



Published in final edited form as:

Curr Genet. 2017 December ; 63(6): 1017–1021. doi:10.1007/s00294-017-0712-9.

3D clustering of co-regulated genes and its effect on gene expression

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Abstract

There are extensive long-distance chromosomal interactions in eukaryotic genomes, but to what extent these interactions affect gene expression is not clear. Recent works have identified several cases where clustering of co-regulated genes leads to enhanced gene expression in budding yeast. Similar phenomenon was also observed in mammalian cells. These results challenge widely held views of gene regulation in yeast and further our understanding of how the 3D organization of the genome contribute to gene regulation in eukaryotes.

Keywords

Long-distance gene regulation; Chromosome configuration capture; Long-distance chromosomal interaction; 3D clustering

Transcription regulation in eukaryotes is a complex process that involves coordination at multiple levels. Besides *cis*-elements (such as transcription factor binding site and TATA box) and nucleosome structure, 3D organization of chromosome also plays an important role in transcription regulation (Babu et al. 2008; Lanctot et al. 2007; Rowley and Corces 2016). The recent advances in chromosome conformation capture techniques (3C, 4C, Hi-C, etc.) allow researchers to study the spatial organization of the genome with unprecedented resolution and output (Dekker et al. 2002; Sandhu et al. 2012; van Berkum et al. 2010; Zhao et al. 2006). Application of these techniques in yeast, flies, and mammalian cells have revealed numerous intra- and inter-chromosomal interactions (Dixon et al. 2012; Duan et al. 2010; Ghavi-Helm et al. 2014; Jin et al. 2013). However, it is important to point out that these methods detect interactions based on their physical proximity, but not on their functional consequences. As a result, among the millions of interactions found in these experiments, it is hard to tell which ones carry active regulatory functions, and which ones are passive consequences of chromosome folding. In fact, computational models of yeast

chromosome as polymers with structural constraints can reproduce the DNA contact frequency measured by the Hi-C experiment to a large degree (Tjong et al. 2012; Wong et al. 2012). For mammalian cells, different cell types often yield very similar interaction maps in spite of their significantly different transcription programs (Dixon et al. 2012; Mifsud et al. 2015; Won et al. 2016). These results suggest that a large fraction of the Hi-C signals may have simple physical bases rather than regulatory roles. Therefore, identifying long-distance chromosomal interactions that regulate gene expression and understanding the underlying mechanism will be one of the main focuses of this field in the coming years.

The relation between long-distance chromosomal interaction and gene regulation is particularly intriguing in budding yeast because it is traditionally thought as a species that lacks gene regulation over long distance. In the yeast genome, regulatory regions and their targeted genes tend to be located closely within a few hundred base pairs (Erb and van Nimwegen 2011; Yan et al. 2015). In addition, by artificially placing the upstream activating sequences (UASs) further and further away from a core promoter, it was shown that UASs quickly lose their ability to activate transcription from that promoter (Dobi and Winston 2007). The idea here is that the constraint of long-distance activation in yeast is essential to ensure UAS-core promoter specificity in its compact genome (Dobi and Winston 2007). Nevertheless, Hi-C experiment in yeast revealed extensive long-distance interactions between sites >20 kb away from each other (Duan et al. 2010). The domain-like configuration of these interactions (Eser et al. 2017), as well as the interaction density (number of interactions per DNA fragment), are similar to those found in mammalian cells. More importantly, statistical analysis of the Hi-C data showed that co-regulated yeast loci tend to cluster, and physically proximal genes tend to co-express (Ben-Elazar et al. 2013; Capurso et al. 2016; Homouz and Kudlicki 2013). These observations strongly raise the possibility that some longdistance interactions play a role in gene regulation in yeast. In particular, a “gene proximity model” has been proposed that the aggregation of specific transcription factors within the nucleus space might function as a recruiter to draw their target genes close in space and probably to nearby transcription factories for coordinated expression (Li and Heermann 2013). Indeed, experimental evidence from a few recent studies support this idea.

The first line of evidence comes from the interaction and regulatory effect between homologous alleles in somatic diploid yeast cells (Fig. 1a). Pairing between homologous chromosomes has been observed in diploid budding yeast for over a decade (Burgess et al. 1999), but its functional significance in gene expression has begun to unravel in recent years only. The studies so far all use the classic activation system, *GALI* promoter, as the model. It was first found that β -estradiol-induced *GALI* promoters at allelic locations could form significantly stronger transinteractions under the activating condition (Mirkin et al. 2013), suggesting that this interaction is intricately related to gene expression. Extending from this observation, Zhang et al. showed that two allelic reporters, one driven by wild-type *GALI* promoter and the other by a mutated *GALI* promoter with delayed response to galactose induction, physically associate upon induction. The wild-type *GALI**pr* triggers synchronized firing of the defective promoter and accelerates its activation without affecting its steady-state expression level (Zhang and Bai 2016). Importantly, the same reporters located at non-allelic locations do not show such interaction and trans-regulatory effect. Brickner et al. also

reported that the wild-type *GALI-10* alleles in diploid yeast cells cluster upon induction, and a *cis*-element in the *GALI-10* promoter, GRS4, is critical for promoting the interaction. Again, this clustering contributes to stronger expression of *GALI* and *GALI10* by increasing the fraction of cells that respond to the inducer (Brickner et al. 2016).

The second line of evidence comes from a recent study of the interaction between non-homologous loci and their function in gene regulation. Du et al. developed a medium-throughput assay to screen for functional long-distance interactions that affect the expression of a reporter gene in the budding yeast genome (Du et al. 2017). An insulated *MET3* promoter flanked by ~1 kb invariable sequences was integrated into thousands of genomic loci, allowing it to make contacts with different parts of the genome. The idea is that, if the *MET3* promoter activity changes, it has to be caused by mechanisms that initiate more than 1 kb away. Their data suggest that a subset of *MET3* co-regulated genes on different chromosomes can physically associate and form 3D clusters, and the activity of the *MET3* promoter increases when inserted near these genes (Fig. 1b). The same phenomenon was also observed for *MET13*, an endogenous gene in the cluster. When translocated to a different genomic locus, *MET13* loses the interactions with other genes in the cluster and shows lower expression, indicating that the endogenous genes also benefit from the cluster for higher activity.

Although the studies above focus on different types of interactions, three common themes emerged. First, co-regulated genes can interact with each other, but only when they are located in certain genomic loci. For example, among the insertion sites tested, *GALI* reporters only interact when they are at allelic locations, or on non-homologous chromosomes but have equal distance to centromere (Mirkin et al. 2013; Zhang and Bai 2016). Similarly, only a small fraction of Met4-targeted genes seems to cluster, and the *MET3* reporter makes contacts with the cluster only when inserted into certain loci (Du et al. 2017). These results indicate that the search for interaction partners is constrained to a nuclear sub-volume imposed by the chromosome context (Noordermeer et al. 2011). Second, the intensity of the interaction changes with transcriptional status. For both *GALI* and *MET3* reporters, the interaction becomes stronger under the activating condition, suggesting that the interactions may be mediated by transcription-related proteins or RNA transcripts. In fission yeast, it was proposed that condensin is used to connect actively transcribed genes (Iwasaki and Noma 2016; Robellet et al. 2016). Third, genes at the cluster show higher expression. The detailed mechanism is not clear, but a simple model is that the clustering may generate a “transcriptional hotspot” with high local concentration of related factors, allowing co-regulated neighboring genes to fire with more strength. Also, factors can quickly bind and rebind among these spatially co-localized genes, making the transcription process more efficient.

Clustering of co-regulated genes has also been observed in the mammalian system, and sometimes referred to as “multigene complexes.” Genomic loci from multigene complexes were shown to associate with Pol II foci or “specialized transcription factories,” suggesting that it may provide a structural framework for co-transcription (Li et al. 2012; Papantonis et al. 2012; Schoenfelder et al. 2010). Consistent with this idea, a few studies have demonstrated that the formation of multigene complexes coincides with alterations in gene

expression (Apostolou and Thanos 2008; Fullwood et al. 2009; Sandhu et al. 2012; Spilianakis et al. 2005). For example, during the differentiation of naïve T cells to effector T-helper cells, there is a drastic change of chromosomal interactions made by two loci, *Ifng* and *T_H2* LCR, and this switch is thought to be critical for establishing the T-helper cell identity (Spilianakis et al. 2005). Most of the studies showed correlations between chromosomal interaction and gene expression, but some works also investigated the causal relationship between them. In Fanucchi et al., for instance, they perturbed the site of contact in the NF κ B-regulated multigene complex, and showed that it reduces the transcription of other interacting genes (Fanucchi et al. 2013). Interestingly, some genes in the complex play more “dominant” roles, and their transcriptions are required for other members to interact and express (Fig. 1c). Collectively, these results show that transcriptional co-association is a wide-spread phenomenon that occurs in many transcriptional programs and has regulatory functions.

Despite recent progress, many important questions about the 3D clustering and transcription regulation require further elucidation. Do all regulons of different transcription factors experience 3D clustering? Among one regulon, how many genes come together? What is special about these genes? What factors are mediating the clustering? At the cluster, how are different genes arranged and orientated? Is there significant cell-to-cell variability of these clusters? Is there a causal relationship between clustering and enhanced gene activity? What is the molecular mechanism underlying the enhanced gene activity? To address these questions, we need more efficient methods to map the 3D localization of all the regulons in the genome, preferably in a more targeted fashion. We also need new methods to selectively perturb the chromosome configuration and examine the biological consequences.

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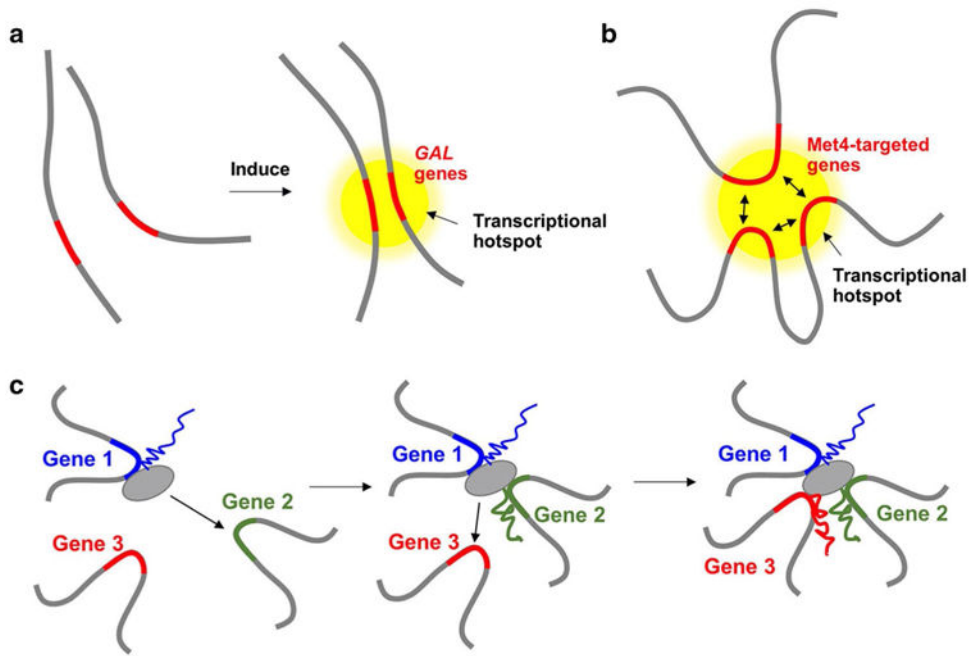


Fig. 1.

3D clustering of co-regulated genes. **a** Two copies of *GAL1* genes (*red*) at allelic locations in diploid yeast cluster together in a transcriptional hotspot (*yellow circle*) upon galactose induction. **b** A fraction of Met4-targeted genes (*red*) form 3D cluster in a transcriptional hotspot. *Double arrows* indicate the physical interactions between different genes in the cluster. **c** Three NF κ B-regulated genes (Gene 1, 2, and 3 shown in *blue*, *green*, and *red*, respectively) form a multigene complex with hierarchy: transcription of Gene 1 is required for the recruitment and expression of Gene 2, and is then followed by the recruitment and expression of Gene 3. *Gray circles* represent the transcription machineries