Dissecting the genomic activity of a transcriptional regulator by the integrative analysis of omics data

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In the study of genomic regulation, strategies to integrate the data produced by Next Generation Sequencing (NGS)-based technologies in a meaningful ensemble are eagerly awaited and must continuously evolve. Here, we describe an integrative strategy for the analysis of data generated by chromatin immunoprecipitation followed by NGS which combines algorithms for data overlap, normalization and epigenetic state analysis. The performance of our strategy is illustrated by presenting the analysis of data relative to the transcriptional regulator Estrogen Receptor alpha (ERα) in MCF-7 breast cancer cells and of Glucocorticoid Receptor (GR) in A549 lung cancer cells. We went through the definition of reference cistromes for different experimental contexts, the integration of data relative to co-regulators and the overlay of chromatin states as defined by epigenetic marks in MCF-7 cells. With our strategy, we identified novel features of estrogen-independent ERα activity, including FoxM1 interaction, eRNAs transcription and a peculiar ontology of connected genes.

DNA regulatory regions represent an important part of the genome, where DNA binding Transcription Factors (TF) and a large number of co-regulators cooperate to convey cellular information and control gene activity. Recent genome-wide analyses, conducted by ENCODE and other projects in a variety of cell lines and tissues, led to the unexpected observation that distant or proximal non-promotorial regulatory regions, defined as enhancers, outnumber gene promoters by a factor of ten1. They appear to serve in a developmentally-regulated fashion, and only a fraction of them is poised or active in a defined cell type at any specific time. Enhancer activity status is quite precisely defined by histone Post Translational Modifications (PTMs), TF and coregulator binding, and enhancer RNAs (eRNAs) transcription2. The genomic activity of a TF or a coregulatory factor (namely collectively TR for Transcriptional Regulators) is studied using Chromatin immunoprecipitation (ChIP) in combination with Next Generation Sequencing (NGS). Binding sites are often taken as a proxy for the regulatory effects of TRs. However, not all binding events are functionally important3. First, the DNA-bound TR may lack a key cofactor or PTMs. Second, it has been shown that only more stable binding events are productive, as opposed to erratic, short-lived events that nonetheless are picked up by ChIP analysis4. Identifying true functional TR Binding Sites (TRBSs) has great relevance not only in regulatory genomics, but also in medical genetics and pathology5. This task can be afforded by leveraging the increasingly wide data available in public repositories concerning, in addition to TR binding, data on chromatin accessibility, histone PTMs, CpG methylation, as well as expression data by microarray and RNA-Seq technologies6. This data can be mined allowing construction of robust cistromes annotated with their activity status, finally obtaining classification of TRBS subsets with coherent functions.

Despite simple rationale, data integration is not trivial due to wide heterogeneity of the data available. The first reason is technical, since data derive from several variants of the ChIP assay or chromatin accessibility assays, or other, run on different NGS platforms at different sequencing coverages, often resulting in quite diverging numbers of binding sites. Second, data have different formats, either as raw sequencing reads or processed data including genomic coordinates (ChIP peak sets), genomic coverage (genomic signal profiles), or reads alignment files.

Thus, when integrating heterogeneous data from different studies, a robust approach is mandatory. Two major issues should be dealt with: first, how binding regions are defined; second, since measurements with ChIP are inherently not quantitative, data normalization is required. Bioinformatics tools to afford these issues exist7-12.

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but, while these tools can be successfully used for comparative analysis of ChIP data, a “start-to-end” strategy to dissect progressively a TR genomic activity by means of genomic and epigenomic data integration still awaits implementation.

A quite impressive number of studies from several labs comprising ours have reported Estrogen Receptor α (ERα, ESR1) genomic binding, ERα-controlled transcriptomes and biological effects of agonists and antagonists in human breast cancer cells\(^a\). Surprisingly though, there is no systematic analysis leading to definition of a reference cistrome and to identification of the differential activity of ERα in different experimental contexts and with different ligands or, notably, in absence of estrogen as we reported previously\(^b\) and that represents possibly one of the most puzzling activity of this TR.

We describe here a “start-to-end” strategy to define a consensus cistrome and dissect it into functional classes, by merging all genomic and epigenomic data available. This procedure, applied to ERα, led to new functional information and, applied to Glucocorticoid Receptor (GR), correctly identified experimentally validated binding sites\(^c\). Our strategy consists in a sequence of integration steps that make it flexible and usable in heterogeneous contexts for any TR of interest.

**Results**

**Dissecting transcriptional regulator cistromes by data integration.** We designed an integrative strategy to analyze heterogeneous genomic datasets, focused on the characterization of three critical aspects of the genomic activity of the TR of interest (TRI): (1) definition of binding sites that are robustly reproducible through different ChIP studies, i.e. a reference cistrome; (2) the co-factors and co-regulators that co-occupy these genomic regions; (3) the epigenetic status of TRI cistrome.

These issues are addressed in separate but converging tasks, as illustrated in Fig. 1a. Results are merged into a coherent analytical approach starting with the definition of a consensus reference cistrome for the TRI. The successive superimposition of co-factors/co-regulators ChIP genomic signal profiles, chromatin states, and other independent genomic features (e.g. transcriptional maps), lead to dissection of the cistrome into classes of TRBSs with different functional activity. In this procedure, we have applied both novel and public methods for ChIP peaks overlapping, normalization and correlation of ChIP genomic signal profiles, and unsupervised prediction of redundant patterns of epigenetic modifications (chromatin states).

The first task (blue boxes in Fig. 1a) is designed to define a TRI reference cistrome, reproducibly measured in a given experimental setting. We retrieve TRI peak sets from public repositories\(^d\) and pre-process them into high-level data structures. Then, we integrate the peak sets into a reference cistrome by progressively overlapping their genomic coordinates. A novel algorithm called RefGen, which applies a majority vote-based approach to identify a reproducible set of TRBSs according to a selected “severity” grade, has been implemented to this purpose (see Materials and Methods).

Cooperative binding of TFs and coregulators is a key feature of genomic regulatory regions. The second task (green boxes in Fig. 1a) is designed to identify TRs binding at regulatory regions defined in the first task. This is obtained by selecting and downloading appropriate datasets from the same experimental context, then by converting ChIP read alignment files into genomic signal profiles. In this analysis, we consider both the shape and the intensity of ChIP signals. Their integration is then carried out in two steps: first, signal profiles are normalized to account for the experimental differences among datasets; then, they are joined into a reference genomic profile, defined for each TR analyzed. To normalize ChIP data, we implemented a novel algorithm called NormChIP which computes a scaling factor to correct each genomic signal profile. Then a correlation between these signals and the signal profile of the TRI is computed. The use of normalized signal profiles allows comparing the genomic occupancy of multiple TRs at specific genomic regions, or at the whole cistrome level. Factors associated with TRI at the highest correlation level are the best candidates as TRI cooperating factors.

The third task (red boxes in Fig. 1a) is focus on the integration of epigenomic data. The epigenome is a pivotal regulatory layer for TF and co-regulator binding, since it reflects the accessibility and activity of chromatin regions. For the epigenetic classification of TRBSs, we collect reads alignment files of ChIP experiments of histone PTMs, histone isoforms, and relevant chromatin-associated proteins from public repositories and pre-process data into binarized genomic signals. These data are used as input for the segmentation algorithm Spectacle\(^e\) that integrates data into a discrete number of chromatin states. We use these states to deconstruct the reference cistrome following the epigenetic context in which TRI chromatin binding occurred.

Finally, the functional classes of TRBSs identified by this strategy are further improved with information derived from independent genomic and gene expression data.

**Definition of an ERα reference cistrome.** The human breast carcinoma cell line MCF-7 is the most used model system for estrogen-dependent breast cancer and was included in ENCODE Tier 2.\(^f\) The number and distribution of ERα Binding Sites (ERBSs) change drastically in response to hormones in these cells\(^g\). The majority of studies concern the genomic response to estrogen or the baseline genomic status in cells exposed chronically to low-dose estrogen. In addition, we described a dataset of ERα activity in MCF-7 cells in absence of hormones\(^h\) that is comparable to data in other datasets, when cells are “vehicle”-treated, as control.

To test the procedure for reference cistrome definition, we focused at first on the most studied condition, i.e. cells cultured continuously in low-dose estrogen (E2-constitutive) and recovered 14 ERα ChIP datasets obtained in six independent studies (Supplementary Table 1a). For each study, we identified the ERBSs detected in each biological replicate, defining a study-specific cistrome. Then, we merged the cistromes into an E2-constitutive reference ERα cistrome by selecting the ERBSs identified in at a least 75% of the studies (Fig. 1b) (see Materials and Methods for selection criterion). This cistrome is composed of 10,779 highly reproducible ERBSs (Supplementary Table 2a), whereas 23,996 were left over (dismissed ERBS). Then, we compared the properties of reference versus dismissed ERBSs (Supplementary Table 2b). Reference ERBSs were definitely more enriched in ERα Response
Element (ERE), centred in the peak sequences (43.89% vs 24.33% of dismissed) (Fig. 1c,d). Reference ERBSs displayed higher overlap with "ERα-bound active enhancer in MCF-7" 21 (9.01% vs 0.74%) and with sites of long-range chromatin interaction 1 (11.43% vs 2.08%). Furthermore, reference ERBSs were enriched in sites detected in primary tumors from patients receiving adjuvant Aromatase Inhibitors (AI) or Tamoxifen (TAM) (1.08–2.33% vs 0.02–0.07%) 22, 23, or metastases (26.86% vs 4.06%) 22 (Fig. 1d). Noteworthy, 14 dismissed ERBSs overlapped ENCODE blacklisted regions of false positive peak calling. Thus, simply applying a majority voting filtering to multiple datasets is sufficient to identify binding sites that are most likely more relevant and less erratic.

As discussed above, ERα binding to chromatin varies depending on the magnitude and duration of the estrogenic stimulus, and evidence exists that these cistromes may have different function, which has not been worked out yet. Therefore, we set out to identify context-specific cistromes, together with a wider "consensus" cistrome. We subdivided available MCF-7 datasets in three groups defined by the experimental context: (i) transient hormone deprivation (E2-Independent); (ii) 45 to 60 minutes E2 treatment (E2-Early); (iii) three to four hours E2 treatment (E2-Late); in addition to the E2-Constitutive described above. We integrated a total of 33 datasets derived from 17 studies, including our own (Supplementary Table 1a, Fig. 2a). By applying the same procedure used above, we defined four context-specific cistromes (Supplementary Table 3a–b, Supplementary Figure 1a) which comprised quite different numbers of ERBSs (Fig. 2b).

Finally, they were merged into a general ERα cistrome for MCF-7 (ERα-Ref), composed by 13,239 ERBSs. These sites present variable degree of co-occurrence in the different experimental contexts. Consequently, we...
subdivided ERα-Ref in four subsets (C1–C4), following the ERBS presence in one, two, three or all the contexts (Fig. 2b,c). 50.1% of the ERBSs (6,726) in ERα-Ref were unique to one experimental context (C1), while only 1,119 ERBSs (8.5%) were common to all the experimental contexts (C4). ERBSs that are detected by ChIP in...
all contexts may represent sites at higher affinity. Therefore, we analysed the differences in the intensity of the normalized ERα ChIP peaks, which revealed a progressive increment in the intensity of ERα binding from C1 to C4 (Supplementary Fig. 1b). Consistently, we measured an enrichment of ERE motifs (ESR1, MA0112.2) in C3–C4, explaining increased affinity of these more constantly bound sites (Supplementary Figure 1c). It should be noted, however, that these sites are not saturated: considering an independent study on ERα binding at 2, 5, 10, and 40 minutes after E2 treatment24, we observed a distinct and rapid increment of the ERα ChIP-Seq signal in C3 and C4 (Fig. 2d,e).

We next explored possible sequence differences in these co-occurrence groups. Using a /+−100 bp interval around ERBSs, we predicted higher representation of c-Fos (FOS, MA0476.1) and GATA Binding Protein 3 (TP63, MA0525.1) motifs were more enriched in C4 ERBSs (chi-square p-value < 0.001) (Supplementary Figure 1d). The motifs of well-known ERα co-factors Forkhead-box protein A1 (FoxA1) and Activator Protein 2γ (AP2γ) were enriched but equally distributed among the ERα-Ref subsets. As seen above for the E2- Constitutive ERBSs, we also evaluated the overlap of ERα-Ref with public data from breast cancer cell lines and tissues (Supplementary Table 3c). Interestingly, ERBSs previously classified as active enhancers, regions involved in long-range chromatin interactions or ERBSs identified by ChIP-Seq in tumor tissues were extensively overlapped to C4 and C3 subsets (Fig. 2f and Supplementary Figure 2a).

As far as the context-specific cistromes are concerned, a high fraction of ERBSs observed in E2-Constitutive and E2-Late contexts belonged to the C1 subset (40.5% and 28.1%, respectively) while E2-Independent and E2-Early were C4 and C3 ERBSs, suggesting that they represent the set with the highest-affinity for ERα.

Then, with our integrative analysis we defined a reference cistrome of ERα chromatin binding in MCF-7 with a joint analysis of multiple ChIP datasets and we identified the binding sites characterized by persistent receptor-chromatin interaction in hormone-deprived and treated cells.

Thus, our integrative strategy was successful in identify subsets of ERα chromatin binding sites in MCF-7 with different features.

Cofactors and coregulators overlay. To the goal of featuring factors that cooperate with ERα on chromatin, we retrieved the datasets relative to nine TFs and eight co-regulators ChIP in MCF-7 cells, in at least two of the four experimental contexts considered for the definition of the ERα-Ref (Supplementary Table 1b). A total of 128 ChIP datasets were included in this analysis. DNase-seq datasets were also included. After re-aligning the datasets, we computed the genomic signal profiles relative to the ERα-Ref regions, for each TR. These were subsequently used to compute a pairwise Pearson correlation with the ERα ChIP signal profile for each ERBS. Results showed in Fig. 2g,h for E2-Early and E2-Constitutive and in Supplementary Figure 2b for E2-Late and E2-Constitutive, report the average correlation computed in ERα-Ref subsets (C1–C4). Interestingly, we found clear differences in cofactor binding in several experimental contexts: E2-Independent ERBSs showed, uniquely, at first ranks the Forkhead box protein M1 (FoxM1), the Double-strand-break repair protein rad21 homolog (Rad21), which is a component of the cohesin complex involved in enhancer-promoter looping25, and the CBP coactivator (Fig. 2g), whereas E2-induced sites presented FoxA1, AP2γ and GATA3 at first places (Fig. 2h), as reported by many studies26–28. In this latter subset, a clear correlation with “active” ERα serine 118 phosphorylation was accompanied by the highest correlation with DNase-seq signals demonstrating increased accessibility of ERBSs upon E2 stimulation (Fig. 2h). It should be stressed that C4 ERBSs consistently showed the highest level of correlation in all the experimental contexts, as expected due to the heterogeneous nature of other subsets.

Epigenetic classification of ERα-Ref. We classified the whole MCF-7 epigenome using 41 ChIP datasets relative to six histone modifications and five regulatory proteins (Supplementary Table 1b). We predicted 15 chromatin states in three experimental contexts (Fig. 3a and Supplementary Figure 2c, Supplementary Table 4a–b). The number of datasets in the E2-late context was not sufficient to generate this classification. Focusing on the different experimental contexts were especially intriguing. Indeed, E2-Independent were predominantly classified as EnhA and Active Enhancer (EnhA), whereas other contexts were mostly classified as EnhA and TssA (Fig. 3b and Supplementary Figure 2d). Figure 3c shows that features of EnhA class in E2-Independent ERBSs are high H3K27ac and RNAPII levels. This suggests transcription at these sites. Thus, we isolated the fraction of ERα-Ref occupied by ERα in the E2-Independent context, which are mostly classified as enhancers (78.5%), specifically, EnhA (30.5%), EnhH (22.7%), and Enhancer-Weak (EnsW, 18.9%) (Fig. 3c and Supplementary Table 5a). Unexpectedly, by examining the cited experiment of GRO-seq26–31, we observed
bidirectional eRNAs transcription around *EnhT* ERBSs, even at point 0, i.e. before estrogen stimulation (Fig. 3c). Four independent public GRO-Seq experiments performed in hormone-deprived cells confirmed this finding (Supplementary Figure 3c). Thus, overlaying context-specific cistromes with epigenetic data allowed us to discover an unexpected feature of sites occupied by unliganded ERα, that is eRNAs transcription, marker of enhancer activity.

Then, our strategy can be applied to narrow down a list of TR binding sites to a subset of interaction that are predicted to be functionally relevant for their cistromic, epigenomic and TR interactions properties.

The glucocorticoid receptor cistrome of A549 cells. We evaluated the performance of our strategy on a second independent case-study concerning the Glucocorticoid Receptor (GR) cistrome in lung cancer A549 cells. Recently, a GR-ChIP-seq library was experimentally validated in reporter assays in response to Dexamethasone (DEX) treatment 16. Thus, experimental classification of GRBSs will provide ideal challenge for our integrative procedure. Following our strategy, we first integrated four GR ChIP datasets from cells treated with DEX for 1 (DEX-Early) or 3 hours (DEX-Late) (Fig. 4a, Supplementary Table 1a), defining a GR cistrome (GR-Ref) composed of 13,466 GRBSs. 5,491 (40.03%) of these were occupied by GR in both the experimental contexts (C2).
Most GRBSs were present only in one experimental context (C1) and were prevalently identified in the DEX-Late context (69.67%). Then, GR-Ref was compared to the validated GRBSs, observing that 95.6% of validated GRBSs overlapped GR-Ref. Importantly, 19.4% of C2 GRBS overlapped validated GRBSs as compared to only 3.2% of C1 GRBSs (Fig. 4d).

For the second step of our strategy, we collected 26 ChIP datasets relative to four TRs from ENCODE experiments in A549 cells treated with vehicle (DEX-Independent) or DEX for 1 hour (DEX-Early). After normalization, we calculated the correlations with GR genomic signals, and we observed a clear increase of the average correlation coefficient from DEX-Independent to DEX-Early context (Fig. 4e,f). Specifically, cAMP Responsive Element Binding protein 1 (CREB1), FoxA1 and Upstream Stimulatory Factor 1 (USF1) were highly correlated ($r > 0.6$) with GR binding signal in DEX-treated cells, while less correlation with CTCF and RNAPII was observed. Interestingly, the rank of GRBSs based on the average correlation computed for these three TFs revealed that the
top quartiles of GRBSs was also associated with the highest superimposition with the validated GRBS set (24.34% of overlapped sites) (Supplementary Table 6c). As a third step, we predicted 15 chromatin states in A549 cells (Supplementary Table 6d and Supplementary Figure 3d), by integrating 32 ChIP datasets relative to eight epigenetic modifications (Supplementary Tables 1b). In DEX-Early, most C2–C1 GRBSs demonstrated a chromatin state (Enh-TssA) shared by both enhancers and promoters (H3K4me3, H3K4me1, and H3K27ac), while only C1 were enriched in gene body marks (H3K79me2 and H3K36me3) (Fig. 4g, Supplementary Figure 3e). Considering the three more represented classes of GRBSs (Enh-TssA, Enh-Tss-Gene-5' and EnhW) we observed that validated GRBSs in the C2 subset were mostly classified as Enh-TssA and to a lesser extent as Enh-Tss-Gene-5' and EnhW (Fig. 4h).

Thus, classification of GR binding through cistrome integration, cofactor analysis and epigenetic features allows identification of functionally relevant sites.

**Discussion**

In this work, we present a strategy to guide the reuse, combination, and post-processing analysis of NGS data describing regulatory protein–chromatin interaction. Analysis of one case-study of ERα in MCF-7 cells, where extraordinarily rich data exist, and GR in A549 cells, where experimentally validated binding sites were published, confirmed the validity of this strategy. In the first case, feasibility with abundant data, i.e. computationally demanding, was verified. In the second case, the paucity of data did not hamper adherence of results to experimental validation. It should be stressed that, in the first case, this analysis provided valuable new information on the less studied experimental context - absence of hormones - which is of great interest since hormone deprivation is the therapeutic strategy of drugs as Aromatase Inhibitors in Breast Cancer. The novelty is assembling all available TEBSs, epigenomic and transcriptomic data in a coherent strategy to functionally classify chromatin binding events of any studied transcriptional regulator. Two novel methodological tools were developed and implemented to define consensus cistromes and to normalize ChIP genomic signal profiles. Moreover, a full integration approach is proposed in association with chromatin states prediction algorithms.

The variability of ChIP measurements is due to many factors, starting from the antibody used to the assay protocol and NGS platform. For cistrome definition we used an algorithm based on the majority voting approach, which allows extrapolating the consensus coordinates of TRI binding. This is a computationally efficient strategy to overlap multiple datasets that does not require a threshold based on the minimal number of overlapped nucleotides or peak centre position, allowing the analysis of NGS datasets in heterogeneous formats. Moreover, this procedure prompts easy and quick update whenever new data is available. In the case of ERα, the efficiency was evaluated by measuring consistency with functionally relevant datasets (e.g. tumors data). For GR, overlap with peaks reportedly active in luciferase reporter assay measured the performance of our analysis.

Regulatory regions are sites of binding of multiple DNA-binding or coregulatory proteins. Describing co-occupation profiles is commonly performed by superposition of genomic intervals, without considering the signal profiles obtained from NGS experiments. Here we propose to combine normalization and correlation analysis of different signal profiles. We adapted the normalization method implemented in DESeq library on a set of NGS experiments, because this method was previously observed to be effective in differential binding analysis on the normalized number of ChIP reads mapping to regulatory regions. Correlation between the normalized signal profiles of TRI and other TRs gives a measure of co-occupancy. The analysis is optimized to compare unimodal genomic signal profiles in the region of chromatin interaction. We are currently working on the extension of our method to multimodal spread signals characterizing some TR complexes. Finally, we propose the classification of TRBSs into functional classes based on redundant patterns of cistromic and epigenomic ChIP signals. Our strategy is to classify the epigenome of the experimental model system into a discrete number of chromatin states, subsequently superimposed to the TRI cistrome, whereas other integrative tools like Seqminer or EaSeq exploit the simple co-occurrence of ChIP-derived patterns, more prone to some bias. In conclusion, our strategy merges new and public algorithms into a coherent process leading to cistrome definition and classification, using extensive integration of genomic and epigenomic data, in the cell/tissue model system considered. Computational tools like EaSeq, HiChIP, Cistrome, or CisGenome assembly validated algorithms to perform restricted single-step analysis.

Considering ERα analysis, although it was hard to imagine adding value to such an extensively studied field, our strategy revealed at least two undescribed features: first, our novel classification in co-occurrence subsets (C1–C4) revealed that ERBSs common to all contexts (C4), i.e. in both presence and absence of hormones, are a peculiar subset, showing the strongest ChIP signal and the most significant co-occupancy by cofactors; they represent undoubtedly lineage-specific, highly accessible chromatin sites for ERα, and in fact they appear the only ones to quickly respond to E2 stimulation by eRNAs transcription. The second unexpected observation was that ERBSs in absence of hormones display features of active enhancers (Enh1) considering both histone PTMs and eRNAs transcription. Furthermore, transcription factor FoxM1, Steroid Receptor Coactivator 1 (SRC1) and the cohesin complex component Rad21 were quite specific to this set. While hormone-independent occupancy of ERBSs by FoxA1, AP-2α and Rad21 at ERBSs was previously demonstrated, our data show FoxM1 as the most correlated protein in hormone-deprived cells. FoxM1 is an important factor for breast cancer cell growth. Importantly, FoxM1 and ERα regulate the expression of each other.

One criticism to our ERα analysis is that the final characterization of differential functions for ERBSs is deducted based on the same kind of data by which it was classified, i.e. indirect data linked to epigenetic features and eRNAs transcription, but lacks stronger experimental proofs, such as target gene regulation. However, ERBSs are generally distant from TSS of regulated genes and rationale matching is not trivial: we are currently working to integrate HiC looping data in our pipeline. We ran preliminary analysis based on proximity using previously published RNA-seq. GREAT analysis showed that EnhT ERBSs-proximal genes are neatly linked to “gland development” and “gland morphogenesis” (Supplementary Figure 4a and Supplementary Table 5b) suggested as

\[\text{Relevant equations and figures could be included here.}\]
a specific function of ERα, whereas other Enh- classes showed more dispersed terms. Consistently, Gene Set Enrichment Analysis (GSEA) demonstrated EnhT ERBSs enrichment in several datasets related to breast cancer and estrogen response (Supplementary Table 5c). It is worth noticing that EnhT ERBSs were also significantly closer to the TSS of genes previously reported to be regulated by ERs in the E2-Independent context, especially for genes that are down-regulated following ERα ablation (Supplementary Figure 4b). Similarly, EnhT ERBSs were associated with the highest number of genes that change upon ERs silencing within 100 kbp (134 genes, 14.3%) (Supplementary Figure 4c and Supplementary Table 5d). E2-Independent EnhT ERBSs were found relatively near or intronic to several genes encoding for Tfs important for mammary gland development, i.e. SPDEF, TTFAP2C, MYB, RARA, ELF3, and the ESR1 gene itself (Supplementary Table 5e), known ERα target genes (FOS, XBP1, TFF1, EGR3) and several long noncoding RNA genes, including DSCAM-AS1, an ERα-dependent IncRNA specific to luminal breast tumors. The analysis of GRBSs was also informative. In all the steps of our procedure we observed enrichment of experimentally validated GRBSs in specific subsets selected in the different steps of our procedure. Again, it would be desirable to obtain direct proofs of gene regulation. Although GRBSs are more frequently proximal to TSS, association to regulated genes is questionable. Indicatively, one RNA-Seq experiment of DEX-treated A549 pointed to 644 responsive genes, three-fourth of them within 100 kbp from a GRBSs belonging to GR-Ref (Supplementary Table 6e–f). Most of these GRBSs co-occurred in the experimental contexts and 42.5% were classified as Enh-TssA (Supplementary Figure 4d). Despite the limited number of datasets integrated in this case, the classification reached is sound with the function.

In conclusion, results obtained in these case-studies suggest that our strategy can be applied to any TR of interest to extract novel information to be tested in experimental settings.

Methods

Datasets. To define the ERα reference cistrome for the MCF-7 cells (ERα-Ref), 33 sets of ERBSs were retrieved from GEO, Array Express, Cistrome, and supplementary material of target publications. To define the GR reference cistrome for A549 cells four sets of GRBSs were retrieved from GEO. All the analysed datasets are reported in Supplementary Table 1a. To make the genomic coordinates of all datasets comparable, they were converted to hg19/GRCh37 human genome assembly using the LiftOver algorithm. Moreover, ERBSs mapped on chromosome Y were removed.

Data of ChIP experiment against Tfs, co-regulators, histone modifications and ERα Serine 188 phosphorylation were collected from Array Express and GEO. Datasets of DNasel-Seq assays were retrieved from GEO. The data of a time-course experiment of ERα ChIP-Seq were downloaded from GSE54855. All the datasets used in the analysis are reported in Supplementary Table 1b.

Reference cistrome definition. The definition of a TR reference cistrome was performed by taking as input a list of genomic intervals corresponding to the TRBSs obtained in a set of ChIP experiments. Then, the reference cistrome is composed by the genomic positions which are shared by a desired number of experiments (√). To efficiently define this reference, an ad-hoc algorithm, namely RefGen, is proposed. In details RefGen first exploits the lists of genomic intervals to generate a genomic coverage (i.e. the intervals overlapping values for each genomic position), then the genomic position characterized by a coverage value greater than or equal to a predefined threshold τ are selected as reference cistrome. The pseudo-code of RefGen is reported in the Supplementary Note section, and its C++ implementation is available at: https://github.com/giuferreno/RefGen.

This program is free software; you can redistribute it and/or modify it under the terms of the GNU General Public License as published by the Free Software Foundation; either version 2 of the License, or (at your option) any later version.

Definition of the ERα and the GR cistromes. In the ERα case study RefGen was first applied to generate a reference cistrome from the biological replicates of 17 independent ChIP experiments (Supplementary Table 1a). In this step, for each experiment, the binding sites identified in all biological replicates were selected. Then, the resulting cistromes (i.e. one for each experiment) were divided into four subsets based on the experimental context in which they were performed: (i) transient hormone deprivation (E2-Independent); (ii) 45 to 60 minutes E2 treatment (E2-Early); (iii) three to four hours E2 treatment (E2-Late); and (iv) continuous cell growth in estrogen-enriched medium (E2-Constitutive).

For each of the four experimental contexts, the cistromes of the experiments belonging to the same experimental context were further processed by RefGen to generate a specific experimental context cistrome. A教程 value equal to 75% of the number of input cistromes was applied on these runs. The τ threshold used in these analyses are reported in Supplementary Table 3a. This threshold was selected after comparison of the number of genomic regions obtained using random datasets (Fig. 1b and Supplementary Figure 1a). Specifically, 1,000 random reference cistromes were defined for each experimental context by considering the same number of input genomic intervals with the same length. These random genomic intervals were generated using the shuffleBed function of bedtools with option -chrmon. The threshold was selected to better balance the rate of false positive/false negative predictions. A main reference cistrome (ERα-Ref) was also derived by the application of RefGen on the four context-specific cistromes. For this analysis, the binding sites identified at least in one experimental context were selected. The same procedure was applied to define the GR cistrome for each of the two analysed experimental contexts (DEX-Early, DEX-Late).

Cistromes overlap with independent genomic features from public datasets. The overlap between ERBSs and ERα GIS-ChIA-PET data from the ENCODE project was performed using the coordinates of ChIA-PET anchor regions retrieved from GSM97021217. An overlap was confirmed valid if observed for two out
of three available ChIA-PET biological replicates. The overlap between ERBSs and a list of 1,248 ERα/H3K27ac co-bound enhancers (Active Enhancers) was performed using the list provided in ref. 21.

ChIP-Seq on primary tumor biopsies from breast cancer patients taken at surgery before treatment with Aromatase Inhibitor (AI) or Tamoxifen (TAM) were downloaded from GSE4086711 and GSE3222246 respectively. Data of patients responsive or not to therapy were considered separately to define an ERα cistrome for each treatment outcome. Three sets of ERBSs defined in metastatic breast cancer samples were also considered. The cistromes were defined using RefGen by setting the τ threshold equal to the number of biological replicates available for each patient group.

Coordinates of amplified and heterozygous deleted regions in the MCF-7 were retrieved from GSE40698. The overlaps between ERBSs and these regions were considered valid if they were observed in all the available biological replicates.

The overlap between the GR cistrome and the set of DEX-Responsive GRBSs was performed by considering the set of 1,376 significant bindings sites provided in ref. 16.

**Ontological analysis.** GREAT algorithm v3.0[6] was exploited to perform the ontological analysis of the genes mapped nearby to ERBSs. Using the default settings of the program, the median distance between ERBSs and associated genes was 93,203 bp. The Gene Ontology Biological Process, Cellular Component and Molecular Function terms significantly enriched for both the binomial and the hyper-geometric by a p-value lower than 0.05 were considered.

**TF binding motif analysis.** Prediction of TF binding motifs was performed using the Centrimo algorithm of the MEME-ChIP pipeline v4.9.1[31] in default settings. A genomic region of +/-100 bp focused on each binding sites center was considered for this analysis.

**ChIP signal profiles normalization.** The normalization of ChIP signal profiles was performed with a new algorithm called NormChIP. This algorithm extends the DESeq normalization method[35] on ChIP signal profiles.

The algorithm initially encodes the ChIP signal profiles on a matrix M so that a cell M[i,j] stores the count of aligned reads in the j interval/bin of experiment i. For each row M[i, *], the NormChIP algorithm computes the geometric mean across the counts of bin j in all the experiments as reported in equation (1):

$$G_{M_{\text{med}}} = \sqrt[N]{\prod_{i=1}^{N} M[i, i]}$$  \hspace{1cm} (1)

where N is the total number of experiments.

Then, each row M[i, *] is divided by the corresponding G_{M_{\text{med}}} value obtaining a new matrix M^0. From the matrixM^0 a vector s, called size factor, is computed as reported in equation (2):

$$\text{med}[^N] M^0[i, i]$$  \hspace{1cm} (2)

where the med operator returns the median value.

Finally, the normalized ChIP signal profiles are obtained by dividing each column j of the initial matrix M with the corresponding size factor s[j].

The pseudo-code of NormChIP is reported in the Supplementary Note section and its C++ implementation is available at: https://github.com/giuferrero/NormChIP.

This program is free software; you can redistribute it and/or modify it under the terms of the GNU General Public License as published by the Free Software Foundation; either version 2 of the License, or (at your option) any later version.

NormChIP was applied to define a reference genomic signal profile for the ERα ChIP experiments. One reference was generated for each considered experimental context. The raw NGS data of the experiments selected for the definition of ERα-Ref were realigned using Bowtie v2.1.0[32] in default settings. The ERα ChIP signal profiles were computed considering a genomic window of ±5 kbp centered on each ERBS of the ERα-Ref. These regions were fractioned in consecutive non-overlapping 50-bp bins and reads aligned within each bin were counted with Seqminer[30] v1.3.3e in default settings. The genomic signal profiles were normalized using NormChIP. Then, a reference for each experimental context was defined by averaging the normalized signal profiles. The same procedure was applied on the GR ChIP-Seq experiments.

NormChIP performance was tested using five ERα ChIP signal profiles obtained by different groups in the same experimental condition (E2-Late). The datasets were selected based on the alignment rate (>90%). NormChIP normalization was compared to i) the raw signal profile and ii) the signal normalized on the number of sequenced reads (count per millions, CPM). Results of the performance test are shown in Supplementary Figure 3.

**Correlation analysis between ChIP genomic signal profiles.** TFs and co-regulators ChIP datasets were aligned with Bowtie v2.1.0[32] in default settings. The available datasets against the same factor and performed in the same experimental context were considered in the further analysis if their percentages of aligned reads were greater than 80%.

For each factor, the genomic signal profile was normalized using NormChIP. The normalized signal profiles obtained in the same experimental contexts were averaged. The resulting signal profiles were also normalized with NormChIP, with respect to different factors measured in the same experimental context. This
Chromatin states prediction. The chromatin state prediction is performed using the Spectacle algorithm19. Raw reads were aligned using Bowtie v2.1.0 in default settings. The ChIP read alignments of the histone modifications were binarized with the BinarizedBed function of Spectacle. The hg19 genome was fractioned in 200 bp non-overlapping bins. Prediction of a 15 chromatin states model, the genome segmentation and the features overlap were performed using LearnModel function with options -i = spectral, -lambda = 1 and -comb.

For the ERα case study, the MCF-7 chromatin states were predicted considering ChIP datasets against six histone marks, histone acetyltransferase p300, RNAPII, Mediator Complex subunit 1 (MED1), and CTCF. The analysis was performed separately to predict the states for E2-Independent, E2-Early and E2-Constitutive experimental contexts. For E2-Early context analysis, the data of H3K9me3 and H3K27me3 of three hours E2-treated MCF-7 were considered, since only these datasets were available at the time of the analysis. For the GR case study, the A549 chromatin states, eight histone marks and CTCF and RNAPII ChIP experiments were used to predict 15 chromatin states for the DEX-Early and DEX-Independent experimental context. Details on the criteria chosen to name the chromatin states defined by Spectacle are reported in the Supplementary Note section and in Supplementary Table 4.

The fraction of epigenomes associated with each chromatin state were overlapped with independent lists of genomic features including: coordinates of Gencode v19 gene body, TSS, Transcription End Sites (TES), CpG islands, Lamin B1 associated domains, and amplified or heterozygous deleted genomic regions. The overlap against the coordinates of different ERe-Ref (or GR-Ref) subsets was also performed. The overlap with these genomic features was computed as previously reported54. Then, enrichments were Z-score converted in order to identify the features enriched or depleted in each of the chromatin state.

Gene expression data analysis. Raw gene expression data were retrieved from public repositories without further reads quality control. Analysis of GRO-Seq datasets was performed using Bowtie v2.1.0 in default settings and the –local option. Three different experiments were considered: GSE45822, 48941324 and GSE27463. The signal profiles of these experiments were computed within a genomic region of ±5 kbp centered on each ERBS of the ERe-Ref. RNA-Seq data of hormone-deprived MCF-7 cells transfected with control or ERα-specific siRNA from GSE53532 were analyzed as previously reported44, and by considering Gencode v19 gene annotations and human genome assembly hg19.

Processed data of a RNA-Seq experiment of DEX- or Veh-treated A549 cells were retrieved from GSE79432. Differential expression analysis was performed on each gene isoforms using the DESeq. 2 R package35. A transcript was considered differently expressed if associated with an adjusted p-value < 0.05.

Gene-set enrichment analysis. The list of EnhT, EnhA and EnhW E2-Independent ERBSs associated with the differently expressed genes was defined by considering the E2-Independent ERBSs mapped within 100 kbp from the differently expressed genes TSS. The GSEA algorithm46 was used to characterize functionally the genes associated with these classes of E2-Independent ERBSs. The preRanked mode of GSEA was applied using 10,000 random permutations and selecting only the gene-sets associated with a p-value < 0.05. The genes were ranked by decreasing number of associated E2-Independent ERBSs and in case of equal number of sites, the absolute log2FC of expression in siEras-treated cells was considered. The MSigDB v4.0 gene set library was used for the analysis.

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Author Contributions

G.F., F.C., M.B., V.M. and M.D.B., designed the research; G.F. performed the data analysis; G.F., F.C. and M.B. wrote the RefGen and NormChIP algorithms; F.C., M.D.B. and G.B. supervised the project; G.F., F.C., V.M., M.B., M.D.B. and G.B. wrote the manuscript.

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