

Chapter 1

Introduction

The nucleus of a human cell contains approximately 2 metres of DNA [1], requiring a huge degree of compaction to fit within the nucleus. In addition to storing DNA the nucleus itself is a highly functional organelle, responsible for the huge variety of complex cellular processes. To achieve this, the contents of the nucleus are very precisely organised into sub-nuclear compartments which specialise in processes such as transcription and replication. In addition to the organisation of the nuclear proteome, chromatin itself is thought to be organised. In this chapter I will discuss the current understanding of nuclear structure and organisation, how we have obtained this knowledge, and how it may play a part in the initiation of certain cancers.

1.1 Chromatin

Chromatin is a general term used to describe the DNA found in the nucleus along with the plethora of proteins that bind to it. There are two principle forms of chromatin: *euchromatin* and *heterochromatin*. These were first described in the early twentieth century by the difference in staining within the nucleus [2].

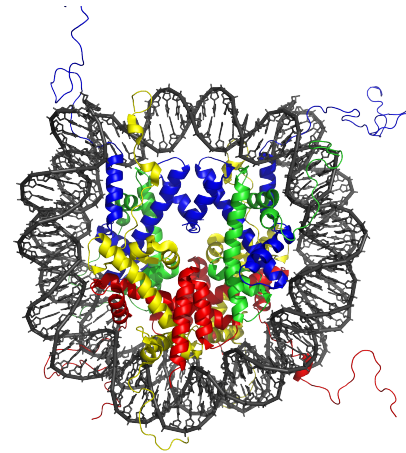
Heterochromatin stains darkly because it remains highly condensed during interphase, typically relocating to the nuclear periphery. can be split into *constitutive heterochromatin* and *facultative heterochromatin*; the former describing heterochromatin found in all cell types typically containing repetitive elements found in centromeres and telomeres; the latter describing less compacted inactive chromatin which may vary between cell types as they differentiate. Heterochromatin is well known as being a repressive environment for gene expression; studies whereby normally active regions are artificially tethered to the inner nuclear membrane show the ablation of gene expression [3, 4] though this effect appears to be locus specific and is not always observed [5]. Euchromatin is the site of most genic transcription, as shown by the presence of RNA Polymerase and nascent transcriptions found in early fractionation studies [6]. Its

looser compaction allows access to the DNA by the cellular machinery; enabling the binding of transcription factors and the initiation of transcription.

The differences between heterochromatin and euchromatin lie within the proteins that they contain. Chromatin acts as a platform for proteins to bind to, differences in histone modifications, variants, nucleosome packing and DNA modifications affect the accessibility and binding profile of the chromatin, and so how the DNA is interpreted by the cell.

1.1.1 Histones

To package DNA, the double helix is wrapped around an octamer of core histones: two H2A, two H2B, two H3 and two H4. 146 base pairs of DNA wrap around these positively charged proteins to form the nucleosome, which is then bound by histone H1 with linker DNA to make a total of 166 base pairs [7]. This packing forms the 10 nm fibre, often known as 'beads on a string' due to its appearance in electron micrographs. At their most basic level, histones function to compact DNA by counteracting the negative charge of the phosphorylated back bone.



1.1.2 Histone modifications

Core histones have flexible tails which extend outside of the nucleosome, and are accessible to proteins within the nuclear matrix.

These tails can be post-translationally modified at a large number of residues and these modifications can affect the packing of chromatin [8] as well as which proteins can bind. The large number of

combinatorial possibilities that result from these modifications have been dubbed the 'Histone Code' [9], different modifications are related to different chromatin states: for example, active promoters are typically enriched for di- and tri-methylation of histone 3 lysine 4 (H3K4), inactive promoters for H3K27me3 and H3K9me3 (Figure 1.2) (for a review, see [10]).

Figure 1.1 – Structure of the nucleosome. X Ray Structure of a Nucleosome core particle at a resolution of 1.9Å. DNA can be seen wrapped around the core histones. PDB structure 1KX5 [7].

As chromatin immunoprecipitation (ChIP) has become a common laboratory technique and been combined with microarray techniques (ChIP on chip) and next generation sequencing (ChIP-Seq), our understanding of how these marks affect chromatin biology on a genome-wide scale has advanced dramatically. Profiling chromatin types using multiple datasets covering a large number of histone modifications is sufficient to predict the identity and function of

regions in the genome with a high degree of accuracy, revealing previously unknown enhancers [11, 12, 13]. Ernst *et al.* used the genome-wide profiles of 9 histone modifications in 9 different cell types to define 15 different chromatin states, describing different states of promoters, enhancers and insulators, amongst others[14]. They integrated data from genome-wide association studies (GWAS) and found numerous enhancer elements which coincide with disease associated mutations. This systems biology approach can reveal the dynamics across different cell types and can be a powerful tool in understanding how the genome is interpreted in health and disease.

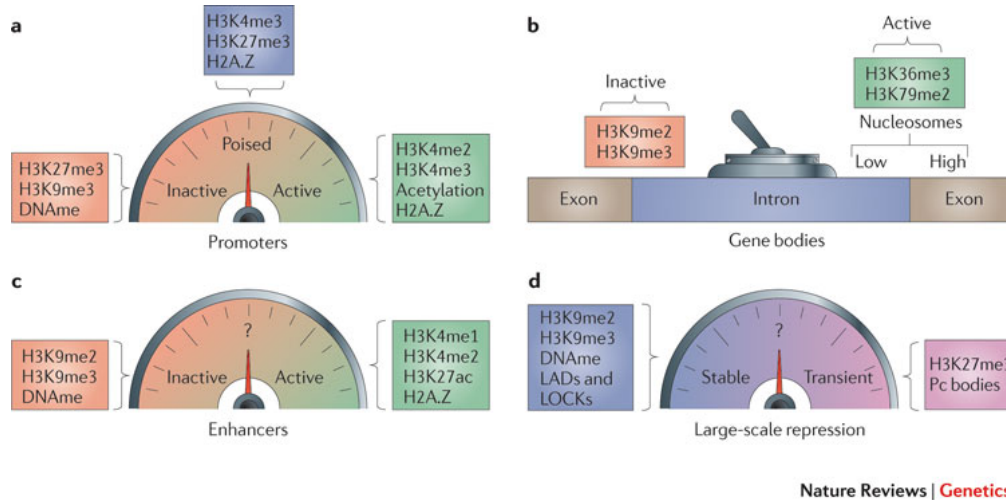


Figure 1.2 – Histone modification associations in chromatin. Diagram showing the range of histone modifications available within mammalian chromatin and how they can specify the behaviour of elements within the DNA. Taken from[10].

1.1.3 Histone variants

In addition to histone tail modifications, chromatin can be modified by the incorporation of histone variants. Canonical core histone genes are found in clustered repeat arrays within the genome, are transcribed during replication and are extremely tightly conserved between species. Histone variants are found as single genes spread through the genome, and are subject to far greater diversity [15].

CENP-A is a human variant of histone H3 which replaces the canonical histone in centromeric heterochromatin. It is a key factor in the establishment of the centromeres and kinetochores required for mitosis. The histone variant is incorporated with the help of a number of chaperone proteins, including HJURP, after replication of DNA has finished [16, 17]. CENP-A is essential for the formation of centromeres.

Another frequent histone variant found in humans is H3.3, which differs from canonical H3 by just four amino acids [15]. This histone variant is found within transcribed genes, promoters and regulatory elements, and is thought to be laid down during transcriptional elongation [18]. Nucleosomes containing H3.3 appear to be less stable than canonical nucleosomes, with a high turnover of H3.3 [18]. It is possible that this increased turnover of the nucleosomal

components helps to keep the chromatin open and accessible to the transcriptional machinery [15].

Other core histone proteins also have variants, such as H2A.Z, a histone variant found on either side of the *nucleosome free regions* found at the transcriptional start sites of active genes as well as insulator regions [19]. H2A.Z promotes the recruitment of RNA Polymerase II through mediating protein interactions [20].

1.2 Two-dimensional organisation

Since the inception of cytogenetics, it has been known that not all chromosomes are the same. The development of chromosome banding techniques in the 1970s allowed detailed human karyotypes to be determined, complete with differential staining of regions within each chromosome, termed 'bands' [21]. Banding assays can show regions of heterochromatin and euchromatin, highlighting the variation in characteristics across regions of the genome [22]. The sequencing of the human genome allowed detailed analysis of GC content, gene density and repetitive sequence content [23]. Versteeg *et al.* integrated a multitude of SAGE tag expression profiles from different cell types into the genomic map and built on earlier work defined regions of high transcriptional activity, called ridges [24, 25]. They found ridges to be gene-dense, highly transcribed, have a high GC content and low LINE repeat density. These features are based purely on the underlying sequence content and so do not vary amongst cell types. The different ridges and anti-ridges were found to contain different classes of genes, with weakly expressed genes clustering within anti-ridges and clusters of highly expressed housekeeping genes found predominantly in ridges [24].

Whilst the two-dimensional organisation of the genome cannot completely explain the degree of complexity found with transcriptional control, these studies demonstrate that the order of sequence within the genome is not random, and has a large effect on the way that DNA content is interpreted.

1.3 Chromosome Territories

The interphase nucleus is a highly structured organelle. As chromosomes decondense after metaphase they retain some degree of structure, forming 'chromosome territories' (CTs) [26]. Circumstantial evidence for interphase organisation of chromosomes has existed for a long time, first suggested by Carl Rabl in 1885 [27]. Observations by Stack *et al.* using microscopy with giemsa-band staining suggested that chromosomes retained some degree of organisation during interphase [28], and in 1982 Cremer *et al.*, showed that interphase chromosomes exist in 'territories' by studying the pattern of DNA damage in metaphase chromosomes after spot irradiation in during interphase [29]. The subsequent development of 'chromosome paints', a method to visualise entire- or part- chromosomes with fluorescence *in-situ* hybridisation (FISH), confirmed these findings [30, 31, 32].

As FISH techniques have developed, so too has the detail with which CT organisation can be studied. Several groups have shown that chromosome territory position within the nucleus is not random and correlates with chromosome size [33, 34, 32], gene-density [34, 35] and replication timing [36, 37]: those near the centre of the nucleus tend to be gene-rich, early replicating and small. The specificity of CT positioning is a topic of debate, though studies have suggested conservation of CT positioning through evolution [38] and shown tissue specificity in chromosome pairing [39, 40].

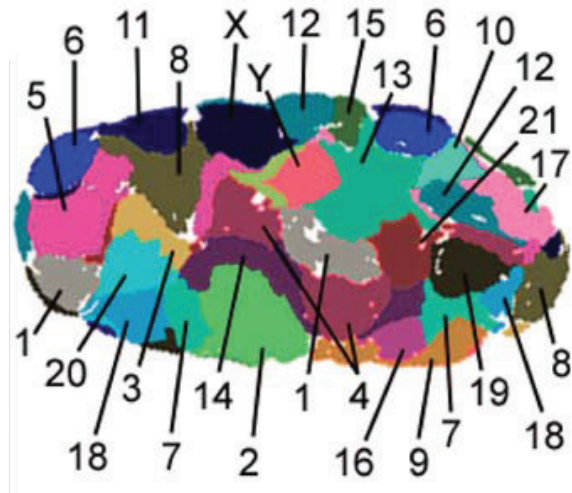


Figure 1.3 – Chromosome Territories. Image of a Human fibroblast cell in G0 with all chromosomes labelled using multi-colour FISH. Adapted from Bolzer *et al.* [32]

1.3.1 CTs and transcription

It's known that for some but not all genes, positioning at the nuclear periphery correlates with reduced gene expression [41, 42, 43]. To investigate whether transcriptional activity is the cause or effect of this positioning effect, two groups published studies which artificially tethered genomic regions to the inner nuclear membrane using Lac operators (*lacO*) [3, 4]. Both groups observed a decrease in the transcriptional activity of the regions when tethered, an effect that was ablated when cells were treated with trichostatin A (TSA) to inhibit class I and class II histone deacetylases [3], though this effect does not appear to apply to all loci [5]. Whilst these studies suggest that it is nuclear positioning that leads to transcriptional effects, a study by Croft *et al.* showed that inhibition of transcription causes a reversible change in CT position [35].

Large scale rearrangements of CTs have also been observed during cell differentiation [44, 45], likely due to changes in transcriptional profiles and chromatin remodelling. As such, it seems that there is a dynamic interplay between CT positioning and transcription - gross transcriptional patterns may drive the position of chromosomes within the nucleus, and those positions may in turn affect the transcription of the genes they contain. Interestingly, derivative chromosomes that result from balanced translocations affect the organisation of CTs within the nucleus [46], raising the possibility that the global changes in gene expression observed after oncogenic translocations could be in part due to changes in genome organisation [46].

1.3.2 Chromosome territory dynamics

The movement of chromatin loci within the nucleus is known to be largely due to local brownian motion, or “constrained diffusion” and seems to be limited by attachment to a nuclear cytoskeleton, nucleoli and the inner nuclear membrane

[47, 48]. Other studies have shown a mixture of local diffusion and larger, active movements [49]. Chuang *et al.* studied Chinese hamster ovary (CHO) cells after stimulation with a transcriptional activator and analysed the movement of the target chromosome from the nuclear periphery to the interior of the nucleus [50]. They found that chromosome movements happened in rapid unidirectional bursts, suggesting an active mechanism. This was supported by a later study in human fibroblast cells by Mehta *et al.* who demonstrated chromosome movement only 15 minutes after serum starvation, a process that was ablated by the inhibition of actin and myosin polymerisation [51].

1.3.3 Chromatin decondensation and the inter-chromosomal space

In addition to the study of whole chromosome positioning, there has been a great deal of research into the positioning of individual sequences relative to their chromosome territory. Early studies suggested that transcribed genes were found at the surface of chromosome territories [52] leading to a model whereby transcriptionally inactive genes are buried within territories and expressed genes contact in an inter-chromosome domain (ICD) where they can contact the transcriptional machinery [53]. This model gained support due to studies showing genes at the periphery of CTs [54] and new techniques to visualise the ICD using microscopy [55]. A number of FISH studies showed genes moving away from their territories in large loops upon activation [56, 57, 58], suggesting that they may be recruited to the ICD for transcription.

Despite these case studies, looping is not a prerequisite for transcription; other DNA-FISH studies have shown transcription within the volume of chromosome territories [59, 60]. Osborne *et al.* showed that actively transcribed *Uros* gene is more frequently outside its CT than the inactive gene *Fgfr2*, though this position was not necessary for transcription, suggesting that actively transcribed genes may preferentially locate to the surface of CTs, but this alone is not sufficient to drive transcription [61]. As such, it has been suggested that the inter-chromosomal domain model should be renamed the inter-*chromatin* domain model, whereby chromosomes are invaginated with channels and subdivided into ~1 Mbp domains of chromatin [26].

1.3.4 Chromosome territory intermingling

A question that followed immediately on from the discovery of loops extending from chromosome territories was that of chromosome intermingling. Mathematical modelling approaches suggested that irradiation induced DNA damage corresponded to the intermingling of chromosome territories [62, 63]. This was later backed up with a study by Branco and Pombo, who studied thin cryosections of nuclei with high resolution light and electron microscopy; they found that there is significant intermingling of chromatin between chromosome territories, and that the extent of this intermingling correlated strongly with previously recorded irradiation induced translocation frequencies [64]. In support of chromosome intermingling, the *HoxB* extra-chromosomal loops found to extend from the chromosome

territory upon gene activation [58] were found to make increased *trans* chromosomal interactions whilst looping out [65], suggesting that the loops contact other chromosomes rather than occupying an empty inter-chromosomal space.

1.4 Nuclear compartmentalisation

- Cajal bodies
- Speckles
- Nucleoli
- Polycomb / silencing bodies

1.4.1 Transcription factories

One nuclear subcompartment which has come to light within the past thirty years is the *transcription factory*, foci of hyper-phosphorylated RNA Polymerase II spread throughout the nucleus. The majority of geneic transcription appears to take place at transcription factories [66, 61, 67], challenging the classical model of transcription found in many text books.

The first study to suggest that Eukaryotic transcription does not take place with a processive RNA Polymerase moving along a DNA template was by Jackson, McCreedy and Cook in 1981. They showed that nascent RNA transcripts labelled with [³H] uridine remained within the nucleus when loops of DNA were removed using a nuclease[69]. They went on to show that RNA Polymerase II and active genes were also resistant to elution after chromatin digestion [70]. The term 'transcription factories' was coined by Jackson *et al.* in 1993. Fluorescence microscopy was used to label the incorporation of bromouridine triphosphate (BrUTP) into nascent RNA; discrete foci of nascent transcription could be seen within the nucleus which did not form in the presence of the RNA Polymerase II inhibitor α -amanitin [66]. Further studies showed that these foci contain RNA Polymerase II along with many other components required for transcription [71, 72]. An ultrastructural study by Eskiw *et al.* used correlative microscopy with both electron spectroscopic imaging (ESI) and fluorescence microscopy to study nuclei sections. ESI can distinguish nitrogen and phosphorous atoms without labelling, and fluorescence light

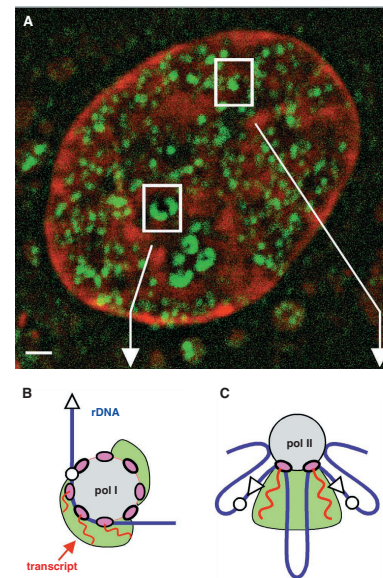


Figure 1.4 – Nascent RNA and transcription factories. (A) Transcription foci in HeLa cells, visualised with labelled Br-UTP in 100 nm cryosections. Nascent RNA (green) is concentrated in punctate foci. (B) Model for a nucleolar factory, showing a transcript with multiple polymerases generating a crescent shaped foci. (C) Model for a nucleoplasmic factory. Multiple transcribed regions each with a single polymerase generate a smaller cloud of nascent transcripts. Taken from Cook *et al.*, Science (2009) [68].

microscopy can visualise transcription through labelling BrUTP in nascent transcripts. The authors found that nascent RNA is almost always associated with the surface of large nitrogen-rich protein structures with a diameter of ~87 nm, comparable in size to that of a transcription factory [67].

The discovery of transcription factories has demanded a new model for the action of RNA Polymerase II [68]. The revised model proposes that instead of RNA Polymerase II freely diffusing to active genes and tracking along the gene body, it is the genes that are recruited to transcription factories and the genes that track through a stationary polymerase. Such a model provides a better explanation for the mechanics of transcription - clustering of transcriptional activity may enable the cell to conduct transcription in a much more efficient manner; HeLa nuclei have a 1 μ M concentration of active RNA Polymerase II, whereas the local concentration within transcription factories is closer to 1 mM [73]. Additionally, a polymerase enzyme moving along a gene would rotate with the helix of the DNA wrapping the nascent transcript around the template. Genes pulled through static transcription factories would extrude their RNA transcripts into the nucleoplasm [71, 68], creating topological loops within the template DNA instead which may be removed through the activity of topoisomerases.

Jackson *et al.* went on to work on a quantitative analysis of transcription factories in HeLa cells calculating the number of active RNA polymerases, the number of transcription sites and the number of polymerases associated with each transcriptional unit [74]. They showed that each HeLa cell nucleus contain ~ 2400 transcription factories, each with approximately 30 active RNA-Polymerase II complexes [74]. Importantly, this study showed that there are more transcribing units than there are foci of transcription, suggesting that genes must colocalise to transcribe. The number of transcription factories varies a great deal amongst cell types, but the observation that genes colocalise within transcription factories has been confirmed by a number of different techniques; Osborne *et al.* showed that genes situated within transcription factories are actively transcribed, whereas those outside are not [61]. They went on to demonstrate that transcription is a discontinuous process, with the frequency of nascent RNA transcription foci related to cellular mRNA concentrations, suggesting that transcription occurs in bursts. Multiple genes both in *cis* and in *trans* were seen to dynamically colocalise in transcription factories, supporting predictions that genes must share transcription factories [61, 74]. A later paper by Osborne *et al.* showed that the immediate-early gene *Myc* and *Fos* are dynamically recruited to existing transcription factories within 5 minutes of B-cell stimulation, suggesting that the recruitment of genes to pre-existing transcription factories may be a method of transcriptional control [75].

1.4.1.1 Specialised transcription factories

After the demonstration that genes shared transcription factories, a number of groups postulated that specific genes may colocalise at a subset of transcription factories. The self-organising principle applied to transcription factories would

predict this; just as the local concentration of active RNA Polymerase II is elevated by the clustering of transcribed units, the local concentration of transcription factors that bind to those transcribed units will also be elevated [76]. Genes that diffuse to a transcription factory already engaged with other genes that share the same factors are more likely to engage and be transcribed themselves [77]. Indeed, genes regulated by common transcription factors appear to cluster within specific chromosomes in yeast [78] and testis-specific genes are found clustered in *Drosophila* [79] - genes clustered in linear sequence are more likely to associate in three dimensions.

Some evidence that such specialised transcription factories may exist within mammalian cells came from Osborne *et al.* [75]. They showed that approximately one quarter of actively transcribing *Myc* alleles shared a transcription factory with *Igh*, over double the rate of colocalisation found with the control genes tested. Using DNA-FISH they showed an overall reduction in spacing between *Myc* and *Igh* alleles upon B cell stimulation, suggesting that *Myc* alleles are being specifically recruited to transcription factories containing transcribing *Igh* [75]. Xu and Cook later demonstrated *in-vitro* that cells transfected with thousands of plasmids containing different transcriptional units clustered together at a handful transcription factories, segregated according to their promoter and the presence of an intron [80].

Perhaps the best evidence for the existence of specialised transcription factories came in a publication by Schoenfelder *et al.* in 2010 [81]. They used a variety of techniques to investigate the nuclear localisation of erythroid genes. A genome-wide screen of genes associating with *Hba* and *Hbb* at transcription factories showed enrichment for genes with CACC motifs capable of binding the erythroid-specific transcription factor Klf1. Using immunofluorescence they found that nuclear Klf1 foci overlap with active RNA Polymerase II foci, suggesting that a subset of transcription factories are enriched for Klf1 in erythroid tissues (Fig 1.5). Co-localising Klf1 dependent genes associated with these Klf1 specific transcription factories at a high frequency, and a number of these gene associations were lost upon the knockout of Klf1 expression. These data highly suggest that a network of Klf1 specific transcription factories exist within mouse erythroid tissues, and that Klf1 specific genes are preferentially recruited to these

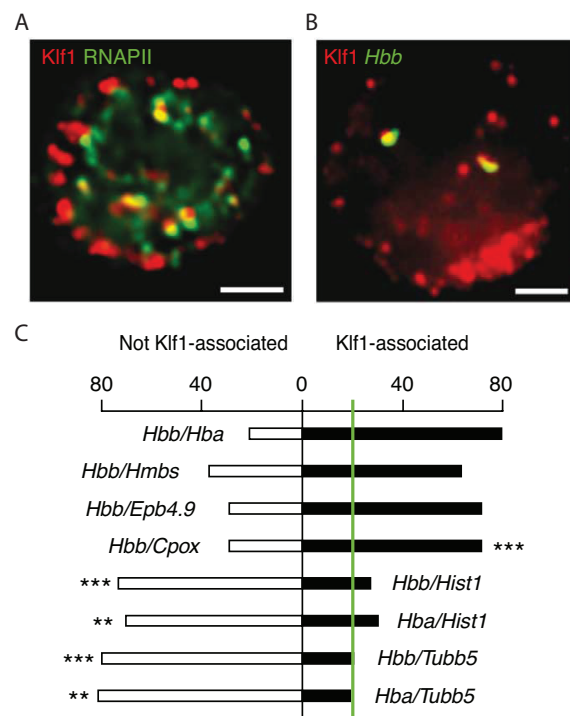


Figure 1.5 – Klf1 specialised transcription factories. (A) Immunofluorescence showing Klf1 (red) overlapping with a subset of RNA Polymerase II transcription factories (green). (B) RNA immuno-FISH showing transcribing *Hbb* alleles (green) co-localising with Klf1 (red). (C) Percentages of co-localising transcript pairs colocalisation with Klf1 associated transcription factories (right, black bars) and non-Klf1 associated factories (left, white bars). Expected level of association due to chance shown by green bar. Scale bars show in (A) and (B) represents 2 μ m. Taken from Schoenfelder *et al.* [81].

sites [81]. If such networks are a general feature in mammalian nuclei, they could be a key driving force in the organisation of the genome.

It is important to note that the evidence above does not describe *exclusively* specific transcription factories, factories which are incapable of transcribing genes which are not controlled by Klf1. Indeed, although statistically significant, the gene associations detected still suggest a population bias only. Specialised transcription factories are certainly not required for the transcription of the eukaryotic genome, but may aid efficiency and affect nuclear organisation and function.

1.5 Chromatin interactions in three dimensions

1.5.1 Promoter and enhancer interactions

The classical model of gene regulation and their promoters began with the characterisation of the *Lac* operon in *Escherichia coli* in 1961 [82]. Three genes required for the digestion of lactose are controlled by the binding of a repressor protein which blocks transcription by RNA Polymerase II. This repressor can form a tetramer and bind two region simultaneously, requiring a topological loop in the chromatin [83].

A number of Eukaryotic genes were later found to require distal enhancer elements [84] prompting speculation that the looping of chromatin to allow direct interaction between sequences may be a common mechanism [85]. Direct evidence for *in-vivo* chromatin interactions was provided in 2002 with two papers studying the mouse *Hbb* locus [86, 87]. The locus contains five beta-like genes β , γG , γA , δ and ϵ , which are arranged in order of their expression. Approximately 50 kilobases upstream is a locus control region (LCR), containing multiple DNase hypersensitive sites (HS1-6) [88].

Carter *et al.* used a novel technique called RNA TRAP (tagging and recovery of associated proteins) which localises horseradish peroxidase to the site of nascent RNA production of a specific gene. The horseradish peroxidase catalyses biotinylation of nearby chromatin which can be purified and quantified by quantitative real-time PCR (qPCR). They showed a 15-fold enrichment of biotin over the HS2 region of the LCR when the actively transcribing *Hbb-b1* transcript was used for the horseradish peroxidase localisation, demonstrating that this region of the LCR is in very close proximity to the *Hbb-b1* gene [86].

Tolhuis *et al.* used chromatin-conformation-capture technique [89] (3C, for discussion of this technique see Section ??) to study the same locus and also found evidence for interaction between the LCR hypersensitive regions and the *Hbb* locus. Importantly they showed that in mouse foetal brain tissue, where the *Hbb* locus is not expressed, the chromatin adopted a linear conformation without any looping [87]. The group went on to show that each gene contacts

the LCR as it is expressed, supporting a system where the developmentally controlled genes must contact an enhancer within the LCR in order to be expressed [90]. Palstra *et al.* also showed that erythroid progenitor cells dedicated to the lineage but not yet expressing beta-like globin genes form a 'poised' structure, contacting the LCR but not engaging strongly with the HS2 enhancer [90].

1.5.2 Chromatin hubs

The beta globin locus has become a model system for the active chromatin hub (ACH) model [91], describing a system where loops of chromatin containing elements capable of controlling the expression of genes are held in close three-dimensional space to the genes that they control. Other gene clusters have been shown to behave in a similar manner, notably the T_H2 locus and *Hox* clusters.

The T_H2 locus control region is involved in the transcriptional control of cytokine genes IL4, 5 and 13 [92]. Spilianakis *et al.* showed that these genes co-associate in T cells, NK cells, B cells and fibroblasts, despite not being expressed in B cells and fibroblasts. In T cells and NK cells this core unit additionally associates with the T_H2 LCR and the genes are expressed [93]. The authors suggest that this mechanism allows the coordinate expression of the gene cluster in a controlled manner.

The *Hox* genes are master regulators of gene transcription and are responsible for the creation of vertebrate segments during development. *Hox* gene clusters *A* to *D* are transcribed in sequence as development of the embryo progresses and have been observed decondensing and looping out from their chromosome territory upon activation [58]. Noordermeer *et al.* studied the *Hox* clusters in three different mouse embryonic day 10.5 tissues; forebrain, anterior trunk and posterior trunk [94]. They found that the *Hoxd* cluster formed a discrete domain in forebrain, where it is inactive. In anterior and posterior trunk the *Hoxd* cluster is active, but different genes are transcribed. In both tissues they found the cluster to form two distinct compartments correlating with an inactive and active regions. Using 4C, a technique based on chromosome conformation capture (3C), they showed that genes move from the inactive to active compartment as they are activated, correlating with active histone marks [94].

1.5.3 Interactions in *trans*

As the role for distal enhancers has become more established, evidence has been uncovered for increasingly distant interactions. Lettice *et al.* used data from known chromosomal translocations resulting in polydactyly to find and characterise an enhancer element for the *SHH* gene over a megabase upstream, within an intron of another gene [95]. Spilianakis *et al.* went on from characterising the *intrachromosomal* interactions of the T_H2 LCR to show that the same locus forms *interchromosomal* interactions [96]. Naïve T cells can differentiate into either TH1 or TH2 cells, depending on the stimulus received. This is defined by the expression of either IFN- γ or IL-4. Spilianakis *et al.* showed

that the *Ifng* gene promoter can interact with the T_H2 LCR to stimulate *Ifng* expression whilst inhibiting *IL4* expression. These two loci reside on chromosomes 10 and 11 and their interaction is the first known interchromosomal interaction to regulate gene expression [96]. A number of studies have since shown specific interchromosomal contacts involved in a number of processes ranging from X-inactivation to genomic imprinting, showing that these interactions play an important role in chromatin biology [97].

1.5.4 Global interaction maps

Our understanding of the three-dimensional organisation of the genome has advanced in leaps and bounds during the last decade largely because of the development of chromatin conformation capture (3C) and its derivatives [98]. There are a large number of 3C variants, but they can be grossly categorised into four classes based on how many loci can be interrogated in a single experiment: one-to-one (3C, quantitative 3C), one-to-all (4C, e4C, ACT), many-to-many (5C) and all-to-all (ChIA-PET, Hi-C, TCC).

The recent development of all-to-all methods has allowed the confirmation of the entire genome to be probed in a single experiment. This approach has many advantages; such an unbiased approach allows the detection of unexpected interactions, additionally many associations can be probed in parallel allowing a far higher rate of data collection. Whilst these techniques have had great impact on the field, they are currently limited by the depth of sequencing that is achievable with today's technology. To address this Sanyal *et al.* recently published a paper as part of the ENCODE project describing the interaction profiles of 628 transcription start sites (TSS) and 4535 surrounding fragments, representing approximately 1% of the genome [99]. To achieve the resolution required for the robust detection of promoter-element interactions, Sanyal *et al.* used 5C, a many-to-many technique that uses a library of oligonucleotides with common adapters to anneal to 3C products and create a library capable of being sequenced. They sequenced libraries from three ENCODE cell lines: K562, HeLa-S3 and GM12878. Only a small proportion of the looping interactions uncovered were shared between the three cell types, with ~60% of interactions being unique to a single cell line. The majority of TSS looping interactions could be classified as interacting with enhancer elements, promoters or regions bound by the structural protein CTCF. Looping interactions with enhancer elements were significantly enriched for actively expressed TSS, demonstrating the importance of three-dimensional chromatin contacts in the regulation of gene expression. Whilst this study provides valuable information in the global patterns of gene interactions in the nucleus, it is also a valuable resource to be used by other research groups. It is becoming increasingly clear that to fully understand the biology of chromatin and gene expression, we must understand the involvement of the structure of the nucleus.

1.6 What drives nuclear organisation?

As our understanding of the structure of the nucleus evolves, an increasing number of structural features and patterns are uncovered. Teasing apart correlation and causation to find the driving forces behind nuclear organisation is not an easy task and remains a hot topic within the field.

1.6.1 Transcription

The discovery of transcription factories has changed our view of nuclear organisation substantially. If genetic templates are mobile and transcription factories are fixed, then the nucleus may be able to use the process of transcription as a tool to fold the genome into specific conformations.

Kimura *et al.* quantified the amount of stable RNA Polymerase II in HeLa cells [100] adding to the work by Jackson *et al.* demonstrating the stability of nascent transcripts and Polymerase in the nucleus [69, 70]. Mitchell and Fraser later demonstrated that RNA Polymerase II transcription factories remain in the absence of transcription, though gene association with factories is ablated if transcription initiation is inhibited [101]. This data, along with the observation that genes are recruited to pre-existing transcription factories upon activation [61], support a model whereby transcription factories are attached to a relatively immobile nuclear substructure. If this is the case, RNA Polymerase II can act as a motor, dragging template chromatin through the nucleus as it is transcribed, powered by the removal of phosphate groups during RNA synthesis [68]. Yin *et al.* measured the force produced by a single *E. coli* RNA Polymerase using an immobilised enzyme transcribing a template bound to a polystyrene bead held by optical tweezers [102]. The force they measured (14 pN) makes RNA Polymerase the most powerful known biological motor. Papantonis *et al.* demonstrated the potential of RNA Polymerase to pull chromatin transcripts through the nucleus *in vivo* by used 3C to measure the change in association between regions of DNA after activation of the TNF α gene [103]. They found that as the gene was transcribed, downstream regions of chromatin progressively came into contact with other transcribing regions at the transcription factory.

1.6.1.1 Not finished - write about transcription stop cassette in T cells

A good example of transcription actively organising a structural motif is within the immunoglobulin heavy-chain locus. The *Igh* locus contains variable (V_H), diversity (D_H), joining (J_H) and constant (C_H) coding elements which must contact each other for the process of recombination. The *Igh* locus spans 3 Mbp and forms a rosette-like structure [104]. This structure appears to be driven by association at a transcription factory, with low level antisense transcription of the V_H gene segments prior to recombination [#REF]. Interestingly, the level of antisense transcription correlates

with the distance of the V_H gene from the C_H region, and if this transcription is inhibited, the V_H gene usage is biased towards V_H genes closer to the C_H region in the linear DNA template [105]. This data suggests a model whereby the antisense transcription of the V_H genes is required to bring them into contact with the C_H region to give equal probability of recombination regardless of the linear spacing of the genes.¹

The role for transcription in the organisation of the genome is further supported by the pervasive nature of transcription. As much as 15% of the human genome is actively transcribed [105], including many enhancers [106, 107]. In the recent ENCODE study of TSS interactions, Sanyal *et al.* found that enhancer elements looping to a TSS were significantly more likely to express enhancer RNAs [99], supporting a model that the transcription of a chromatin template could be responsible for the formation of chromatin loops.

An attractive model for larger scale genome organisation revolves around the transcription of housekeeping and tissue-specific genes. Lercher *et al.* have shown that genes with high expression in multiple tissue types have a propensity to be present in clusters within the genome [108] and the mouse alpha globin locus has been shown to assemble at a transcription factory already transcribing a cluster of housekeeping genes in erythroid cells [109]. These housekeeping genes have been shown to bind stably to the nuclear matrix, whereas the alpha globin genes are eluted in a high salt extraction [110]. These studies may point to a model of the nucleus where persistently transcribed housekeeping genes are responsible for forming transcription factories, with tissue specific genes being recruited to these sites upon activation [110]. Such a system could have far reaching consequences, such as chromosome conformation being determined by housekeeping gene hubs.

1.6.2 CTCF

- Insulators. Chicken beta globin.
- ChIA-PET - [111]
- CTCF not always marking insulators - interactions skipping CTCF sites in Sanyal *et al.* [99]
- Talk about CTCF bordering domains, Hi-C papers

1.6.3 Cohesin

- Introduction to standard stuff about sister chromatid adhesion
- Basically all of Suzie's papers showing that Cohesin binds CTCF in interphase

¹This will need to be re-written after I've found the right references.

1.6.4 Actin and myosin

- Nuclear actin and myosin and stuff.

1.6.5 Replication

- Should probably write quite a bit here as haven't talked about it at all yet
- Similarity to transcription / factories etc

1.6.6 The self-organisation model

- Cook paper on entropy basics, nature of everything wanting to cluster. [76, 71]
- Transcription factor networks (Bertie).
- Mention specialised transcription factories again?

1.7 The annotated genome

- Talk about how wonderful systems biology is
- Integration of datasets - ChIP Seq is two dimensional and needs interaction data to be understood
- Correlation between different marks gives the full picture
- Genome wide studies paving the way for a different type of analysis
- Talk about the systems biology of networks
 - This is where I extoll the virtues of Bertie's work looking at transcription factor networks
 - Use *cis* regulatory modules as an example? Then shoot down with Dekker 5C paper? Check Bertie doesn't love them first (I think he might)

1.7.1 The ENCODE project

- Description of ENCODE project
- Couple of key findings
- Mention value of ENCODE datasets beyond initial publications

1.8 Chromosomal translocations

1.8.1 Formation of chromosomal translocations

Since the development of chromosome banding microscopy, it has been known that the genetic material within cancerous cells is frequently disrupted [112]. Chromosomal translocations involve the rearrangement of genetic material between non-homologous chromosomes through the formation and aberrant repair of two double strand breaks (DSBs). The resulting product can be a straight swap without any loss or gain of sequence (balanced translocation) or can result in deletions or gain of material (unbalanced translocation).

1.8.1.1 DSB repair

Double strand breaks can be caused by exogenous damage (ionizing radiation, free radicals) and endogenous damage (physiological programmed DSBs). As many as 1 million DNA lesions are formed per cell per day [113] and can be highly deleterious to the cell, causing loss of genetic material, translocations and ultimately cell death if left unrepaired. Mammalian cells have an array of different DSB repair pathways that depend on the cellular context and type of lesion [114].

Homologous recombination (HR, Fig 1.6B) repairs double strand breaks by using long regions of homology on undamaged sister chromatids or homologous chromosomes. HR is primarily active during cell replication and rarely leads to serious chromosomal rearrangements or translocations, although ‘crossing-over’ can sometimes occur, whereby two sections of homologous chromosomes switch to their partner chromosomes (a process that is encouraged in meiosis with the formation of chiasmata).

Single strand annealing (SSA, Fig 1.6C) creates a short region of single stranded DNA at the site of the DSB which is used to search for regions of microhomology, usually repeat sequences in the same orientation. These regions anneal and any excess single stranded tails are removed to allow the nicks to be ligated. SSA is prone to introducing deletions into the genome, and can form chromosomal translocations if two sequences with similar repeat regions are nearby, and both suffer simultaneous DSBs.

Non-homologous end joining (NHEJ, Fig 1.6A) is a last resort for the cell, whereby any two adjacent DSBs are ligated, needing little or no sequence homology. This pathway is extremely prone to errors as it is capable of joining any two strands of DNA without knowledge of their identity.

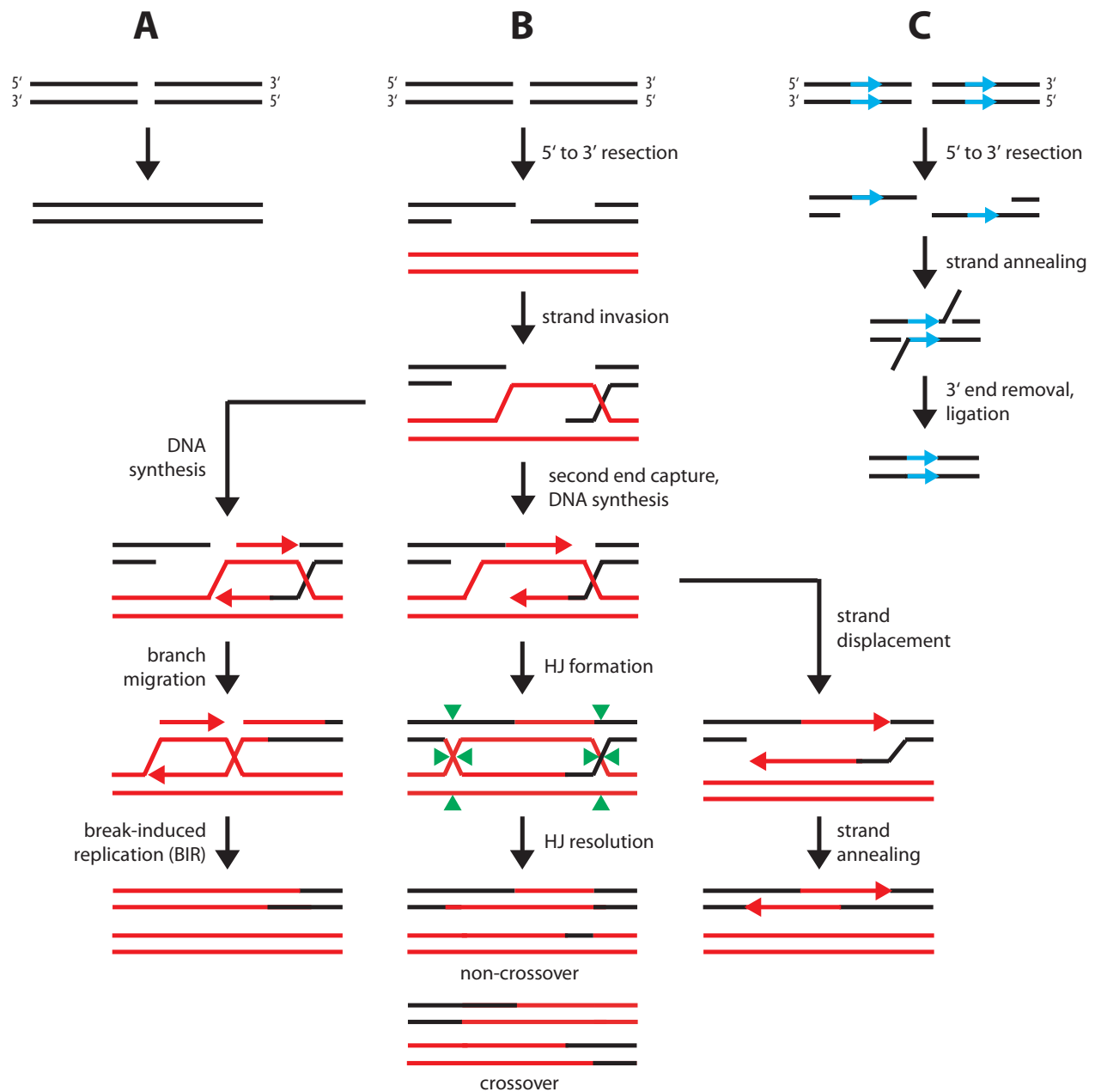


Figure 1.6 – DSB repair. Three double strand break repair pathways. **(A)** NHEJ (Non-Homologous End Joining) requires little or no sequence homology and joins any two nearby ends. **(B)** HR (Homologous Recombination) uses a strand of homologous DNA on a duplicated interphase chromosome or sister chromatid to reconstitute sequence before joining. Although it only requires a single double strand break to generate a crossover, the extensive region of homology needed makes errors rare. **(C)** SSA/NHEJ (Single Strand Annealing / Non-Homologous End Joining) searches for regions of homology, such as repeat sequences, and deletes any sequence between. From Longhese *et al.*, 2006 [114].

1.8.2 Chronic myeloid leukaemia

Chronic myeloid leukaemia (CML) is one of the best studied cancers, despite having a relatively low occurrence [115].

The term 'leukaemia' was based on CML patients in the mid 1800s [116], and it was the first cancer for which a direct

causative genotype was found [117]. A number of standardised therapies exist for CML and up to 87% of patients achieve complete cytogenetic remission [115], making it one of the success stories of cancer research.

1.8.2.1 The Philadelphia chromosome

The genetic abnormality associated with CML was first identified in 1960 by Nowell *et al.* [117]. They noticed the presence of an abnormal chromosome and called it the Philadelphia chromosome, or Ph chromosome. The chromosome was initially thought to be the result of a deletion until chromosome banding enabled its identification as a product of a translocation in 1972 [118]. The breakpoints within chromosomes 9 and 22 were subsequently identified as being within the *c-ABL* and *BCR* genes, respectively. The *c-ABL* gene was named as such because of its similarity to the Abelson murine leukaemia virus gene *v-ABL* [119]. Less is known about the *BCR* gene, so named because it was found at the Breakpoint Cluster Region of chromosome 22 (not related to the B-cell receptor, often referred to as the BCR).

1.8.2.2 *BCR* and *ABL*

The *ABL* gene encodes a tyrosine kinase which is ubiquitously expressed in mammalian cells. Involved in cell cycle regulation, it is thought to be involved in several cell signalling pathways [120]. Less is known about the endogenous function of the *BCR* protein; it contains a serine-threonine kinase and has GTPase activity but its function is not known [120]. The t(9;22)(q34;q11) translocation creates a fusion protein lacking the SH2 domain of *ABL* which normally regulates its activity. This results in constitutively active tyrosine-kinase activity which drives oncogenesis [120]. Relatively little is known about the mechanism of translocation formation, though ionising radiation is known to be a risk factor [121, 122].

Identification and characterisation of the *BCR-ABL* translocation and its product led to the development of a number of kinase inhibitors such as the drug imatinib, which achieve excellent success in the treatment of CML [115].

1.8.3 Mixed lineage leukaemia

Acute leukaemia is typically categorised as either Acute Lymphoblastic Leukaemia (ALL) or Acute Myeloid Leukaemia (AML) according to which lineage of blood cells are cancerous. ALL is defined by the uncontrolled proliferation of lymphoblasts, precursors to lymphocytes which differentiate into B cells, T cells and NK cells. ALL was estimated to account for approximately 12% of all leukaemia cases in the US in 2008 with around five and a half thousand cases occurring annually. ALL is the most common form of cancer in children aged under fourteen [123]. AML is a cancer of the myeloid lineages of blood cells, responsible for approximately 30% of all cases of leukaemia in the US for 2008 with a higher mortality rate than ALL (66% and 27% mortality for AML and ALL, respectively) [123]. The incidence

of AML increases with age; the median age of AML diagnosis between 2002 and 2006 in the US was 67 years, and the median age of mortality was 72 [123].

The biology of leukaemia is not always as discrete as the classification however, some patients present with expansion of both lymphoid and myeloid lineages [124]. Common genetic abnormalities found in these patients include the t(9;22)(q34;q11) Philadelphia chromosome and structural changes in 11q23 [124]. Chromosomal translocations within band q23 of chromosome 11 have been implicated in both AML and ALL, all involve the gene *MLL* (also known as *ALL-1*, *Htrx*, *HRX*) identified by Ziemer-van der Poel *et al.* in 1991 [125]. Translocations involving the *MLL* gene are found in over 70% of all infant leukaemias [126] and approximately 10% of adult AML cases [127]. The translocation correlates with poor patient prognosis and is of high clinical interest [128].

In 2002, Armstrong *et al.* showed that acute lymphoblastic leukaemias containing a translocation within the *MLL* gene have a unique expression profile that is different to ALL and AML, and suggests an origin within a less committed progenitor cell which can produce cells in both the myeloid and lymphoid cell lineages [129]. They suggest that these leukaemias are substantially different from AML and ALL and deserve a new, distinct, class of leukaemia called Mixed Lineage Leukaemia (MLL). Further cytogenetic studies have supported this theory; leukaemic cells in MLL have been found to express cell surface antigens normally present on both myeloid and lymphoid cells, such as CD14 and CD19 [127].

The suggestion that MLL is initiated within a progenitor cell is in line with previous evidence showing that some cases of ALL and AML are initiated within undifferentiated haematopoietic stem cells (HSCs) [130, 131]. HSCs are present within the CD34⁺ progenitor cell population, accounting for approximately 3% of normal human bone marrow and 0.3 to 0.5% of human cord blood mononuclear cells [132]. They are long lived and capable of self-renewal, differentiating into lineage restricted progenitors and eventually mature terminally differentiated white blood cells. HSCs are necessary for the long term maintenance of the haematopoietic system and are commonly used to repopulate bone marrow after myeloablative therapy, as well as in the treatment of a number of other disorders such as autoimmune, cardiac and vascular diseases [133]. Cancer stem cells are thought to be present in both leukaemias and solid tumours [134] and HSCs are a probable founder population due to their ability to self-renewal [135]. The concept of a small pool of cancer stem-cells driving the large heterogeneous pool of cancer cells has a number of implications for treatment - these are the cells that must be targeted for the efficient and long lasting cure of cancer [135].

1.8.3.1 The MLL protein

The MLL protein is a H3K4 methyltransferase involved in the positive regulation of global gene regulation, including the maintenance of expression of the *Hox* genes [136]. *MLL* is required for embryonic haematopoiesis [137] and adult bone marrow maintenance [138]. MLL is a mammalian homologue of the *Drosophila melanogaster* trithorax complex and is thought to bind DNA via an AT-hook domain [139] and zinc finger domain [140]. It's thought that the zinc

finger domain targets MLL to unmethylated CpG islands DNA [140] and ChIP studies have shown that MLL binds to a subset of transcribed genes [141]. MLL binds promoters and gene bodies, associating tightly with RNA Polymerase II [141].

All known MLL fusion proteins contain exons 8-13 of *MLL* and in-frame exons of a partner gene [127]. Fusion proteins always retain their AT-hook and zinc-finger CxxC motifs, which are essential for their transforming potential [142]. The H3K4 methyltransferase domain of MLL is often lost in fusion proteins [127], despite this the fusion proteins can drive constitutive expression of *HOXA9* and *MEIS1*, which if over-expressed together in the absence of a MLL fusion protein give a similar phenotype [143].

1.8.3.2 The *MLL* gene

Translocations within the *MLL* gene are found within approximately 10% of all human leukaemias [144]. There are 87 documented *MLL* translocation partners of which 51 have been characterised at the molecular level [145]; the five most frequent translocation partners, *AF4*, *AF9*, *ENL*, *AF10* and *AF6* account for approximately 80% of cases [145]. Translocations within the *MLL* gene usually occur within an 8.3 kb *BamHI* fragment known as the breakpoint cluster region [146]. This region contains exons 5-11 as well as a number of repeat regions, notably 8 direct *Alu* SINE repeats, 5 direct L1 and L2 LINE repeats and 2 MER elements, as well as a number of putative Topoisomerase II binding sites and a SAR/MAR [147]. An internal promoter is also present within the murine *Mll* breakpoint cluster region, correlating with etoposide-induced DSBs [148], DNase I hypersensitive sites [149] and histone modifications associated with transcription [150].

1.8.3.3 Mechanisms of *MLL* translocation formation

A clue to how translocations form within the *MLL* gene comes from the observation that 11q23 translocations are especially prevalent in therapy-related leukaemias - secondary leukaemias that develop in patients after treatment for a primary cancer with topoisomerase II inhibitors [127]. Topoisomerases are found in all eukaryotic nuclei and are able to relieve supercoiling and promote chromosome disentanglement [151]. They function by binding DNA, forming a transient double strand break and passing another strand of DNA through the gap, before ligating the double strand break.

Topoisomerases are important in transcription, which creates supercoils as template DNA is processed through static RNA Polymerase II enzymes [152]. DNA topoisomerase II associates with gene promoters [153] and is required for the transcription of genes longer than 3 Kbp in yeast [154]. Topoisomerase II induced double strand breaks have been implicated in the regulation of certain genes via the assembly of transcription complexes and changes in chromatin structure [155].

DNA topoisomerase II inhibitors are commonly used as chemotherapeutic agents and work by decreasing the

ligation rate, disrupting the cleavage / ligation equilibrium. This leads to an increase in DNA cleavage and an accumulation of double strand breaks, triggering the cell DNA damage response and leading to cell death by apoptosis [156]. Topoisomerase II inhibitors are widely used chemotherapeutic agents, effective against a range of malignancies including small-cell lung cancer and gonadal tumours [157].

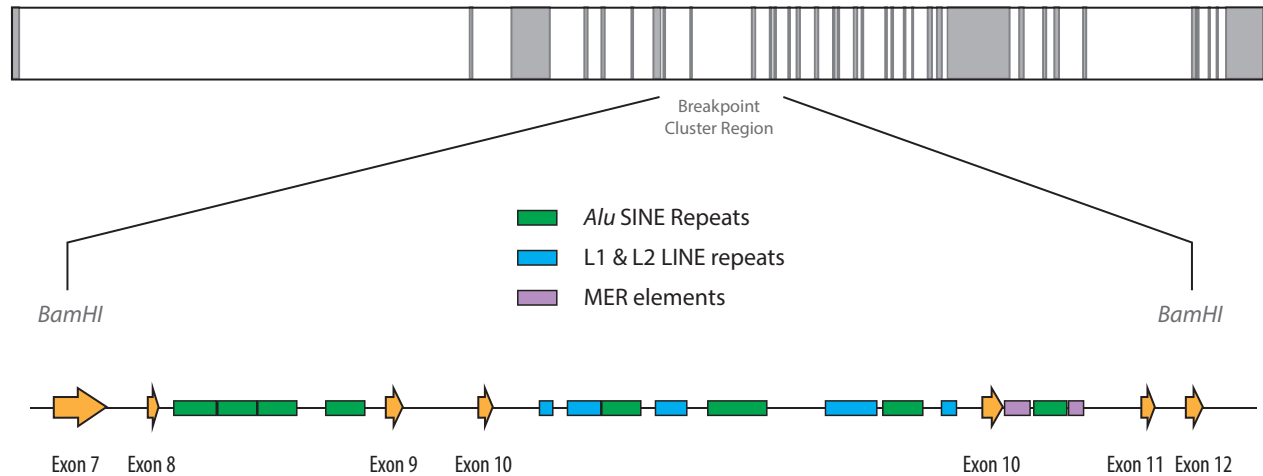


Figure 1.7 – Diagram of the *MLL* breakpoint cluster region.

The *MLL* breakpoint cluster region contains a number of putative topoisomerase II binding sites, prompting a number of groups to hypothesise a mechanism of topoisomerase inhibitor related translocations. Libura *et al.* showed that CD34⁺ HSPCs exposed to etoposide formed 11q23 chromosomal aberrations typical of those seen in clinical samples [158]. They went on to show that such exposure to etoposide increased the proliferative potential of the cells in a bone marrow graft assay using immune-deficient mice [132]. This data supports a model whereby poisoned topoisomerase II may create double strand breaks within the *MLL* gene, allowing the NHEJ machinery to aberrantly repair the locus due to microhomology found in the nearby repetitive elements (Fig 1.7).

1.8.4 Effect of nuclear organisation on translocation formation

1.8.4.1 Breakage first and contact first models

For a chromosomal translocation to form, two double strand breaks must exist simultaneously and be adjacent in three-dimensional space. Two models have been described to explain how DSBs may meet in the nucleus – the breakage first model and the contact first model. The breakage first model states that double strand breaks can form anywhere and are able to freely diffuse in the nuclear space. They undergo large scale movement through the nucleus until they meet and are joined. The contact first model states that the two double strand breaks form in sequences already close to each other within the nuclear space. Large scale chromatin movements are not required for the two DSBs to meet.

In support of the breakage first model Aten *et al.* showed that a linear series of DSBs formed simultaneously in

HeLa nuclei clustered [159], however, others have argued that this may be the result of a higher degree of chromatin mobility along the ion beam trajectory used to generate the DSBs [160]. A larger body of evidence supports a contact first model; DSBs created with ultra-soft X-rays by Nelms *et al.* remained in a fixed position for several hours after the damage was caused [161] and Jakob *et al.* used live cell microscopy to visualise proteins involved in DNA damage signalling and repair. After accumulation at sites of DNA damage caused by heavy ion impacts the foci exhibited only a small degree of movement [160]. Soutoglou *et al.* showed a high degree of positional stability of DSB ends created by endonuclease digestion, with a greater degree of local diffusion seen in the absence of the Ku80 DNA-end binding protein [162].

1.8.4.2 Chromosome Territories

The large scale organisation of chromosome territories has been implicated in translocation frequency by a number of studies. Kozubek *et al.* showed that chromosomes 9 and 22 were found in the centre nuclei more frequently than would be expected by chance in lymphocytes, T- and B-cells, HL60 cells and bone marrow cells [163]. They used neutron irradiation to show that transfer of genetic material was much higher than that found with chromosome 8, which was found towards the nuclear periphery [163]. Parada *et al.* investigated the positions of chromosomes 12, 14 and 15 in a mouse lymphoma cell line and mouse splenocytes [164]. They found that two translocated chromosomes preferentially paired together in the nucleus of the cell line as well as in normal cells not containing the translocation. [164]. Parada *et al.* went on in a further study to examine the positioning of a larger range of chromosomes in a number of different tissue types [40]. They found that chromosome pairing was tissue specific and correlated with the occurrence of tissue-specific translocation events [40].

In 2006, Branco and Pombo published a study where they developed a new technique known as cryo-FISH to enhance the resolution of chromosome territory detection [64]. This technique had sufficient resolution to study the degree of intermingling between chromosome territories and they found that the degree of intermingling between chromosome pairs correlated strongly with their propensity to form translocations when subjected to radiation [64].

1.8.4.3 Transcription Factories

Mounting evidence has shown that in addition to CT pairing, the position of specific genes involved in chromosomal translocations are frequently found in close proximity [165, 166]. As discussed in Section 1.6.1, localisation at transcription factories can drive organisation within the genome. Osborne *et al.* used DNA- and RNA-FISH to study the localisation of the *Myc* proto-oncogene upon activation of B cells [75]. They showed that *Myc* on mouse chromosome 15 is dynamically recruited to an existing transcription factory and that this transcription factory was preferentially occupied by *Igh* on mouse chromosome 12. This preferential association at transcription factories leads to a model whereby genes sharing transcription factories are predisposed to the formation of chromosomal translocations [75].

Similar transcriptional associations have been observed for *MLL* and its frequent translocation partners *AF4* and *AF9* [167].

1.9 Hypothesis and aim of thesis

- State hypothesis - Preferential gene co-association within transcription factories predisposes specific gene pairs to the formation of chromosomal translocations.
- Aims - investigate the genome-wide association networks of specific oncogenes and see if gene pair co-localisation frequency matches up with gene-pair translocation frequency
 - Do this in an unbiased manner (not purely by RNA- and DNA-FISH), mention biased nature of such studies
 - Do this with a genome wide technique
 - Do this in the cell type where we think the translocations are taking place

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