A Dynamical Model Reveals Gene Colocalizations in Nucleus

Jing Kang¹,², Bing Xu³, Conor Hennessy¹, Peter Fraser¹*, Jianfeng Feng³,²*

¹ Laboratory of Chromatin and Gene Expression, the Babraham Institute, Cambridge CB22 3AT, UK

² Centre for Scientific Computing, Warwick University, Coventry CV4 7AL, UK

³ Centre for Computational Systems Biology, Fudan University, Shanghai, PR China

*Corresponding authors: peter.fraser@bbsrc.ac.uk, jianfeng.feng@warwick.ac.uk

JK and BX contribute equally.
Abstract

Colocalization of networks of genes in the nucleus is thought to play an important role in determining gene expression patterns. Based upon experimental data, we built a dynamical model to test whether pure diffusion could account for the observed colocalization of genes, within a defined subnuclear region. A simple standard Brownian motion model in two and three dimensions shows that preferential colocalization is possible in a wide parameter range and suggests the occurrence is mainly due to the limited number of available transcription factors. Experimental data of chromatin movements demonstrates that the fractional rather than the standard Brownian motion is more appropriate to model gene mobilizations, and we tested our dynamical model against recent static experimental data, using sub-diffusion process by which the genes tend to colocalize more easily. Moreover, in order to compare our model with recently obtained experimental data, we studied the association level between genes and factors, and exhibit the validation of this dynamic model. Finally, our model further demonstrates that increasing transcription factor number, rather than factory number and nucleus size, might be the key mechanism to prevent gene colocalization.
Author Summary

Transcription is a fundamental step in gene expression, yet it remains poorly understood at cellular level. Transcription requires a variety of transcription factors binding to promoters to switch on transcription, as well as RNA Polymerases which catalyze the transcription of DNA to synthesize precursors of mRNA. Textbooks are full of description that promoter bound transcription factors recruiting RNA polymerase, which initiates transcription before sliding along the transcription unit. However, increasing evidence shows that it is the DNA template bound with transcription factor that slides through the immobile RNA polymerase in discrete nuclear sites (known as transcription factories), rather than RNA polymerase sliding along DNA template.

Based on this transcription factory model, we build a virtual space in which genes and transcription factors move randomly while transcription factories are immobile. We find that under a large number of parameter range (number of genes, factors, and factory and nucleus size), this simple dynamical model are valid for a number of experimental observations. Moreover, we suggest the occurrence of gene colocalization might be mainly due to limited numbers of transcription factors, rather than other factors such as nucleus size or transcription factory number. This work offers insight into the general principles of regulation of transcription and gene expression by simulating the translocation of transcriptional units (genes and transcription factors) using purely random diffusion process.
Introduction

A central theme in the regulation of transcription is the binding of transcription factor proteins to specific sites along the DNA. Though these sites can be several tens or hundreds of kilobases from a target gene promoter, regulation is achieved by the formation of chromatin loops that bring the sites together to form transcriptional hubs. It is thought that proximity between distal regulatory elements and their target genes increases the local concentration of specific regulatory factors to affect transcriptional control. Recent studies have also shown that active genes colocalize in the nuclear space at focal concentrations of the active form of RNA Polymerase II (RNAPII) called transcription factories [1,2,3,4,5,6]. A genome-wide enhanced 4C (e4C) screen demonstrated that specific combinations of genes from different chromosomes share factories with a high frequency, suggesting that active genes have preferred transcription partners. Colocalization of these spatial gene networks at transcription factories was found to be dependent on the transcription factor Klf1, which co-regulates many of the partners. Just as distal regulatory elements affect gene regulation by spatial clustering, intra- and inter-chromosomal associations between co-regulated genes may affect expression by creating specialized hotspots that are optimized for their transcription. Thus, the transcriptional program of a cell may be reflected by, or may even dependent upon, the spatial organization of the genome. The appreciation that a very large proportion of the genome is transcribed with relatively few transcription sites suggests that the organization of the transcriptional machinery plays a major role in shaping the nuclear organization of the genome. The positioning of genes, regulatory sequences and transcription factors in relation to each other and to landmarks in the nucleus, such as nuclear bodies and lamina, are important determinants in gene expression [7].

Little work has assessed the role of transcriptional factories in developmental switches in gene expression. The emerging experimental evidence has shown that transcription factors may bind the active genes in the region of transcription factories [8], but how these transcription factors and active genes position and reach to the factories and how they bind together are still unknown. Gaining an understanding of the emergence of complex spatiotemporal patterns of behavior from the interactions between genes in a regulatory network poses a huge scientific challenge with potentially high industrial pay-offs [4,9,10]. Experimental techniques to dissect regulatory interactions on the molecular level are critical to this end. In addition to experimental tools, mathematical modeling and computer tools will be indispensable. As most genetic regulatory systems of interest
involve many genes connected through interlocking feedback loops, an intuitive understanding of their behavior is hard to obtain. By explicating hypotheses on the topology of a regulatory network in the form of a computer model, the behavior of possibly large and complex regulatory systems can be predicted and explained in a systematic way. One of such recent examples is described in Misteli [11] and Rajapakse et al [12], where the authors developed a model based upon self-organization. It is probably the most successful model in the area, as confirmed in Misteli [11]. However, the model is phenomenological with an oversimplified system of Kuramoto oscillators and the random effect is largely ignored.

Here we have developed a model based upon known experimental data, aiming to account for experimental results and can be used for predicting and guiding further experiments. Colocalization ratio is introduced to characterize the gene colocalizations in transcription factories. Within a wide parameter region, we demonstrate that gene colocalization is plausible for both two and three dimensional cases. Experimental data tells us that sub-diffusion is observed in various cell-cycle phases (S and G phase) in yeast [13], which implies that fractional Brownian might be required to model gene movement, at least locally in time and space. Using fractional Brownian motion, gene and gene pairs colocalized with transcription factors are estimated, and tested against with experimental data obtained from RNA-immuno-FISH experiments. We find that the model, albeit simple, can account for observed experimental data.

Previous research [1] showed that mouse embryonic fibroblasts (MEFs) of a flattened nucleus had relatively higher number of transcription factories (2,000-2,5000) while in erythroid cells (such as E10, E14, AS, Sp, Th, and Br, which are embryonic blood, fetal liver erythroid, adult anemic spleen erythroid, normal adult spleen , adult thymus , and fetal brain, respectively) of a spherical nucleus, a relatively lower number of transcription factories (100-300) were observed. Therefore, one may wonder the effects of the number of transcription factories and the shape of the nucleus on the level of gene colocalization. Our simulation using both flattened and spherical nucleus, showed that colocalization is not very sensitive to the number of transcription factories, but is sensitive to the number of transcription factors.
Results

Colocalization between Genes

Recent studies show that active genes dynamically colocalize to shared transcription sites and that specific networks of genes share factories at high frequencies [1,14]. We built a model by randomizing the movement of genes and transcription factors, with a defined number of immobile transcription factories [6]. Live cell studies have shown that chromatin is highly mobile but regionally constrained within eukaryotic nuclei. We therefore created a defined space for random diffusion of genes (genes restricted to a square in the 2D model or a cubic in the 3D model) based on the observed mobility of chromatin in vivo. Each gene regulatory element is regarded as a point for simplicity, rather than as a polymer. The transcription process is simulated as follows: when a gene and its transcription factor come into close proximity, they bind and diffuse together for a limited time (which is called binding time). If the bound complex encounters a transcription factory before their binding time elapses, the gene engages with the polymerase and becomes active, remaining associated with the transcription factory until finished (the transcription time); if the gene-factor complex does not encounter a transcription factory during the binding time, the gene and factor dissociate and continue their brownian motion separately. During a productive transcriptional event, the transcription factor may be released from the gene before it finishes transcription and is available to randomly interact and bind another gene of the same family.

We examined the behavior between genes of the same family (X genes) and two different families of genes (X genes and Y genes) (see Fig. 1A). Genes and their corresponding factors (X factors and Y factors) are allowed to move randomly within the restricted region. However, to simulate the constrained diffusion of chromatin, the genes cannot exit the defined space. Transcription factors are allowed to exit the space, but exit of one factor is followed by entry of an identical factor on the opposite/same side of the space (We have simulated both cases and found no significant difference in the simulation). This maintains the concentration of factors within the space and simulates the observed behavior of factors to explore the entire nucleus. The number of genes and factors for each family are equivalent. X factors can only bind to X genes and Y factors to Y genes, and the two families of genes and factors are independent of each other. We also assume that each gene has only two states (Fig. 1B and 1C) — either it is being transcribed (X gene: $u(t) = 1$; Y gene: $v(t) = 1$) or not being transcribed (X gene: $u(t) = 0$; Y gene: $v(t) =$.
Fig. 1B and 1C illustrate X-X gene colocalization and X-Y gene colocalization, respectively, where \( u(t) \) is the state function of X gene and \( v(t) \) the state function of Y gene.

*Standard Brownian motion: 2D model*

We start with the 2D model using standard Brownian motion to simulate the translocation of genes and factors (simulation parameters are presented in Table I). All parameters were obtained from experimental data \([4,15]\). When a gene engages with a factory for a productive transcription event, the presence of other genes transcribing in that factory results in a colocalization. The simulation is run until both X-X gene colocalization events and X-Y gene colocalization events have happened over 5000 times, for each set of parameters. As a result, the colocalization ratio (Eq. (13)) becomes rapidly stable with a small fluctuation over long simulation time (Fig. 2A), as pointed out in the Methods section. The parameter values in Fig. 2A are specified to be 40 X genes, 40 Y genes, 5 X factors, 5 Y factors, and 10 sec binding time. The yellow triangles are the colocalization ratio obtained from Eq. (15), which used the inter-colocalization interval for the calculation rather than the counting of events. It shows that the two methods (Eq. (13) and Eq. (15)) for calculation of the colocalization ratio are equivalent.

In Fig. 2B, the histogram of inter-colocalization interval \( T_{C,xx} \) and \( T_{C,xy} \) is plotted for X-X and X-Y gene colocalization, which can be well fitted by gamma distributions. The distribution of X-X gene inter-colocalization interval \( T_{C,xx} \) is much narrower than \( T_{C,xy} \), showing that X-X gene colocalization events are more likely to happen over time. This means colocalization is more often observed between genes of the same family. Fig. 2C further demonstrates the asymmetry of colocalization for homogeneous and heterogeneous families of genes, by pseudocolor plotting the mean values of inter-colocalization interval \( ET_{c,xx} \) and \( ET_{c,xy} \) for various parameters (4-11 factors and 5-100 seconds binding). The mean of inter-colocalization interval of genes from the same family \( T_{C,xx} \) are generally smaller compared with that from different gene families \( T_{C,xy} \) for each parameter set, indicating the same family of gene tends to colocalization more often. It is also observed that the inter-colocalization interval increases dramatically as the binding time increases, for both cases, when the number of factors is fixed.

Moreover, Fig. 2D demonstrates the probability of gene-gene colocalization in the 2D case for one specific parameter combination (7 factors, 1sec binding time). The
colocalization probability can be understood as the chance when a gene starts to transcribe that another gene is already being transcribed in the same factory. The histograms of colocalization probability between X-X genes (Eq. (18), pink) and X-Y genes (Eq. (19), blue) show that X-X genes colocalize preferentially, compared to X-Y genes, because the chance for X-X gene colocalization are clearly higher than that of X-Y gene (i.e., \( r_2(x_1, x_2 | x_1) = 0.033 > r_2(x_1, y_1 | x_1) = 0.02 \), see Methods section). Finally, Fig. 2E illustrates the colocalization ratio \( r_{xx/xy} \) (Eq. (13)) for various parameters, calculated from counting of colocalization events for X-X gene and X-Y gene. The red dashed line indicates the classification threshold for colocalization region where \( r_{xx/xy} > 0.52 \) (region above). Interestingly, although the model is based on completely random movements of genes and factors, there is a region of parameters in which the system behaves non-randomly, where X genes tend to colocalize preferentially with other X genes.

The interesting question is where the asymmetry (i.e., tendency for genes of the same family to colocalize) comes from. Consider an extreme case in which the number of transcription factors for each family is severely limited (i.e., one X factor and one Y factor). In this case, only family member genes in the immediate vicinity of their factor have a chance of transcribing and colocalization of the family member genes in the same factory would have a high probability. On the other hand, if there were enough factors to bind every gene simultaneously, gene transcription events would be totally independent and a random distribution of factory sharing would be expected. Therefore, for a fixed number of factories, the colocalization ratio is a decreasing function of the number of transcription factors, as indicated in Fig. 2E. Hence, limited resources (finite number of transcription factors) could be the main reason for colocalization in the simulation.

*Standard Brownian motion: 3D Model*

Little information is available about how gene regulatory components are organized within the three-dimensional space of the nucleus from experiments. Our 3D model can provide a temporal-spatio simulation of the translocation of genes and transcription factors, and their interactions with factories. Video demonstration for genes and factor translocation, binding process and transcription in the 3D case (when the locations of transcription factories are fixed) is available from our group’s website (www.dcs.warwick.ac.uk/~feng/gene-factory). Fig. 3A illustrates the positioning of each transcription element at a specific time \( t \) (big green dots represent the location of each factory, asterisks are the transcription factors, and genes the small spots. Red and blue
represent different family of genes and factors). The parameters of 3D simulation can be found in Table I. Most of the parameters are the same as the 2D case, except the genes and factors translocate randomly in a cubic space rather than a square plane (Fig. 3A).

Fig. 3B-E are obtained similarly as Fig. 2B-E. Interestingly, in comparison with the 2D case, the area of the colocalization ratio is considerably enlarged (Fig. 3E). This could be easily understood from the dynamics of the 3D Brownian motion. The main reason for the bias (colocalization) is again due to the limited number of transcription factors, as discussed in the 2D case previously. With Brownian motion in the 3D cube, it is more difficult for genes and factors to collide and combine with each other, and then engage with a factory compared to the 2D square. Hence the probability that transcription factors of two different families cluster in the same factory is lower. This explains why the colocalization regions in Fig. 3E are larger than Fig. 2E.

**Fractional Brownian motion**

Is standard Brownian motion good enough to simulate preferential colocalization in nucleus? Experimental data of chromatin movements reported in yeast cells in Sage et al [13], Gasser [16] and Heun et al [17] demonstrates that fractional rather than standard Brownian motion may be more appropriate to model gene mobilizations. In S and G1 phase of the cell cycle, the trajectory of the chromatin movement is negatively correlated (one long increment followed by a short one), and therefore fractional Brownian motion is a sub-diffusion process (see Methods section). This means the trajectory of the chromatin movement is more localized, enabling the genes to colocalize more easily.

With the information from experiments, we expect that the fractional Brownian motions play an important role in the nuclear dynamics. To test the hypothesis, we ran simulations in 2D using fractional Brownian motion to simulate the translocation of genes and factors. Fig. 4A is the inter-colocalization interval histogram for the sub- and super-diffusion with 5 factors and 10 seconds binding time. Clearly, the inter-colocalization interval of sub-diffusion ($H = 0.1$) is much smaller than super-diffusion ($H = 0.9$), indicating that colocalization between members of the same family of gene occurs more readily for sub-diffusion ($H = 0.1$, as $T_{C,xx} < T_{C,xy}$ for both cases). Intuitively, the target hitting time for sub-diffusion movement should be larger than super-diffusion, but for sub-diffusion, the particle will stay locally and hit the target once again more easily. This may explain why sub-diffusion is preferred in gene translocation biologically (as indicated from the data
obtained from [13]). Fig. 4B shows the colocalization region for sub-diffusion \((H = 0.1)\) and super-diffusion \((H = 0.9)\) movement. It clearly reveals that the colocalization region is very large for sub-diffusion as the colocalization ratio is always above the threshold 0.52, but for super-diffusion, colocalization is hardly observed.

**Colocalization (association level) between genes and factors**

All results above tell us that there exist colocalization regions between genes; even though the model is set up completely symmetric (equal number of genes and factors for each family). In order to further validate our model and make our simulation results comparable with recently obtained experimental data (Fig. 5A), we studied the gene-factor colocalization (also called association level). Here, we say that genes and factors are associated with each other as long as they are both presented in the same factory, and it is not necessary for the gene to bind the factor, consistent with the measurement used in experiments.

Schoenfelder et al [14] reported the intra- and inter-chromosomal colocalization frequencies of 33 mouse genes relative to the Hbb and Hba globin genes in erythroid tissues (Fig. 5A). Gene regulated by the transcription factory Klf1 preferentially clustered in factories containing high levels of Klf1. Fig. 5Aa shows the spatial distribution of transcription factor Klf1 (Kruppel like factor) relative to RNAPII factories by immunofluorescence in mouse erythroid cells. The data exhibits nearly all nuclear Klf1 foci overlapped with RNAPII-S5P foci, indicating 10 - 20% of transcription factories contain high levels of Klf1. Therefore, we restrict 20% of Klf1 associated level with RNAPII (as the background association level) by selecting the factor number and binding time in our simulation (Fig. 5B). Fig. 5Ab is the double-label RNA immune-FISH of nascent transcripts (Hbb, green) and Klf1 foci (red). This image shows the positions of transcriptionally active, Klf1-regulated gene (e.g., Hbb) relative to Klf1 foci. It is found that majority (59% - 72%) of actively transcribed alleles of Hbb, Hba, Hmbs and Epb4.9 (regard as X genes) were preferentially associated with Klf1 transcription factories. Cpox genes (regard as Y gene) associate with Klf1 factories at marginally higher frequencies (26%) than expected by a purely random distribution. For actively transcribed alleles of the Klf1-independent Tubb5 and Hist1 genes (regard as Z genes), they show no preferential localization to Klf1-containing factories (20%). Hence, we regard Klf1 as the
X factor, and the X gene - X factor association level is estimated to be around 64% in experiments, while Y gene – X factor association level is around 20% from this experiment, matching the Klf1 background association level (20%). Therefore, we understand that X factors and Y genes are independent to each other. Fig. 5Ac is the triple-label RNA immune-FISH for pairs of nascent transcripts (Hbb and Hist1, blue and green, respectively) and Klf1 foci (red). From experimental observation, this colocalizing pair of genes relative to Klf1 foci reveal that colocalizing pairs of Klf1-regulated genes are associated with Klf1 transcription factories at very high frequencies (63-79%), and the colocalizing Klf1-independent gene pairs show no preferential association with Klf1 transcription factories.

In simulation, we calculated the association level between X gene (Hbb, Hba, Hmbs, and Epb4.9) and X transcription factor (Klf1), the association level between Y gene (Cpox) and X factor (Klf1), when confining the Klf1 background association level as 20%. Using sub-diffusion ($H = 0.4$) to simulate the translocation of regulatory elements, we calculated the X gene – X factor association level (Eq. (25)) both in 2D and 3D, by fixing gene number but varying factor number and binding time (Fig. 5Ba), or by fixing factor number but varying gene number and binding time (Fig. 5Bb). It is clearly shown that the experimental data can be well matched with our model with one set of parameters (gene number 5, factor number 2, and binding time 130 s) in 3D case. Next we examine the colocalization between a pair of genes (X genes, Y genes or Z genes) and Klf1 (X factor). Our simulation result shows that the colocalization of paired X genes with Klf1 is 0.8, and paired X-Y genes with Klf1 is 0.2, again in agreement with experiments [14]. All experimental results and our simulation results except for Z genes are summarized in Table II.

However, experimental data shows that X-Z gene pairs (Hbb/Tubb5, Hba/Tubb5, Hbb/Hist1, Hba/Hist1) are not associated with Klf1 factories. Therefore, we need to understand what mechanism the transcription of Z genes is governed. The simplest possibility is that either there are interactions between X gene transcription factors and Z gene transcription factors or X genes and Z genes themselves. In other words, Z genes (or Z factors) might be negatively correlated with X genes (or X factors), while Y genes and X genes are independent. To assess this, we run simulations with the following exclusive rules: if an X gene (factor) is in a factory, it will prevent the entry of a Z gene (factor) with a probability $p$. Essentially, this introduces a negative correlation between factors or
genes. In factor case, it simply implies that Z gene is co-regulated by X and Z factors. The simulation results on the gene-factor association level (Fig. 5Bc) did not show much difference after including the preventing probability among genes or factors, and it is not easy to simultaneously fit the experimental data which implies that Z gene – X factor association level as 0.2, indicating the negative correlation between different families of genes (factors) might not be the primary reason for different values of Klf1 association rate among different families of genes, as observed from experiments [14]. Hence, more sophisticated interactions are required, and we will further investigate this phenomenon in our future work.

**Nucleus size, factory numbers and factor numbers**

Osborne et al. [1] shows approximately 2000 transcription factories in the extended and flattened nuclei of mouse embryo fibroblast. In contrast (Fig. 6A), they found that erythroblast, B cell, T cell and fetal brain cells, which have spherical nuclei with significantly smaller radii and nuclear volumes, have dramatically fewer transcription factories (100-300 per nucleus). It was argued that the large differences in factory numbers seen in nuclei from tissues versus cells grown on a surface appear genuine and may be a consequence of a reduced potential for intra-chromosomal and especially interchromosomal sharing of factories in flattened cell nuclei [18]. To test how the changes in nuclei shape and transcription factory number will effect on gene colocalization, we ran simulations with flattened cells, squashing the original cubic from \(2 \times 2 \times 2\) to \(0.5 \times 4 \times 4\) but maintaining its volume (Fig. 6A). We have also tested the situation when the flattened cell is of volume five times bigger than the spherical cell \((0.5 \times \sqrt[3]{80} \times \sqrt[3]{80})\), according to the experimental observation (data available by request). We partitioned the flattened cell into four subunits \((0.5 \times 2 \times 2)\) and restricted the translocation of genes within each subunit, so all genes are restricted locally for consistency with experiments while transcription factors are free to move within the entire space. We found, consistent with our ‘limited resource’ theory, that no matter if we increase the volume of the flattened cell or not, the colocalized transcription is increased rather than reduced in the flatten cell, and is almost independent of the number of transcription factories (Fig. 6A). Moreover, it is also observed from Fig. 6A that increasing the volume of nucleus might increase the colocalization ratio. We will investigate the reason mathematically in our future research work.
From the analysis above, we propose the possibility to reconcile the facts observed in experiments and our model simulations: increasing the transcription factors might be the only possible mechanism to prevent gene colocalization. To confirm this, we then run the simulations with different number of transcription factors (Fig. 6B), for different volumes of the flattened cell. The left panel of Fig. 6B (unchanged volume) clearly demonstrated that when the number of transcription factors is around 30, the colocalization ratio is reduced to around 0.5 (non-colocalization) and is independent to the number of transcription factories. When the nucleus volume is enlarged 5 times bigger than the original spherical cell (Fig. 5B right panel), colocalization is even easier to happen for various cases, but increasing the number of transcription factory can hardly be the only reason for higher chance of colocalization.
Discussion

In the current paper, we have investigated whether a simple diffusion model can account for the colocalization observed in experiments, based upon parameters measured from experiments. We first assess the ratio of gene colocalization. It is found that the colocalization ratio is determined by the inter-colocalization intervals and is biased. We then applied the theory and numerical simulations to two and three dimensional cases with standard and fractional Brownian motion. We have shown that the experimentally observed colocalization is possible in both two and three dimensional cases and concludes that our dynamical model can match many experimental data.

However, a direct comparison with experimental data is still not easy since we do not have data of the dynamics of multi-genes. All experimental results are static results [14]. With the development of new experimental techniques, we expect that the dynamic data should be available soon. Such data would be valuable for us to understand the interactions between genes.

It is clear that our model is a simplified version of the gene mobilization in the nucleus: each gene is treated as independent (fractional) Brownian motion which is only true in local loci and small time intervals (Fig. 1A) [19,20]). The transcription process is also a simplified process. Moreover, we tried to introduce negative correlation between different families of genes and factors with preventing probability $p$. However, the simulation results on the gene-factor association level (Fig. 5Bc) did not show much difference after including the preventing probability among genes or factors. Hence, more sophisticated interactions are required, and we will further investigate this phenomenon in our future work.

In the models above, all genes are treated as a point (point model) and we should model 3D chromosomes as polymer chains [21]. The 3D whole genome conformation will be based on ChIA-PET data (see Lieberman-Aiden et al [22]). The dynamics of each polymer chain will be modelled according to the well known polymer physics [23], in collaborating with our experimental data. The simulation would be computational very expensive and we will run it on state-of-art clusters. The interactions in the model between genes (chromosomes) etc. should fit well with the known experimental data accumulated in our experimental teams for the past years.
After having a biophysically realistic model (with some coarse-grain approaches), we would expect to use the model to predict some key stages of hematopoietic differentiation. These predictions will then be tested by our experimental groups. Certainly this would be a very challenging task and it is a multi-scale spatio-temporal dynamics. Ideally we should be able to predict key decision making mechanisms at the molecular and cellular level that control genome function and may lead to the lymphoid versus myeloid differentiation. The transcription factories story might fit well with some general computational principle as reviewed in Oehler et al [24].

In general, we have to take into account the interactions between genes, both in cis and in trans, between genes and transcriptions, and between genes and transcription factories. As mentioned in Methods section (Eq. (1)), we can include the interactions between transcription units in the drift terms [25]:

\[
\begin{align*}
m \frac{d^2 u_i^T(t)}{dt^2} &= -\varsigma^T \int_{-\infty}^{t} \frac{d u_i^T(t)}{dt} K_H(t - s) \, ds \\
&\quad - m \Psi \sum_{j=1}^{N_f} \left( u_i^T(t) - c_j \right) + \sqrt{2 \varsigma^T k_B T_e} \frac{dB_H(t)}{dt}
\end{align*}
\]

where \( m \) is the mass, \( \Psi \) is the shape parameter of the harmonic potential, \( c_i \) is the centre of each factory and \( N_f \) is the number of transcription factories, \( k_B \) is the Boltzman constant, \( \varsigma^T \) is a friction constant, \( T_e \) is the temperature and \( K_H(t) \) is a kernel so that the fluctuation-dissipation theorem holds true. For each gene, it obeys similar equation

\[
\begin{align*}
m \frac{d^2 u_i^T(t)}{dt^2} &= -\varsigma \int_{-\infty}^{t} \frac{d u_i(t)}{dt} K_H(t - s) \, ds \\
&\quad - m \Psi \left( u^T(t) \right) \sum_{j=1}^{N_f} \left( u_i(t) - c_j \right) + \sqrt{2 \varsigma k_B T_e} \frac{dB_H(t)}{dt}
\end{align*}
\]

but with a potential depending on whether its corresponding transcription factor is in a factory or not (the term \( \Psi(u^T(t)) \)). How to find the right parameters of the interactions in the equations above would be an interesting issue. In the past decades, many techniques have been developed, mainly using the idea of Markov chain Monte Carlo and Bayesian
approaches (for example, Pavliotis and Stuart [26]). With the drift term introduced here, we could expect that sub-diffusion has a larger colocalization region than super-diffusion.

Recent research [27] shows that some transcription factors translocate into the nucleus in response to external signals (e.g., calcium flow), exhibiting short bursts of nuclear localization, suggesting that frequency-modulation regulation of localization bursts may be a general control strategy used by cell. Therefore, we could introduce external signal, such as calcium frequency or duration, to regulate gene cluster in specialized transcription factories [14] and study the effect on gene colocalization in our future work.
Methods

Standard Brownian motion

Assume that we have $m$ X genes and $k$ Y genes, with $n$ transcription factors of X gene and $l$ transcription factors of Y gene. Denote their positions at time $t$ as

- **X genes**: $x_1(t),...,x_m(t)$;
- **X factors**: $x_1^T(t),...,x_n^T(t)$;
- **Y genes**: $y_1(t),...,y_k(t)$;
- **Y factors**: $y_1^T(t),...,y_l^T(t)$.

All genes and transcription factors move according to diffusion processes, i.e.

$$dz(t) = b(\Theta)dt + \sigma^z dB^z(t),$$

where $b$ is the drift term depending on the global activity in the nucleus $\Theta$, $\sigma^z$ is the diffusion coefficient of transcription element $z$ (where $z$ can be $x$, $y$, $x^T$ or $y^T$) and $B^z(t)$ is the independent fractional Brownian motion. The genes and transcription factors move around with a constant diffusion coefficient

- **Gene**: $\sigma^x = \sigma^y = \sigma^\theta$;  
- **Factor**: $\sigma^{x^T} = \sigma^{y^T} = \sigma^\tau$.

The drift term summarizes the interactions between association of centromeres, clustering of co-regulated genes, association of a regulatory element and its target genes, interaction of a genome region with the nuclear envelope etc [11]. For a gene $h$, define a sequence of stopping (binding) times for X genes and Y genes as

- **X gene**: $\tau^i_h = \inf\{t: d(x_h, X^T) \leq \epsilon_0, t > \tau^{i-1}_h + T_b\}$,  
- **Y gene**: $\omega^i_h = \inf\{t: d(y_h, Y^T) \leq \epsilon_0, t > \omega^{i-1}_h + T_b\}$,
with \(\tau_h^0 = 0, \omega_h^0 = 0\), and \(j = 0, 1, \ldots\). Moreover, \(T_b\) is the binding time of a transcription factor, \(d(\cdot, \cdot)\) is the distance and \(X^T\) and \(Y^T\) are the sets of all available (unbound) transcription factors at time \(t\), i.e.,

\[
X^T = \{x^T_i \mid d(x^T_i, \{x_1, \ldots, x_m\}) > \epsilon_0\};
\]

\[
Y^T = \{y^T_i \mid d(y^T_i, \{y_1, \ldots, y_n\}) > \epsilon_0\}. \quad (6)
\]

Once a transcription factor binds to a gene of the same family, they will move together with gene diffusion coefficient \(\sigma^g\) (which is much slower than transcription factor diffusion coefficient \(\sigma^T\)), i.e.,

\[
x_i(t) = x^T_j(t) = X_i(t) \text{ if } d\left(x_i(\tau^h_i), x^T_j(\tau^k_i)\right) \leq \epsilon_0. \quad (7)
\]

When the bound gene-factor enters a factory, transcription starts. The transcription time for both X and Y gene are given by

\[
X \text{ gene: } \xi^j_{h,s} = \inf\{t : d(x_h, F_s) \leq \epsilon_0, t > \xi^j_{h} + T_t\}; \quad (8)
\]

\[
Y \text{ gene: } \eta^j_{h,s} = \inf\{t : d(y_h, F_s) \leq \epsilon_0, t > \eta^j_{h} + T_t\}. \quad (9)
\]

where \(j = 0, 1, \ldots\), \(\xi^0_{h,s} = 0\), \(\eta^0_{h,s} = 0\), \(F_s\) is the \(s\)th factory, \(s = 1, 2, \ldots, N_f\), and \(T_t\) is the transcription time length. We have

\[
X_h(t) = X_h(\xi^j_{h,s}); \quad (10)
\]

\[
Y_h(t) = Y_h(\eta^j_{h,s}). \quad (11)
\]

The physical meaning is clear: when the transcription starts, the gene is frozen and stays in the factory. For a given factory \(s\), we can calculate the colocalization event. Define the
colocalization event as the counting process of the inter colocalization interval $T_{c,xx}$ between one X gene and another X gene (see Fig. 1b) as

$$N_{xx}(s, [0 T]) = (\# \text{ colocalization events in time window } [0 T]).$$

We can define the colocalization event between X gene and Y gene $N_{xy}(s,[0 T])$ similarly. Let $u_{h,s}(t), v_{h,s}(t) \left( or \ u_{h,s}^T(t), v_{h,s}^T(t) \right)$ be the indicator function of the gene (or factor) transcription event of the $s$th factory for the $h$th gene (X or Y). Note that each process $u_{h,s}(t), v_{h,s}(t) \left( or \ u_{h,s}^T(t), v_{h,s}^T(t) \right)$ is a dichotomous random process.

The quantity we intend to calculate is

$$\lim_{T \to \infty} \frac{N_{xx}(s, [0 T])}{N_{xy}(s, [0 T]) + N_{xx}(s, [0 T])}.$$  \hspace{1cm} (12)

However, there is a problem if we calculate the ratio as above. When we count the events of $N_{xx}(s,[0 T])$, the population size is $m$ ($m - 1$), but for $N_{xy}(s,[0 T])$, it is $mk$. Hence we define

$$r_{xx/xy} = \lim_{T \to \infty} \frac{N_{xx}(s, [0 T])}{\frac{m - 1}{k} \cdot \frac{N_{xy}(s, [0 T])}{N_{xx}(s, [0 T])} + \frac{m - 1}{m - 1}}$$ \hspace{1cm} (13)

as the colocalization ratio. When $r_{xx/xy}$ is larger than 0.5, an X gene tends to transcript with another X gene more often in a factory. Let us first confirm that $r_{xx/xy}$ is independent of time and converges to a constant rapidly. From the definition of $N_{xx}(s, [0 T])$, it is the counting process of a renewal process with the inter colocalization interval $T_{c,xx}$. From the renewal theorem [28,29] we know that

$$E \left( \frac{N_{xx}}{T} \right) = \frac{1}{E(T_{c,xx})}, \quad E \left( \frac{N_{xy}}{T} \right) = \frac{1}{E(T_{c,xy})}$$ and
Hence, as $T$ is large enough we should have

\[
E\left(\frac{N_{xx}(s,[0,T])}{T} - \frac{1}{E(T_{c,xx})}\right)^2 = \frac{E(T_{c,xx} - E(T_{c,xx}))^2}{T \left(E^3(T_{c,xx})\right)}
\]

\[
E\left(\frac{N_{xy}(s,[0,T])}{T} - \frac{1}{E(T_{c,xy})}\right)^2 = \frac{E(T_{c,xy} - E(T_{c,xy}))^2}{T \left(E^3(T_{c,xy})\right)}.
\]  

(14)

Therefore whether there is a colocalization event in the nucleus is completely determined by the inter colocalization interval distribution $T_{c,xx}$ and $T_{c,xy}$.

**Fractional Brownian motion**

Is the standard Brownian motion good enough to match the experimental data? Chromatin loci are highly mobile but their motion is restricted within confined volumes. Each gene is constrained by interactions with immobile nuclear structures. Although chromosomes are relatively static, individual chromatin domains undergo Brownian motions and can extend far beyond the edges of their chromosome territory.

A normalized fraction Brownian motion $B_H(t)$ is a continuous-time Gaussian process starting at zero, with mean zero, and having the following covariance function

\[
E[B_H(t)B_H(s)] = \frac{1}{2}(|t|^{2H} + |s|^{2H} - |t - s|^{2H})
\]  

(16)

where $H$, called the Hurst index or Hurst parameter associated to the fractional Brownian motion, is a real number in $[0,1]$.

The value of $H$ determines what kind of process the fraction Brownian motion is:

- if $H = 0.5$, the process is in fact a standard Brownian motion;
o if $0.5 < H < 1$, the increments of the process are positively correlated (super-diffusion);
o if $0 < H < 0.5$, the increments of the process are negatively correlated (sub-diffusion).

According to Eq. (16), when $H$ is greater than 0.5, it moves faster than the normal diffusion ($H = 0.5$), hence it is called superdiffusion. We also use fractional Brownian motion in the model developed in the previous subsection.

**Colocalization moment and ratio between genes**

To compare with experimental data, next we introduce some quantities which are experimentally measurable. The transcription rate of an X gene is

$$r_1(u_1) = \lim_{S \to \infty} \frac{\int_0^S \left( \sum_{j=1}^{N_f} u_{1,j}(t) \right) dt}{S} = N_f E[u_{1,1}(t)], \quad (17)$$

where $E[\cdot]$ stands for the expectation, and $N_f$ is the number of transcription factories. This means that for an X gene, its transcription rate depends on the number of factories and the probability that this gene is being transcribed over time inside each factory. Similar definition can be given for Y gene transcription rate. The colocalization between two X genes is defined as

$$r_2(u_1, u_2 | u_1) = \lim_{S \to \infty} \frac{\int_0^S \left( \sum_{j=1}^{N_f} u_{1,j}(t)u_{2,j}(t) \right) dt}{\int_0^S \left( \sum_{j=1}^{N_f} u_{1,j}(t) \right) dt} = \frac{E[u_{1,1}(t)u_{2,1}(t)]}{E[u_{1,1}(t)]}, \quad (18)$$

describing the probability that when an X gene is being transcribed in a factory, another X gene is also being transcribed in the same factory at the same time. Similarly the colocalization ratio for an X and a Y gene is given by

$$r_2(u_1, v_1 | u_1) = \frac{E[u_{1,1}(t)v_{1,1}(t)]}{E[u_{1,1}(t)]}. \quad (19)$$

When X genes and Y genes are independent, we have
The X-X genes colocalization ratio defined before is simply given by

\[ r_{2}(u_1, v_1 | u_1) = E[v_{1,1}(t)] = \frac{r_1(v_1)}{N_f} \]  

(20)

This should give us a clear explanation why we call \( r_{xx/xy} \) the colocalization ratio. Therefore, when two X and X genes are colocalized, it should have

\[ r_{xx/xy} > 0.5 \].

**Colocalization moment and ratio between genes and factors**

Defined

\[ r_{1}(u^T) = \lim_{S \to \infty} \int_{0}^{S} \chi \left( \sum_{j=1}^{N_f} \sum_{i=1}^{N_T} u_{i,j}^{T}(t) \right) dt = E \left[ \chi \left( \sum_{j=1}^{N_f} \sum_{i=1}^{N_T} u_{i,j}^{T}(t) \right) \right] \]  

(22)

as the X transcription factor association rate with all factories, where \( N_f \) is the number of transcription factors and \( \chi(w) \) is the characterization function, i.e. \( \chi(w) = 1 \) whenever \( w > 0 \), and \( \chi(w) = 0 \) otherwise. We assume that all processes are stationary. When \( u_{i,j}^{T}(t) \) is sparse, we have

\[ E \left[ \chi \left( \sum_{j=1}^{N_f} \sum_{i=1}^{N_T} u_{i,j}^{T}(t) \right) \right] = E \left[ \sum_{j=1}^{N_f} \sum_{i=1}^{N_T} u_{i,j}^{T}(t) \right] . \]  

(23)

Hence the transcription factor association rate \( r_{1}(u^T) \) is simply \( N_f \cdot N_T \cdot E[u_{1,1}^{T}(t)] \). The advantage of our approach over the experimental is that we have a dynamical model and we can concentrate on each individual transcription factory. To this end, we will concentrate on the dynamic behaviour of a single transcription factory: peer through one single factory. Under the ergodicity assumption, we intend to match the modelling results with experimental results which are obtained with spatio average. Hence we drop the
transcription factory subscript $j$ from now on. In our simulations, for a fixed number of transcription factors, we find a binding time so that X factors (as Klf1 in Schoenfelder et al [14]) are colocalized with the factories with a rate

$$r_1(u^T) = \mathbb{E}[\chi\left(\sum_{i=1}^{N_T} u_i^T(t)\right)].$$

(24)

The colocalization ratio between the first X gene and X factors is given by

$$r_2(u_1, u^T|u_1) = \lim_{s \to \infty} \frac{\int_0^s \mathbb{E}\left[\chi\left(u_1(t)\left(\sum_{j=1}^{N_T} u_j^T(t)\right)\right)\right] dt}{\int_0^s \mathbb{E}\left[\chi(u_1(t))\right] dt} = \frac{\mathbb{E}\left[\chi\left(u_1(t)\left(\sum_{j=1}^{N_T} u_j^T(t)\right)\right)\right]}{\mathbb{E}[u_1(t)]}$$

(25)

This gives us the Klf1-associated ratio for the first X gene (say, Hbb, Hba, Hmbs and Epb4.9 in Schoenfelder et al [14]) per factory. Again, when the event is sparse, we have

$$r_2(u_1, u^T|u_1) = \frac{\mathbb{E}[\chi\left(u_1(t)\left(\sum_{j=1}^{N_T} u_j^T(t)\right)\right)\]}{\mathbb{E}[u_1(t)]} = \frac{N_T E[u_1(t)u_i^T(t)\]}{\mathbb{E}[u_1(t)]}$$

(26)

Similarly,

$$r_2(v_1, u^T|v_1) = \frac{\mathbb{E}\left[\chi\left(v_1(t)\sum_{i=1}^{N_T} u_i^T(t)\right)\right]}{\mathbb{E}[v_1(t)]}$$

(27)

is the Klf1-associated rate with Y genes or Z genes. When they are sparse and independent, it equals $N_T E[u_{1,1}(t)]$. A pair of X genes colocalized ratio per factory with Klf1 is

$$r_3(u_1, u_2, u^T|u_1, u_2) = \frac{\mathbb{E}\left[\chi(u_1(t)u_2(t)\sum_{i=1}^{N_T} u_i^T(t)\right)}{\mathbb{E}[u_1(t)u_2(t)]}$$

(28)

Since X and Y genes are independent in our model, the colocalization ratio of an X and a Y gene pair with Klf1 becomes

$$r_3(u_1, v_1, u^T|u_1, v_1) = \frac{\mathbb{E}\left[\chi(u_1(t)v_1(t)\sum_{i=1}^{N_T} u_i^T(t))\right]}{\mathbb{E}[u_1(t)v_1(t)]}$$

(29)
When they are sparse and independent, we have

\[
 r_3(u_1, v_1, u^T|u_1, v_1) = \frac{N_T E[v_1(t)] E[u_1(t)u^T_{1,1}(t)]}{E[v_1(t)] E[u_1(t)]}
\]

\[= \frac{N_T E[u_1(t)u^T_{1,1}(t)]}{E[u_1(t)]} = r_2(u_1, u^T|u_1) \tag{30} \]
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Figure Captions.

**Fig. 1.** Demonstration of positionings and states of gene and transcription factory. (A) Schematic representation of chromatin loops (black) extruding from a chromosome territory (gray). Transcribed genes (white) in RNAP II factories (black circles). Potentiated genes (free loops) that are not associated with RNAP II factories are temporarily not transcribed. Potentiated genes can migrate to a limited number of preassembled RNAP II factories to be transcribed (dotted arrows). Both cis and trans associations are possible. If a piece of gene is being transcribed at a particular time $t$, we define the state of that piece of gene as 1 (e.g., $u_{1,1}(t) = 1$, where the first subscript indicates the specific gene, and the second subscript indicates the specific factory that the gene is associated with), otherwise, the state of the gene is defined to be 0. Hence, $u_{1,1}(t) = 1$ means X gene 1 is being transcribed at factory 1, $v_{1,1}(t)$ means Y gene 1 is being transcribed at factory 1, and $u_{3,2}(t) = 1$ means X gene 3 is being transcribed at factory 2. (B) Illustration of the colocalization events (red lines) among the same family of genes (X gene) within time window $[0,T]$ (in dashed line window) in factory 1. Once there is a gene start being transcribe inside factory 1 while there are one or more than one genes already being transcribed inside the same factory at time $t$, we say there is a colocalization event happened at time $t$. Therefore, we have 4 colocalization events among X genes in total over time $T$, i.e. $N_{xx}(1,[0,T]) = 4$. The inter-colocalization interval $T_{c,x,x}$ tells the timing of X-X gene colocalization. (C) Illustration of colocalization events between different families of genes (X gene and Y gene) within time window $[0,T]$ in factory 1. The X-Y gene colocalization events are similarly defined as X-X gene, as shown by the figure, where we have 4 colocalization events between X gene and Y gene over time $T$, i.e. $N_{yx}(1,[0,T]) = 4$, and $T_{c,y,x}$ is the inter-colocalization interval between Y gene and X gene.

**Fig. 2:** Simulation results for 2D case. (A) One example of colocalization ratio convergence. The ratio rapidly becomes stable as time increases. Yellow triangle is obtained from Eq. (15), showing the consistency of calculation of colocalization ratio using different variables. (B) The inter-colocalization (ICI) interval distribution for $T_{c,xx}$ and $T_{c,xy}$. The parameters are the same as displayed in (A). The inter-colocalization interval distribution can be fitted with gamma distributions. (C) The mean values of ICI distribution by varying the factor number and binding time between genes and factors. The mean ICI increases as the binding time increases, and decreases as the factor number increases. (D) The colocalization probability for X-X gene and X-Y gene. (E)
Colocalization ratio for various combinations of factor number and binding time. The ratio threshold is set to be 0.52. The red dashed curve is the threshold boundary to distinguish whether the colocalization is significant or tends to be random.

**Fig. 3:** Simulation results for 3D case, similar as Fig. 2. (A) Demonstration of the 3D framework in a cubic, where the big green dots represent the location of the transcription factories, asteroids the transcription factors, and small dots the genes. Different families are represented by either red or blue. (B) The inter-colocalization interval distribution for $T_{c,xx}$ and $T_{c,xy}$. The parameters are the same as in (D). (C) The mean values of ICI distributions by varying the factor number and binding time between genes and factors. The mean ICI increases as the binding time increases, and decreases as the factor number increases. (D) The distribution of the colocalization ratio for X-X genes and X-Y genes. (E) The colicalization ratio for various combinations of factor numbers and binding times. The critical ratio is 0.52. The red dashed curve is the threshold boundary to distinguish whether the colocalization is significant or tends to be random.

**Fig. 4.** Colocalization regions for sub- and supperdiffusion Brownian motions. (A) The inter-colocalization intervals for super- and sub-diffusion for 5 transcription factors and 10 sec binding time. (B) Colocalization regions for different factors and binding times for sub-diffusion case ($H = 0.1$) and super-diffusion case ($H = 0.9$).

**Fig. 5.** Colocalization between genes and transcription factors. (A) Experimental results with immuno-RNA FISH reveal colocalization between transcription factors and factories (RNAPII), genes and transcription factors, and gene pairs and transcription factors. This is the contour plot from original experimental data. (Aa) Immuno-fluorescence detection of Klf1 (red) and RNAPII-S5P (green) in definitive erythroid cells, with a scale bar of 2 µm. This shows the colocalization between Klf1 and transcription factories RNAPII. This Klf1 background association rate (level) is estimated to be 20%. (Ab) The colocalization between transcription factor Klf1 and Hbb gene. (Ac) The colocalization between factor Klf1 and genes pairs (Hbb and Hist1). (B) Simulation results when we hold the Klf1 background association level as 20%, while the translocation of transcription units (genes, factors and factories) are following sub-diffusion process ($H = 0.4$). (Ba) Gene-factor association level (numbers indicated behind the stars) with various factors and binding time, both for 2D and 3D cases. The number of genes (for each family) is fixed to be 5 and the Klf1 (X factor) background association level is fixed to be 20% (the stars indicate the parameter values when this condition is
satisfied). The detailed Klf1 association level for each X gene and Y gene are presented in the figure below, revealing the fact that the simulation results for 3D case (gene-factor association level = 0.6) match the experimental result (gene-factor association level = 0.64) quite well for a specific set of parameters (5 genes, 2 factors and 130 sec binding time). (Bb) Gene-factor association level (numbers besides the stars) with various genes and binding time for 2D and 3D cases. The number of factors is fixed to be 2 and the Klf1 background association level is fixed to be 20%. (Bc) The association rate of Klf1 (X factor) with Z gene and paired X-Z genes when there is a negative correlation between X and Z gene, or X and Z factors, under 3D case. The parameter used here are 5 genes, 2 factors, and 130 sec binding time. Preventing probability means the chance for stopping another gene (factor) to enter the factory when there is already one gene (factor) in that factory. When $p = 0$, it represents the independent situation of X factor (gene) and Y factor (gene).

**Fig. 6.** The simulation results for ratio of colocalization in flattened and spherical nucleus. (A) The colocalization ratio increases as the degree of flatness of the nucleus increases when there are 5 factors and 10 sec binding time, and is independent of the number of transcription factories, at least in the flattened nucleus, such as E10 (embryonic blood), E14 (fetal liver erythroid), AS (adult anemic spleen erythroid), Sp (normal adult spleen), Th (adult thymus), Br (fetal brain), mouse embryonic fibroblasts (MEFs) in experiments. Scale bar = 10 µm. The cubic of spherical nucleus and rectangular block of flatten nucleus demonstrate the positioning of transcription factories, factors and genes (refer to Fig. 3A). Note that the volumes of the cubic spherical nucleus and the rectangular flattened block are either the same (solid line), or the volume of the flattened nucleus is 5 times bigger than that of the spherical nucleus (dash line). (B) The colocalization ratio is a decreasing function of the number of transcription factors for both flattened nucleus of the same volume as the spherical one, and the nucleus of 5 times larger volume. No matter the volume, the colocalization ratio is independent of the transcription factories.
### Tables

|                           | Brownian motion          | Fractional Brownian motion |
|---------------------------|--------------------------|-----------------------------|
|                           | 2D                       | 3D                         | 2D                       | 3D                       |
| Gene family               | X gene (Hbb, Hba, Hmbs, Epb4.9), Y gene (Cpox) | X gene; Y gene             | X gene; Y gene           | X gene; Y gene           |
| Factor family             | X factor, Y factor       | X factor, Y factor          | X factor, Y factor       | X factor, Y factor       |
| # of genes for each family| 40                       | 100                        | 40                       | 100                      |
| # of factors for each family | 5-12                   | 5-25                       | 5-12                    | 5-25                     |
| # of factories            | 9                        | 35                         | 9                       | 35                       |
| Binding Time $T_b$ (sec)  | 10-50                    | 10-150                     | 10-50                   | 10-150                   |
| Gene diffusion rate $\sigma^g$ | $\sqrt{0.001}$        | $\sqrt{0.001}$             | $\sqrt{0.001}$          | $\sqrt{0.001}$          |
| Factor diffusion rate $\sigma^T$ | $\sqrt{0.01}$       | $\sqrt{0.01}$              | $\sqrt{0.01}$           | $\sqrt{0.01}$           |
| Transcription time $T_t$ (sec) | 300                     | 300                        | 300                     | 300                      |
| Gene radius $r_g$ (µm)    | 0.01                     | 0.025                      | 0.01                    | 0.025                    |
| Factor radius (µm)        | $r_g / 3$                | $r_g / 3$                  | $r_g / 3$               | $r_g / 3$               |
| Factory radius (µm)       | $r_g * 10 / 3$          | $r_g * 10 / 3$             | $r_g * 10 / 3$          | $r_g * 10 / 3$          |
| Gene radius after binding with factor (µm) | $1.5 * r_g$            | $1.5 * r_g$                | $1.5 * r_g$             | $1.5 * r_g$             |

**Table I.** Parameter values used for 2D and 3D Brownian motion and fractional Brownian motion.
|                  | Klf1 | X   | Y   | Z   | X-X | X-Y | X-Z |
|------------------|------|-----|-----|-----|-----|-----|-----|
| **Simulation**   | 0.2  | 0.6 | 0.2 | 0.2 | 0.8 | 0.6 |     |
| **Experiment**   | 0.2  | 0.64| 0.2 | 0.2 | 0.8 | 0.64| 0.2 |

Table II: Comparison of sub-diffusion 2D simulation results (Fig. 5b, 3D case) and the experimental results.
A

B

C

Inside factory 1

start

end

X gene

1 2 3 4

T_{c,xx}

time

T

Y gene

1 2 3

T_{c,yx}

time

T
**Colocalization convergence**

2D simulation of gene translocation in a cell.

Gene: X−Gene (40), Y−Gene (40),
Factor: X−Factor (5), Y−Factor (5),
Binding Time: 10 sec.

\[x\text{Gene} \rightarrow x\text{Gene} = 0.559\]
\[y\text{Gene} \rightarrow y\text{Gene} = 0.561\]

\[x\text{Gene} \rightarrow y\text{Gene} = 0.441\]
\[y\text{Gene} \rightarrow x\text{Gene} = 0.439\]

| X−Gene | Y−Gene |
|--------|--------|
| 12341  | 10001  |
| 10022  | 12485  |

**T\text{c} distribution**

\[T\text{c, XX} \]
\[T\text{c, XY} \]

**Mean of T\text{c} distribution**

\[T\text{c, XX (sec)}\]
\[T\text{c, XY (sec)}\]

**2D colocalization probability**

Factor number = 7
Binding Time = 1

\[\text{mean} = 0.02\]
\[\text{mean} = 0.033\]

**Colocalization ratio**

X−X gene

Factor Number: 0.52, 0.54, 0.56, 0.58
Binding Time: 20, 40, 60, 80, 100

Counts: 0, 2000

Counts: 0, 2000
A  

**T<sub>c</sub> distribution**

H = 0.1  

![Graph showing distribution](image1)

H = 0.9  

![Graph showing distribution](image2)

\[ T_{c,xx} \text{ mean}=162 \]

\[ T_{c,xy} \text{ mean}=344 \]

\[ T_{c,xx} \text{ mean}=485 \]

\[ T_{c,xy} \text{ mean}=500 \]

---

B  

**2D fBM Colocalization ratio**

H = 0.1  

![Heatmap showing colocalization ratio](image3)

H = 0.9  

![Heatmap showing colocalization ratio](image4)

**Factor Number**

**Binding Time (sec)**

---

**mean=162**

**mean=344**

**mean=485**

**mean=500**
A  Experiments

b. Gene number is fixed to be 5

| Factor Number | 0  | 2  | 4  | 6  | 8  |
|---------------|----|----|----|----|----|
| Binding Time  | 0  | 2  | 4  | 6  | 8  |
| 0             | 0.22 | 0.23 | 0.25 | 0.28 | 0.31 |
| 40            | 0.32 | 0.41 | 0.45 | 0.48 | 0.60 |

b. Factor number is fixed to be 2

| Gene Number | 0  | 20 | 40 | 60 |
|-------------|----|----|----|----|
| Binding Time| 0  | 0.2 | 0.4 | 0.6 |
| 0           | 0.28 | 0.31 | 0.29 | 0.30 |
| 0.2         | 0.29 | 0.30 | 0.30 | 0.31 |
| 0.4         | 0.46 | 0.36 | 0.33 | 0.30 |
| 0.6         | 0.60 | 0.31 | 0.30 | 0.30 |

B  Simulations results

Klf1 background association level = 20%, sub-diffusion process

a. Gene number is fixed to be 5

b. Factor number is fixed to be 2

c. negative correlation between factors or genes
A) Colocalization ratio of spherical and flattened nucleus (10 sec binding time, 5 factors)

B) Colocalization ratio of flattened nucleus

The same size of volume as spherical nucleus

Volume 5 times bigger than spherical nucleus