not cluster with any of the above samples, putting the biological integrity of this sample in doubt. Corroborating this, the CMYC motif is enriched in the Stanford, but not the UTA sample (Figs. S3G and H).

Although further analysis is indeed a way forward as demonstrated above, it does not always provide sufficient evidence to prefer one experiment over other for the same condition. For instance, the two CHD1 ChIP-seq experiments in H1-hESC are dissimilar (Fig. S9A). Interestingly, the “common” peaks did not select for peaks of higher coverage for one of the replicates (Fig. S9C). De novo motif discovery detected different primary (Figs. S3D and E) and secondary motifs in each experiment, but neither appears more biologically relevant than the other. When peak overlap of all CHD1 experiments in the ENCODE resource is clustered, neither of the two experiments appear closer than the other to the CHD1 ChIP-seq experiments done in GM12878 and in K562. Taken together, we can only conclude that both experiments are dissimilar, cannot be intersected or merged, and that our current knowledge is insufficient to select the most biologically relevant experiment.

4. Discussion

The ENCODE ChIP-seq data is of great value to computational and non-computational biologists alike and is widely used by the scientific community [1,2]. One great strength of this consortium is that its transparency and extensive data release policy. Taking advantage of this, we noted that the data contains several different replicate experiments for the same factor in the same cell line under the same treatment (or absence of treatment), without indication about the consistency between replicates or recommendations about which peak list to use. We performed an independent assessment of the consistency between these replicate experiments by categorizing the conditions with replicates in three groups: similar, sensitive and dissimilar. We found 18 of 57 showed a very low overlap between peak lists from replicate experiments. Assuming that a discordance is due to only one faulty sample amongst the replicate experiments, a simple extrapolation of these results to all 690 ENCODE ChIP-seq samples puts the biological validity of about 14% of them into question. To examine this, we compared the clustering of binding sites of multiple factors studied by ChIP-seq across cell types. Thirty-eight factors have been studied by ENCODE in both H1-hESC and K562 cell lines. NRSF in K562 clusters together with P300 while it does not cluster together with P300 in H1-hESC. SP1 and SP2 cluster together in K562 while SP1 shows highest peak overlap with P300 in H1-hESC cells. Due to the absence of replicates, it is impossible to know from this data alone whether these factors have a context-specific (cell type specific) binding profile, or if the observed differences are emerging from variable quality between experiments.

The practical question still remains which replicate to use or what is the best way to merge replicates. Intersecting peak lists is the most conservative approach, and retains a small number of high quality peaks. Although the most reasonable approach in most situations, it suffers a drawback: replicating a ChIP-seq experiment will effectively reduce the number of peaks detected. We therefore suggest other strategies, on a case-by-case basis (Table S1). For most conditions in the similar or sensitive category,
we suggest to use the union of peak lists. For conditions in the dis-
similar category, we used additional analyses or information to
suggest one peak list likely to be more reliable than the other(s).
For example, for the CMYC ChIP-seq in H1-hESC, we recommend
using replicate 1 based on de novo motif discovery (Figs. S1E and
F). For RNA pol2 phosphoS2 ChIP-seq in K562, we suggest replicate
2 with a higher proportion of peaks downstream of a TSS. In other
cases, we prioritized replicate(s) based on the peak overlap with
ChIP-seq of the same factor in other cell line(s) or under a different
treatment (Fig. S3).

Taken together, this study demonstrates that the high standards
set for technical quality control achieved by the ENCODE consor-
tium does not guarantee the robustness of the sample. This high-
lights a potential caveat concerning mining of the vast amounts
of data deposited in the public domain in general, where even high
technical quality of the data is not always documented.

Competing interests

The authors declare no competing interests.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in
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027.

References

[1] Lan, X., Witt, H., Katsumura, K., Ye, Z., Wang, Q., Bresnick, E.H., Farnham, P.J.
and Jin, V.X. (2012) Integration of Hi-C and ChIP-seq data reveals distinct types
of chromatin linkages. Nucleic Acids Res. 40, 7690–7704.
[2] Heidari, N., Phanstriel, D.H., He, C., Grubert, F., Jahanbani, F., Kasowski, M.,
Zhang, M.Q. and Snyder, M.P. (2014) Genome-wide map of regulatory
interactions in the human genome. Genome Res. 24, 1905–1917.
[3] Albert, I., Mavrich, T.N., Tomsho, L.P., Qi, J., Zanton, S.J., Shuster, S.C. and Pugh,
B.F. (2007) Translational and rotational settings of H2A.Z nucleosomes across
the Saccharomyces cerevisiae genome. Nature 446, 572–578.
[4] Ong, C.-T. and Corces, V.G. (2014) CTCF: an architectural protein bridging
genome topology and function. Nat. Rev. Genet. 15, 234–246.
[5] Landt, S.G., Marinov, G.K., Kundaje, A., Kheradpour, P., Pauli, F., Batzoglou, S.,
Bernstein, B.E., Bickel, P., Brown, J.B., Caying, P., Chen, Y., DeSalvo, G., Epstein,
C., Fisher-Aylor, K.L., Euskirchen, G., Gecstein, M., Gertz, J., Harremokk, A.,
Hoffman, M.M., Iyer, V.R., Jung, Y.L., Karmakar, S., Kellis, M., Kharchenko, P.V.,
L, Q., Liu, T., Liu, X.S., Ma, L., Milosavljevic, A., Myers, R.M., et al. (2012) ChIP-
seq guidelines and practices of the ENCODE and modENCODE consortia.
Genome Res. 22, 1813–1831.
[6] Wang, J., Zhuang, J., Iyer, S., Lin, X.-Y., Greven, M.C., Kim, B.-H., Moore, J., Pierce,
H. and Weng, Z. (2013) Factorbook.org: a Wiki-based database for transcription factor-binding data generated by
the ENCODE consortium. Nucleic Acids Res 41 (Database issue), D171–D176.
[7] Quinlan, A.R. and Hall, I.M. (2010) BEDTools: a flexible suite of utilities for
comparing genomic features. Bioinformatics 26, 841–842.
[8] Li, H., Handsaker, B., Wysoker, A., Fennell, T., Ruan, J., Homer, N., Marth, G.,
Abecasis, G. and Durbin, R. (2009) The sequence alignment/map format and
SAMtools. Bioinformatics 25, 2078–2079.
[9] Statham, A.L., Strimenac, D., Coolen, M.W., Struzaker, C., Clark, S.J. and Robinson,
M.D. (2010) Reptools: an R package for the analysis of enrichment-based
epigenomic data. Bioinformatics 26, 1662–1663.
[10] Shao, Z., Zhang, Y., Yuan, G.-C., Orkin, S.H. and Waxman, D.J. (2012) MNorm: a
robust model for quantitative comparison of ChIP-Seq data sets. Genome Biol.
13, R16.
[11] Harrow, J., Frankish, A., Gonzalez, J.M., Tapasani, E., Diekhans, M., Kokocinski,
F., Aken, B.L., Barrett, D., Zadissa, A., Searle, S., Barnes, I., Bignell, A., Boychenko,
V., Hunt, T., Kay, M., Mukherjee, G., Rajaian, J., Despacio-Reyes, G., Saunders,
G., Steward, C., Harte, R., Lin, M., Howald, C., Tanzer, A., Derrien, T., Churat, J.,
Walters, N., Balasubramanian, S., Pei, B., Tress, M., et al. (2012) GENCODE: the
reference human genome annotation for the ENCODE project. Genome Res. 22,
1760–1774.
[12] Heinz, S., Benner, C., Spann, N., Bertolino, E., Lin, Y.C., Laslo, P., Cheng, X.J.,
Murre, C., Singh, H. and Glass, C.K. (2010) Simple combinations of lineage-
determining transcription factors prime cis-regulatory elements required for
macrophage and B cell identities. Mol. Cell 38, 576–589.
[13] Jung, J.J., Lusquette, L.J., Ho, J.W.K., Ferrari, F., Tolstoroukov, M., Minoda, A.,
Issner, R., Epstein, C.B., Carpen, G.H., Kuroda, M.J. and Park, P.J. (2014) Impact
of sequencing depth in ChIP-seq experiments. Nucleic Acids Res. 42, e74.
[14] Ross-Innes, C.S., Stark, R., Teschendorff, A.E., Holmes, K.A., Ali, H.R., Dunning,
M.J., Brown, G.D., Gojo, O., Ellis, T.O., Green, A.R., Ali, S., Chin, S.-F., Palmieri,
C., Caldas, C. and Carroll, J.S. (2012) Differential oestrogen receptor binding is
associated with clinical outcome in breast cancer. Nature 481, 389–393.
[15] Wang, J., Zhuang, J., Iyer, S., Lin, X., Whitfield, T.W., Greven, M.C., Pierce, B.G.,
Dong, X., Kundaje, A., Cheng, Y., Rando, O.J., Birney, E., Myers, R.M., Noble, W.S.,
Snyder, M. and Weng, Z. (2012) Sequence features and chromatin structure
around the genomic regions bound by 119 human transcription factors.
Genome Res. 22, 1798–1812.
[16] Price, D.H. (2010) Regulation of RNA polymerase II elongation by c-Myc. Cell
141, 399–400.