The three-dimensional (3D) organization of telomeres during cellular transformation

By

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Abstract

Statement of Problem

Telomere dynamics in the three-dimensional (3D) space of the mammalian nucleus plays an important role in the maintenance of genomic stability. However, the telomere distribution in 3D nuclear space of normal and tumor cells was unknown when the study was initiated.

Methods

Telomere fluorescence in situ hybridization (FISH) and 3D molecular imaging, deconvolution, and analysis were used to investigate telomere organization in normal, immortalized and tumor cells from mouse and human cell lines, and primary tissues.

Results

Telomeres are organized in a non-overlapping manner and in a cell-cycle dependent fashion in normal cells. In the late G2 phase of cell cycle, telomeres are assembled into a flattened sphere that is termed the telomeric disk. In contrast, the telomeric disk is disrupted in the tumor cells. Moreover, telomeric aggregates (TAs) are found in tumor cells. Conditional c-Myc over-expression induces telomeric aggregation leading to the onset of breakage-bridge-fusion cycles and subsequent chromosomal abnormality.

Conclusions
Telomeres are distributed in a nonrandom and dynamic fashion in the 3D space of a normal cell. Telomeric aggregates are present in cells with genomic instability such as tumor cells and cells with deregulation of c-Myc. Consequently, TA can be a useful biomarker for research in cancer and other disease processes.
Acknowledgements

First of all, I want to thank my family for standing behind me without failing during the great turbulences I endured in my pursuit of my training and research.

“If I have seen further it is only by standing on the shoulders of giants.” Sir Isaac Newton, in letter to Robert Hooke.

The work done in my thesis is made possible due to the collaboration with many giants, from different scientific disciplines and different parts of the world, with the same spirit of answering fundamental questions of any novel scientific observation – “what is it” and “why is it”.

Much gratitude goes to my supervisor and committee members, Drs. Sabine Mai, Paul Kerr, James Davie, Robert Shiu, and Edwin Kroeger. You have given me different sets of skills, from critical thinking to performing head and neck surgery, that will be forever treasured by me.

Support from the Department of Physiology is much appreciated. Gail, I thank you for all your time and encouragement in getting me to completion of my study.

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Dedication

To members of my family, friends and mentors from various institutions including University of Manitoba, The Johns Hopkins University, National Institutes of Health, and Delft University.
Lists of Tables and Figures

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List of figures – see publications Chuang et al, 2004 and Louis et al, 2005


**List of Abbreviations**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>3D</td>
<td>Three-dimensional</td>
</tr>
<tr>
<td>AIDS</td>
<td>Acquired immune deficiency syndrome</td>
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<tr>
<td>ALT</td>
<td>Alternative lengthening of telomerase</td>
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<td>APB</td>
<td>ALT-associated promyelocytic leukemia nuclear bodies</td>
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<td>ATM</td>
<td>Ataxia telangiectasia mutated</td>
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<tr>
<td>BAX</td>
<td>BCL2-associated X protein</td>
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<td>BBF cycle</td>
<td>Breakage-bridge-fusion cycle</td>
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<td>BCL2</td>
<td>B-cell CLL/lymphoma 2</td>
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<td>b-FGF</td>
<td>Basic fibroblast growth factor</td>
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<tr>
<td>BIR</td>
<td>Break-induced replication</td>
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<tr>
<td>CCD</td>
<td>Charge-coupled device</td>
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<tr>
<td>CDKN2A</td>
<td>Cyclin-dependent kinase inhibitor 2A</td>
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<tr>
<td>CLEM</td>
<td>Controlled light exposure microscopy</td>
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<tr>
<td>COX-2</td>
<td>Cyclooxygenase-2</td>
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<td>CT</td>
<td>Chromosome territory</td>
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<tr>
<td>DNA</td>
<td>Deoxyribonucleic acid</td>
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<tr>
<td>EBV</td>
<td>Epstein Barr virus</td>
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<tr>
<td>EGFR</td>
<td>Epidermal growth factor receptor</td>
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<tr>
<td>FISH</td>
<td>fluorescence <em>in situ</em> hybridization</td>
</tr>
<tr>
<td>GAR1</td>
<td>GAR1 ribonucleoprotein homolog (yeast)</td>
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<tr>
<td>H cells</td>
<td>Hodgkin cells</td>
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<tr>
<td>HER2/neu</td>
<td>Human epidermal growth factor receptor 2</td>
</tr>
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<td>HPV</td>
<td>Human papilloma virus</td>
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<tr>
<td>H-Ras</td>
<td>Harvey sarcoma viral oncogene homolog</td>
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<td>hTERC</td>
<td>Human telomerase RNA gene</td>
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<td>Abbreviation</td>
<td>Description</td>
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<tr>
<td>hTERT</td>
<td>Human telomerase reverse transcriptase</td>
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<td>IC</td>
<td>Interchromatin compartment</td>
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<tr>
<td>K-Ras</td>
<td>Kirsten sarcoma viral oncogene homolog</td>
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<tr>
<td>MALT</td>
<td>mucosa-associated lymphoid tissue</td>
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<tr>
<td>MMP</td>
<td>Metalloproteinase</td>
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<tr>
<td>NHP2</td>
<td>NHP2 ribonucleoprotein homolog (yeast)</td>
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<td>NOP10</td>
<td>NOP10 ribonucleoprotein homolog (yeast)</td>
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<tr>
<td>PEV</td>
<td>Position effect variegation</td>
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<td>PML</td>
<td>Promyelocytic leukemia</td>
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<td>POT1</td>
<td>Protection of telomeres protein 1</td>
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<tr>
<td>PTEN</td>
<td>Phosphatase and tensin homologue</td>
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<tr>
<td>RAD52</td>
<td>RAD52 homolog (S. cerevisiae)</td>
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<tr>
<td>RAP1</td>
<td>Repressor activator protein 1</td>
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<td>RB</td>
<td>Retinoblastoma</td>
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<tr>
<td>RNA</td>
<td>Ribonucleic acid</td>
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<tr>
<td>RS cells</td>
<td>Reed-Sternberg cells</td>
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<tr>
<td>snRNP</td>
<td>Small nuclear ribonucleoprotein</td>
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<td>TA</td>
<td>Telomeric aggregates</td>
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<tr>
<td>TGF-α</td>
<td>Transforming growth factor alpha</td>
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<td>TIN2</td>
<td>Second TRF1 interacting partner</td>
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<td>TMM</td>
<td>Telomere maintenance mechanism</td>
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<td>TP53</td>
<td>Tumor protein 53</td>
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<td>TP73</td>
<td>Tumor protein 73</td>
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<tr>
<td>TRF1</td>
<td>Telomere repeat-binding factor 1</td>
</tr>
<tr>
<td>TRF2</td>
<td>Telomere repeat-binding factor 2</td>
</tr>
<tr>
<td>T-SCE</td>
<td>Telomere-sister chromatid exchange</td>
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<td>VEGF</td>
<td>Vascular endothelial growth factor</td>
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I. INTRODUCTION

1.1 Historical Perspective of Nuclear Organization of Tumor Cells

David Paul Hansemann (1858-1920) made the first documentation of tumor initiation as the result of aberrant chromosome numbers in tumor cells. He observed asymmetric mitoses (Hansemann, 1890) and multipolar mitosis (Hansemann, 1891; Hansemann, 1893) in tumor cells. Theodor Boveri (1862-1915) extended Hansemann’s views experimentally by using sea urchin eggs as his experimental model. He published this seminal work, “On Multipolar Mitosis as a Means to Analyze the Cell Nucleus” in 1902 (Boveri, 1902). He concluded that tumor formation was the result of multipolar mitosis which led to chromosome anomalies. He postulated that aberrant nuclear organization was linked to tumor initiation.

1.2 Tumorigenesis

Tumorigenesis is a multistep process that recapitulates Darwinian evolution, in which successive genetic alterations lead to transformation of normal cells into cancer cells (Foulds, 1957; Nowell, 1976). Hanahan and Weinberg defined six hallmarks of cancer: self-sufficiency in growth signals, insensitivity to anti-growth signals, evading apoptosis, limitless replicative potential, sustained angiogenesis, and tissue invasion and metastasis (Hanahan and Weinberg, 2000). Different genetic alterations that contribute to the characteristics of cancer have been identified in various tumor types. Genes
such as **EGFR** (Gibault et al., 2005; Hirsch et al., 2009; Rimawi et al., 2010; Ryott et al., 2009), **HER2/neu** (Choi et al., 2009; Hirsch et al., 2009; Skagias et al., 2009; Slamon et al., 1989), **cyclin D1** (Hadzisejdic et al., 2010; Lee et al., 2010; Liu et al., 2008) and **TGF-α** (Baek et al., 2009) can be over-expressed in tumor cells and thus impart self-sufficiency in growth signals. Tumors are insensitive to anti-growth signals from tumor suppressor genes such as **TP53** (Green and Kroemer, 2009; Riley et al., 2008; Yee and Vousden, 2005), **Rb** (Longworth and Dyson, 2010), **CDKN2A** (Bradley et al., 2006; Haller et al., 2008), and **PTEN** (Guney et al., 2007; Li et al., 2009). Genes involved in the regulation of apoptosis include **BAX**, **BCL2**, **Bcl-L** (Porichi et al., 2009; Zhang et al., 2009), and **Survivin** (Krepela et al., 2009; Lin et al.). Overexpression of genes such as **hTERT** give limitless replicative potential (Califano et al., 1996). Genes involved in angiogenesis include **b-FGF** (Marzioni et al., 2009), **COX-2** (McCormick et al., 2010), and **VEGF** (Zhang et al., 2010). The invasion-metastasis cascade is a series of steps involving primary tumor formation, local invasion, intravasation, extravasation, micrometastases and eventually macroscopic metastases (Fidler, 2003; Gupta and Massague, 2006; Thiery, 2002). Reduced expression of **beta-catenin** (Hsu et al., 2010) and **E-cadherin** (Hsu et al., 2010), or overexpression of **metalloproteinases (MMPs)** are associated with tumor invasion and metastasis (Del Casar et al., 2009; Germani et al., 2009; Szarvas et al., 2010).

Since Hanahan and Weinberg introduced the concept of the six hallmarks of cancer in 2000, additional hallmarks such as inflammation have been
proposed. The concept of a seventh hallmark, an inflammatory microenvironment, was first introduced in the 19th century by Rudolf Virchow (Balkwill and Mantovani, 2001) and, while its popularity waned for more than a century, it has become a generally accepted paradigm (Balkwill et al., 2005; Balkwill and Mantovani, 2001; Coussens and Werb, 2002; Karin et al., 2006; Mantovani et al., 2008). Recent studies have revealed that chronic inflammatory conditions have triggered cancer development. Examples of infections include human papilloma virus (HPV) for head and neck and cervical cancers (Gillison et al., 2000), Epstein Barr virus (EBV) for nasopharyngeal cancer (Gullo et al., 2008) and Hodgkin’s lymphoma (Kapatai and Murray, 2007), *Helicobacter pylori* for gastric cancer (Machado et al., 2010) and mucosa-associated lymphoid tissue (MALT) lymphoma (Zullo et al., 2010), *Schistosomiasis* for bladder cancer (Casella et al., 2009), and hepatitis virus B and C for hepatocellular carcinoma (Neuveut et al., 2010). Autoimmune diseases involved in cancer development include Hashimoto’s disease for thyroid cancer (Lee and Hasteh, 2009) and inflammatory bowel disease for colon cancer.

### 1.3 Genomic Instability

Genomic instability is defined as genetic alterations that affect the normal genetic and chromosomal organizations and functions. Genomic instability is broadly classified into structural and numerical genomic instability, and it ranges from single nucleotide mutations to gross chromosomal aneuploidy (Mai and Mushinski, 2003).
Structural genomic instability comprises point mutations, microsatellite instabilities, deletions, amplifications, duplications, translocations, and inversions of genes and chromosomes. On the other hand, numerical genomic instability, or aneuploidy, corresponds to an abnormal number of a chromosome in a cell, and extends from nullisomy to polysomy. When both structural and numerical genomic instabilities occur concurrently, it is termed karyotypic instability (Mai and Mushinski, 2003). In addition to karyotypic instability, more than one type of structural genomic instability is seen in various tumors throughout the progression of tumors.

Genomic instability is commonly seen in malignant transformation of cells, even though it is also found in normal and premalignant cells. Genomic instability can be induced by drugs, clastogens, deregulation of oncoproteins such as RAS and c-MYC, growth factors and genes responsible for cell cycle machinery, and double-stranded DNA breaks, which can induce gene amplification, deletions, and rearrangement, leading to loss of heterozygosity and translocations (Wright et al., 2009).

1.3.1 Cancer and Genomic Instability

The available evidence suggests that acquisition of the hallmarks of cancer during initiation and progression of cancer is through changes in the genomes of cancer cells, either directly or indirectly. In normal cells, the genome maintenance machineries such as apoptosis, cell cycle control and DNA repair machinery ensure that mutations are rare events that are unlikely to occur within
a human life span. However tumor genomes have acquired increased mutability during tumor progression (Loeb, 1991). Malfunctioning of the genome maintenance machinery can explain this increased mutability (Lengauer et al., 1998). These phenotypes, including microsatellite instability and chromosomal instability, are often seen in the karyotype of cancer cells, where there is an imbalance between the mechanisms of cell-cycle control and mutation rates within aberrant genes. Examples include ataxia telangiectasia, a disease that results from mutations in the ATM gene, a cell cycle checkpoint gene. Nijmegen breakage syndrome is also a disease characterized by chromosomal and genomic instability. Moreover, DNA damage repair genes are lost in different tumors. Their loss of function allows genomic instability and the generation of consequently mutant cells with selective advantages. Mutations of tumor suppressor genes such as Rb (Shay J Biochim Biopys Acta, 1072:1-7), TP53 (Zilfou 2009) and TP73 (Anselmo et al., 2009) can also increase genomic instability

1.4 c-Myc

c-Myc is a proto-oncogene involved in ≥ 70% of human cancer (Beckman and Loeb, 2005; Nesbit et al., 1999). It was first identified as the human homologue of avian retroviral oncogene in 1981. Since its discovery, it has been studied intensively in a wide spectrum of fields, including but not limited to molecular biology of cancer (Kuttler and Mai, 2006; Meyer and Penn, 2008). The overexpression of c-Myc in many human cancers is accomplished by a variety of molecular strategies. These include chromosomal translocations in Burkitt and
AIDS-related lymphoma, gene amplification in breast and prostate cancer, enhanced transcription of structurally normal c-Myc gene in colorectal cancer and increased transcriptional efficiency of c-Myc transcripts in multiple myeloma (Nesbit et al., 1999). Although point mutations in the c-Myc coding region are occasionally seen, particularly in Burkitt lymphoma, the vast majority of tumors express only the wild type protein, thus attesting to its primacy in mediating transformation (Bhatia et al., 1993; Lindstrom and Wiman, 2002; Nesbit et al., 1999).

1.4.1 c-Myc and Genomic Instability

c-Myc is a transcription factor and replication factor that alters the expression of hundreds of downstream target genes, many of them are either oncogenes or tumor suppressor genes (Guffei et al., 2007; Guijon et al., 2007; Kuschak et al., 2002; Prochownik, 2008). There are 2 types of transformation by c-Myc with overlapping pathways. The first type, “acute” transformation, is rapid, efficient, and requires no cooperating oncogenes (Stone et al., 1987) in in vitro transformation that only found in certain immortalized rodent cell line such as Rat1a fibroblasts (Stone et al., 1987). While transformation of primary rodent fibroblasts in vitro is rapid, over-expression of c-Myc alone is not sufficient and requires the contribution of another cooperating oncogene such as mutant H-Ras or K-Ras (Lee et al., 1985).

The second type, “chronic” transformation, is observed in vivo. The “chronic” form of in vivo transformation by c-Myc is a rare event that requires the
acquisition of multiple mutations in other genes affecting cell cycle, senescence, and apoptosis. By greatly accelerating the intrinsic mutation rate at several levels, c-Myc increases the likelihood that these additional mutational “hits” will occur. c-Myc deregulation is insufficient and other non-Ras oncogenes and/or the inactivation of tumor suppressors are required.

In addition to its direct action on chronic transformation, c-Myc also plays an indirect role in inflicting DNA damage through “mutator phenotype” (Beckman and Loeb, 2005), which accelerates the rate of acquisition of these additional genetic hits. Among the types of genomic instability mediated by c-Myc are: 1) single nucleotide substitutions and double-stranded breaks arising via the induction of reactive oxygen species, 2) gene amplification and generation of extrachromosomal elements, 3) numerical chromosomal defects resulting from aberrant DNA synthesis and defects in the mitotic spindle checkpoint, 4) initiation of illegitimate replication of the ribonucleotide reductase R2 gene (Kuschak et al., 2002) and 5) regulation of differentiation and maintenance of neural stem cell as well as tumorigenic potential of glioblastoma when both p53 and PTEN are deleted (Zheng et al., 2008). These non-mutually exclusive activities ensure a constant and varied source of genotoxic insults and suggest that c-Myc overexpression imposes a “mutator phenotype”. This may be an early and necessary requirement for the initial steps in chronic transformation as well as for subsequent evolutionary changes that produce important tumor behaviors such as invasiveness, metastasis, and acquisition of chemotherapy resistance.
1.5 Telomeres

Telomeres are specialized chromatin structures located at the end of the linear chromosome containing repetitive DNA sequences (5’ – TTAGGG- 3’) and a nucleoprotein complex. This complex is termed the telosome and protects the chromosomes from rearrangements and triggering DNA repair pathways. There are 6 proteins associated with the telomeres which are collectively termed the shelterin (de Lange, 2005). It consists of TRF1, TRF2, and POT1 that will directly recognize the TTAGGG repetitive sequence. They are interconnected to the other three proteins TIN2, TPP1 and RAP1.

1.5.1 The End Replication Problem

Semiconservative replication of DNA presents a unique problem: the process only works in the 5’ to 3’ direction, and DNA polymerase requires binding of an RNA primer. Olovnikov and Watson predicted the consequences of this long before the telomere was characterized and termed it the end-replication problem. It anticipated the loss of a small 5’ nucleotide segment as DNA synthesis took place, with progressive replication-induced telomere shortening (Olovnikov, 1973; Watson, 1972).

The 3’ end of the DNA molecule is replicated continuously to the very end, having started at the opposite 5’ terminus. However, synthesis of the lagging strand (also in the 5’ to 3’ direction) is discontinuous because of the requirement for RNA primer binding and unidirectional growth of the new strand. Upon
removal of the RNA primers, the gaps in the discontinuous lagging strand are filled in and ligated. However, there is no provision for such a process at the immediate 5' end, thus leaving a gap. Since the DNA duplex is antiparallel, each daughter molecule will be shortened on its 5' end after replication, with successively shorter daughter chromosomes resulting from additional cycles of cell division.

The end-replication problem, along with other endogenous or exogenous factors, causes shortening of telomeres by ~50-200 base pairs in each cell division (Lansdorp, 2000). Telomere erosion limits the replication capacity of somatic cells that do not express telomerase (Harley et al., 1990; Lindsey et al., 1991). Once the telomeres are shortened to a critical length, cells enter a state of replicative arrest called senescence (Harley et al., 1992; Hayflick, 1965). Senescence is an aging process that protects cells from genomic instability and transformation as a result of telomere dysfunction. Stem cells and germ cells express telomerase and therefore are able to maintain their telomere length and escape senescence. Senescence can be bypassed during tumor development (Campisi, 2000).

Various lines of evidence have shown that several diseases including cancer can be driven by shortening of telomeres in the absence of telomerase. When telomere end protection is compromised, DNA damage response occurs at the chromosome end. This explains replicative senescence. Telomeres can also become dysfunctional through another mechanism, alterations in telomere-
binding proteins, the shelterins. And finally, conditional c-Myc deregulation leading to the formation of telomeric aggregates (Louis et al., 2005) was described previously and is part of my thesis.

1.5.2 Telomerase and Alternative Lengthening Pathway (ALT)

One of the hallmarks of cancer is limitless replicative potential, which could be accomplished through activation of telomerase (Hanahan and Weinberg, 2000). Cancer cells can maintain telomere length through one of two telomere maintenance mechanisms (TMM): active telomerase or alternative lengthening of telomerase (ALT). Kim et al. surveyed several tumor types and have shown that ~85% of tumors express telomerase and maintained a stable and homogenous telomere length and therefore avoid replicative senescence (Kim et al., 1994). The remaining ~15% of tumors either do not maintain telomere length or activate the ALT mechanism (Bryan et al., 1995). Utilization of different TMM is tumor type-specific. Kammori et al. had demonstrated that ~100% of colon adenocarcinomas express telomerase (Kammori et al., 2002). In contrast, a high proportion of sarcomas utilized ALT mechanism, such as 47% in osteosarcomas, 25% in liposarcomas, and 34% in astrocytomas (Costa et al., 2006; Henson et al., 2005; Johnson et al., 2005). There is evidence that both TMMs can be activated in the same cell line and a small number of tumors (Costa et al., 2006; Johnson et al., 2005). Similarly, ALT positive (ALT +) primary tumors can give rise to telomerase positive (telomerase +) secondary tumors and vice versa.
Human telomerase is a ribonucleoprotein complex that is required for overcoming the end-replication problem. It adds the six nucleotide repeats (TTAGGG) to the chromosome ends by utilizing the reverse transcriptase (hTERT) and the RNA template (hTERC), as well as the associated proteins including dyskerin, NOP10, NHP2, and GAR1.

The recombination-based ALT mechanism was first described in *Saccharomyces cerevisiae* that lack one of the essential components of telomerase (TLC1 or EST1) (Le et al., 1999; Lundblad and Blackburn, 1993; McEachern and Blackburn, 1995). A majority of telomerase negative yeast died after entering crisis. Only few clones with RAD52, which is required for double strand break repair and homologous recombination (Lundblad and Blackburn, 1993), were selected.

A telomere tagging experiment performed by Dunham et al. (Dunham et al., 2000) was the first evidence demonstrating that the mechanism underlying ALT in human cells is based on a recombination-like process. The phenotype of human ALT+ cell lines was first described by Murnane et al. and Bryan et al. (Bryan et al., 1995; Murnane et al., 1994). A proportion of ALT+ cells contain large specialized ALT-associated promyelocytic leukemia nuclear bodies (APBs) (Yeager et al., 1999). These APBs contain telomeric DNA, telomere repeat binding factors 1 and 2 (TRF1 and TRF2) and associated proteins, which can be distinguished from other promyelocytic leukemia (PML) bodies in the same cell and in other cell types.
ALT+ cell lines and tumors show heterogeneous telomere length, extrachromosomal circular and linear telomeric DNA (Tokutake et al., 1998), APBs (Yeager et al., 1999), a high frequency of post-replication exchanges in telomeres, also known as telomere-sister chromatid exchange (T-SCE) (Laud et al., 2005), and high instability at a GC-rich minisatellite, MS32 (D1S8) (Jeyapalan et al., 2005). There is a link between the minisatellite instability and the mechanism that underpins ALT. Single molecule analysis of telomeric DNA from ALT+ cell lines and tumors has revealed complex telomere mutations that have not been seen in cell lines or tumors that express telomerase. These complex telomere mutations cannot be explained by T-SCE but must arise by another inter-molecular process. The break-induced replication (BIR) model may explain the observed high frequency of T-SCE and the presence of complex telomere mutations (Dunham et al., 2000).

1.5.3 Telomere and Genomic Instability

Defects in DNA repair and the DNA damage response pathway, as well as structural and functional disruption of telomeres are among the leading causes of genomic instability in cancer and aging (Stewenius et al., 2005). Telomeres are crucial for the preservation of chromosome integrity and controlled cell proliferation (Blackburn, 2001; de Lange, 2002). Telomere-driven instability can promote both structural and numerical chromosomal aberrations (Pampalona et al.). A minimal length of telomeric DNA repeats and proper recruitment of telomere binding proteins are necessary to preserve telomere function (de
In addition, the acquisition of a heterochromatic structure at telomeres is essential for the maintenance of telomere length homeostasis (Blasco, 2007). Over-expression of c-Myc leads to telomeric dysfunction and the subsequent formation of telomeric aggregates and dicentric chromosomes, eventually inducing breakage-bridge-fusion (BBF) cycles (Louis et al., 2005).

1.6 Three-Dimensional Organization of Chromosomes

The mammalian nucleus has a non-random organization of the genome, nuclear compartments, and nuclear proteins (Misteli, 2007; Schneider and Grosschedl, 2007). Studies have shown that the mammalian nucleus is temporally, spatially, and functionally organized and containing numerous nuclear bodies and non-randomly positioned genome domains (Dundr and Misteli, 2001; Kress et al., 2010). The 3D arrangement of the distinct chromosome territories is linked to genomic function and the global regulation of gene expression (Berezney, 2002; Kumaran et al., 2008; Lanctot et al., 2007; Marella et al., 2009; Misteli, 2005; Misteli, 2007; Stein et al., 2003a; Stein et al., 2003b). However, its physiologic function has not been well understood until recently when Solovei et al. demonstrated for the first time that non-random nuclear order plays an important role in physiologic function of retina of nocturnal animal (Solovei et al., 2009).

1.6.1 Nuclear Compartments and Functional Relevance
The genomic function can be spatially organized into three hierarchical levels in a cell: the organization of the chromatin fiber into higher order, the spatial and temporal organization of nuclear processes, and the spatial arrangement of genomes within the nuclear space (Misteli, 2001; Misteli, 2007). The double-stranded DNA helix is folded into higher-ordered structures that eventually form the three-dimensional conformation of chromosomes (Misteli, 2007; Woodcock, 2006). In addition to DNA arrangement, the cellular factors such as replication and transcription machineries are also spatially compartmentalized and distributed in specific nuclear domains (Misteli, 2001).

Most nuclear events occur in the specific, spatially defined compartments that are generally proteinaceous nuclear bodies or chromatin domains (Lamond and Spector, 2003; Misteli, 2005). The most prominent nuclear bodies include: the nucleolus, which is the site of transcription and processing of ribosomal RNA; the splicing factor compartments, which act as storage assembly sites for spliceosomal components; the Cajal body, which is the proposed site of small nuclear ribonucleoprotein (snRNP); and the promyelocytic leukemia (PML) body, which is of unknown function. These nuclear bodies are self-organized into well-defined yet dynamic compartments (Misteli, 2005). Different types of cells and tissues have different preferential positions of nuclear bodies within the nucleus. Similarly, the positions of the nuclear bodies change in different physiological and pathologic processes.
Genomes are non-randomly arranged in the 3D space of the nucleus. Several studies have suggested that the nucleus is compartmentalized and alteration of the 3D nuclear organization affects genomic function (Dechat et al., 2008; Goldman et al., 2002; Gonzalez-Suarez et al., 2009; Gruenbaum et al., 2005; Misteli, 2007). Thomas Cremer described that chromosomes occupy discrete territories in the cell nucleus and contain distinct chromosome-arm and chromosome-band domain (Cremer and Cremer, 2001). Interphase chromosomes are organized into chromosome territories (CTs) in the nucleus (Cremer and Cremer, 2001; Cremer et al., 2006; Spector, 2003). Each individual CT has a non-random location and orientation within the interphase nucleus and no overlapping between CTs. Each CT occupies a spatially defined subvolume in the nuclear space. Using fluorescence in situ hybridization, the CTs are visualized in situ (Lichter et al., 1988). CTs with different gene densities occupy distinct nuclear positions. Euchromatins are less-condensed and gene-rich. On the other hand, heterochromatins are open chromatin regions with gene-poor regions. Gene-rich chromosomes are positioned preferentially towards the center of the nucleus while the gene-poor chromosomes are located toward the periphery. In addition, the arrangement of chromosomes within the nucleus appears to be tissue-specific (Cremer et al., 2003; Meaburn and Misteli, 2007; Parada et al., 2004). Gene-poor, mid-to-late-replicating chromatin regions on chromosomes are enriched in nuclear compartments that are located at the nuclear periphery and at the perinucleolar region. A compartment for gene-dense, early-replicating chromatin is separated from the compartments for mid-
to-late-replicating chromatin. Chromatin domains with a DNA content of ~1 Mb can be detected in nuclei during interphase and in non-cycling cells. The interchromatin compartment (IC) contains various types of non-chromatin domains with factors for transcription, splicing, DNA replication and repair. The CT-IC model predicts that a specific topological relationship between the IC and chromatin domains is essential for gene regulation. The transcriptional status of genes correlates with gene positioning in CTs. A dynamic repositioning of genes with respect to centromeric heterochromatin has a role in gene silencing and activation. Various computer models of CTs and nuclear architecture make different predictions that can be validated by experimental tests. Comprehensive understanding of gene regulation requires much more detailed knowledge of gene expression in the context of nuclear architecture and organization (Cremer and Cremer, 2001).

1.6.1.1 3D Nuclear Organization and its Cellular Functions

It has been proposed that clustering of genes in transcription hot spots plays a role in their regulation and expression (Fraser and Bickmore, 2007; Lanctot et al., 2007), the association of translocation and the relative positioning of chromosomes (Misteli, 2007), replication timing and gene positioning (Gilbert, 2001), and X-inactivation through physical interactions between X chromosomes (Erwin and Lee, 2008).

Gene positioning is found in non-random patterns. Positioning-induced silencing is the phenomenon of position effect variegation (PEV). A gene is
silenced when inserted into heterochromatin genome region. In human cells, the translocation partners for several lymphoma have been found in closer spatial proximity (Roix et al., 2003). These observations suggest that non-random spatial positioning of genome can contribute to what genome regions interact with what other regions (Parada et al., 2002).

1.6.1.2 3D Nuclear Organization and its Physiologic Functions

Solovei et al. has shown that the physiologic functions of rod cells are closely related to the 3D organization of the interphase nucleus (Solovei et al., 2009). The nuclear architecture of rod photoreceptor cells is different in nocturnal and diurnal mammals. In diurnal retinas, the rods possess the canonical architecture with euchromatin located in the nuclear center while the heterochromatin is situated at the nuclear periphery. On the other hand, the rods of nocturnal retinas have an inverted pattern, where heterochromatin resides toward the nuclear interior while euchromatin, nascent transcripts, and splicing machinery line the nuclear border. The differences of the two patterns suggest that the canonical architecture in eukaryotic nuclei result in more flexible chromosome arrangements, facilitating positional regulation of nuclear functions.

1.6.2 Telomere Compartmentalization

Telomere compartmentalization for telomere function has been studied in yeast (Akhtar and Gasser, 2007). However, the underlying mechanism for the spatial organization of telomeres in the mammalian nucleus still remains elusive.
In addition, the relationship between the arrangement of mammalian telomeres and telomere biology is not well understood. Gonzalez-Suarez et al. showed that loss of A-type lamins changes the nuclear distribution of telomeres away from the nuclear centre and towards the periphery. The latter results in telomere shortening, defects in telomeric heterochromatin, and increased genomic instability (Gonzalez-Suarez et al., 2009).

### 1.7 Molecular Imaging

Light microscopy has been used for centuries to visualize structures too small to see with the naked eyes. Optical microscopy is an important tool in the investigation of cellular process and structures. It allows one to work with intact samples including living cells and to see samples with the naked eye.

#### 1.7.1 Light Microscopy

Robert Hooke (1638-1703) published the Micrographia in 1665, which is the first book on microscopy (Hooke, 1665). He refined compound microscopy, which consisted of a stage, a light source and three optical lenses. Among numerous detailed observations made with his microscope, he termed the pores of the cork as “cells”. The introduction of multiple lenses increased issues with spherical and chromatic aberration. A simple microscope containing a single convex lens was first developed by Van Leeuwenhoek (1632-1723) (van Leeuwenhoek, 1800). He was the first to observe sperm cells and bacteria and
protozoa in water. Using a simple microscope was superior for observation until the advent of achromatic lens became widely available in nineteenth century.

1.7.2 Resolution of Light Microscopy

Diffraction occurs as the light enters the lens aperture. A diffraction pattern in the image which has central white spot with surrounding white rings separated by dark rings is known as the Airy disc, as described by George Airy in 1835 (Airy, 1835). The diffraction barrier dictates the limit of resolution of light microscopy.

Resolution is the ability to discern 2 points distinctively, and it is described by Abbe in 1873 (Abbe, 1873). The Abbe's formula published in 1873 stated:

\[ D = \frac{\lambda}{2} n \sin a \]

\( D = \) minimum distance between 2 objects that reveals them as separate entities
\( \lambda = \) wavelength of the light
\( n \sin a = \) numerical aperture of the lens.

The limit of optical resolution of a conventional light microscope is approximately ½ of wave-length. This is approximately 200 nm.

1.7.3 Fluorescent Dyes and Fluorescent Microscope

Fluorescence was first defined by Stokes who described it as light emission induced during excitation (Stokes, 1852). Fluorescence provides
infinite contrast with the right equipment. The fluorescent microscope was first constructed by Oskar Heimstädt in 1911 (Heimstädt, 1911). Initial limitations of fluorescence microscopy included reliance on auto-fluorescence of the images object and the need for transmitted illumination and darkfield condenser. The first hurdle was addressed via development of fluorochrome or secondary fluorescence (Freund, 1969). This involves attaching exogenous fluorescence chemicals to the samples. Epi-fluorescence microscope was developed in 1929 (Ellinger and Hirt, 1929). The light source lies on the same side of the sample as the objective, and excitation and emission light pass through the objective. This allows more efficient sample excitation and imaging of opaque objects. Another advance occurred in 1967 with development of dichromatic mirrors or beam splitters (Ploem, 1967). Dichromatic filters reflect a narrow width of wavelengths while transmitting all others, allowing illumination of the sample with precise wavelength. The reflected shorter wavelength excitation light enters the objective which also functions as a condenser to allow even excitation of the sample. The emitted longer wavelength fluorescent light is collected by the objective and passed through the dichromatic mirror and a barrier filter to the eyepiece.

1.7.4 The Charge-Coupled Device (CCD)

The charge-coupled device (CCD) was invented and published in 1970 by Boyle and Smith (Boyle and Smith, 1970). The CCDs are solid-state image sensors that make use of the photoelectric effect. This allows transformation of
light or photons into electric signals so that light can be captured and stored electronically. This technology is used in many fields including digital photography for research or recreation, and endoscopy for medicine or industrial purposes. Use of CCD in digital imaging of fluorescence was first described in 1986 (Connor, 1986). The study was to image free calcium levels in individual central nervous system cells from rat embryo. From digital images captured, the author was able to show high levels of calcium at the growing tips of the central nervous system cells.

1.7.5 Deconvolution Algorithm

The conventional fluorescence microscope contains light from throughout the specimen. The out-of-focus fluorescence confounds what is present in the focal plane. A means to achieve clarity and to minimize artifact of images is by applying a mathematical processing method known as deconvolution algorithm to the stack of images. The deconvolution algorithm determines how much out-of-focus light is expected given the optics in use and then seeks to reassign the diffracted light to its points of origin in the specimen. With the use of deconvolution, reconstruction of a three-dimensional object is possible by taking a sequential stack of images captured on films or digital image sensors. The deconvolution algorithm was first used in 1983 and allowed for the recovery of detail of the polytene nucleus of the *Drosophila melanogaster* salivary gland from fluorescent images captured on films (Agard and Sedat, 1983).
Different deconvolution algorithms have been developed. The constrained iterative deconvolution algorithm was chosen for the 3D work performed in this thesis (Schaefer et al., 2001) to restore 3D images.

1.7.6 3D Microscopy

With the convergence of different disciplines of science including the development of the CCD digital camera, deconvolution algorithm, and automated FISH microscope, the ability to visualize microscopic objects in 3D space is greatly enhanced. Through collaboration with physicists from the Netherlands, the Teloview program has been developed to perform quantitative analysis of three-dimensional images of telomeres and chromosomes (Vermolen et al., 2005).

The importance of telomeres and the development of CCD technology is highlighted by the Nobel prize committee members who awarded the pioneers in the fields of telomeres and CCD the Nobel prizes in 2009. Elizabeth Blackburn, Jack Szostak, and Carol Greider were awarded the Nobel Prize in Physiology or Medicine for their discovery of the structure and function of telomeres. Willard S. Boyle and George E. Smith were awarded the Nobel Prize in Physics for the invention of the CCD sensor.
II. Published Manuscripts

2.1 The three-dimensional organization of telomeres in the nucleus of mammalian cells.

Access the publication online is available via the following link:

http://www.biomedcentral.com/1741-7007/2/12

Rationale of the Investigation

Genomic instability is a characteristic of the tumor cell. It includes numerical and structural alterations of chromosomes. Numerical changes such as amplifications, deletions, and duplications have been widely studied. However, aberrations of temporal and spatial organization of chromosomes in three-dimensional (3D) space and dysregulation of transcription, replication, and stability of the genome in tumor cells have yet to be investigated further.

Objectives

To establish the telomere organization in 3D space of interphase nuclei.

To evaluate for differences in telomeric organization in normal and tumor cells.

Hypothesis

Telomeres are arranged in a non-random and dynamic fashion in the 3D space of a normal cell. In tumor cells, telomere organization is disrupted as a result of genomic instability.
Summary of Findings

Nonrandom dynamic 3D telomere organization in a normal nucleus

The 3D telomere organizations of normal, immortalized and tumor cells have been examined in different cell lines and various human primary tissues. Telomeres are organized in non-random and non-overlapping arrangement in the 3D space of a normal nucleus.

3D telomere organization is disrupted in the nucleus of a tumor cell

Telomeres are distributed non-randomly in the 3D space of a mammalian nucleus throughout the cell cycle. However, this organization is disrupted in tumor cells with the formation of telomeric aggregates of various numbers and sizes.

My Contributions to the Work

I was one of the first ones to make the initial observation of non-random telomere organization in normal and immortalized cell in 3D space. I had also made additional observation that telomeric aggregations were a common denominator in various tumor cell lines and tissues.

To study the 3D telomere organization throughout the cell cycle of a normal mammalian nucleus, I planned the experiments and performed 3D telomere fluorescence in situ hybridization (FISH) followed by the quantitative analysis of telomere distribution using the TeloView programme developed by
our collaborators in the Netherland. I discovered that telomeres were arranged in a cell-cycle dependent fashion, with the formation of the telomeric disk in the late G2 phase.

I obtained all head and neck cancer tissues from patients. Prior to the surgery, I described the study to the patients, answering their questions, and obtained the consents prior to the surgery. I also processed the tissues for the study by preparing the fresh frozen sections for FISH analysis. I also prepared clinical data and correlated the experimental result with the clinicopathological parameters.
Research article

The three-dimensional organization of telomeres in the nucleus of mammalian cells

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Abstract

Background: The observation of multiple genetic markers in situ by optical microscopy and their relevance to the study of three-dimensional (3D) chromosomal organization in the nucleus have been greatly developed in the last decade. These methods are important in cancer research because cancer is characterized by multiple alterations that affect the modulation of gene expression and the stability of the genome. It is, therefore, essential to analyze the 3D genome organization of the interphase nucleus in both normal and cancer cells.

Results: We describe a novel approach to study the distribution of all telomeres inside the nucleus of mammalian cells throughout the cell cycle. It is based on 3D telomere fluorescence in situ hybridization followed by quantitative analysis that determines the telomeres' distribution in the nucleus throughout the cell cycle. This method enables us to determine, for the first time, that telomere organization is cell-cycle dependent, with assembly of telomeres into a telomeric disk in the G2 phase. In tumor cells, the 3D telomere organization is distorted and aggregates are formed.

Conclusions: The results emphasize a non-random and dynamic 3D nuclear telomeric organization and its importance to genomic stability. Based on our findings, it appears possible to examine telomeric aggregates suggestive of genomic instability in individual interphase nuclei and tissues without the need to examine metaphases. Such new avenues of monitoring genomic instability could potentially impact on cancer biology, genetics, diagnostic innovations and surveillance of treatment response in medicine.
Background
Cancer is characterized by multiple alterations that affect the modulation of gene expression and the stability of the genome. These interconnected changes occur within the nuclei of cells that affect their three-dimensional (3D) organization during tumor initiation and progression [1,2]. It seems reasonable to assume that the highly organized mammalian interphase nucleus is the structure that ascertains genomic stability. In line with these concepts, oncogenic activation remodels this nuclear order and sets the stage for genomic instability as we have recently measured for conditional c-Myc deregulation. The deregulated expression of c-Myc alters the 3D nuclear space of chromosomes and telomeres, and makes genomic rearrangements topologically feasible (Chuang et al., in preparation).

Defining the structural organization of the interphase nucleus is therefore essential to our understanding of the 3D genome organization in the interphase nucleus. Such a study can be performed by fluorescence in situ hybridization (FISH). Two of the most attractive features of FISH measurements of the 3D nuclear organization are the ability to simultaneously visualize multiple targets and the structural organization of nucleus and cells, something that cannot be achieved by array-based methods.

The organization of the interphase nucleus has been studied since the late nineteenth century [3]. It is now well accepted that the position of chromosomes in the nucleus plays an important role in gene regulation [4]. Nevertheless, some controversy exists. Most laboratories have observed a non-random organization of chromosome territories [2,5,6] that has been conserved during evolution [7]. This has been further supported by studies that demonstrate an architectural stability of the chromosomal positions in the nucleus [8,9]. There are, however, different observations on chromosomal positions [10-15] as well as on positional changes of chromosomes during the cell cycle [16,17].

Recently, interest has also focused on telomeres, whose importance to genomic stability was recognized as early as the 1980s [16]. Capping the chromosomes, telomeres are responsible for chromosomal integrity [19] to prevent genomic instability [20]. Some reports have been published on the 3D organization of telomeres in the nucleus, mainly with regard to the distances of telomeres from the nuclear shell. Telomeres have been previously found at the nuclear edge [21], at the nuclear periphery [22], throughout the entire nucleus [18,23], in non-radial association [11], in association with the nucleolus [24] or in the nuclear matrix [25].

Telomere dynamics also have been studied in living human U2OS osteosarcoma cells [26]. Individual telomeres showed significant directional movements and telomeres were shown to associate with promyelocytic leukemia bodies in a dynamic manner. This means that telomere structure is dynamic, and may be important for both transcriptional processes and for stabilizing chromosome positions in the nucleus.

We have developed a method of studying the organization of the genome by analysis of the 3D organization of telomeres in the nucleus and their positional changes along the cell cycle, using flow-sorted living cells. This method enables us to determine, for the first time, that telomere organization is cell-cycle dependent, with assembly of telomeres into a telomeric disk in the G2 phase. Moreover, we show for tumor cells that the 3D telomere organization is disordered and that telomeric aggregates are formed. These results emphasize a non-random and dynamic 3D nuclear telomeric organization and its importance to genomic stability.

Results and discussion
To study the organization and structure of the genome in the nucleus, we took the approach of labelling only the telomeres and measuring their 3D organization as indicators for chromosomal distribution. After the 3D fluorescent measurements, the data were analyzed with a programme that was developed for this study. The programme finds all the telomeres in the nucleus their size, intensity and shape, and determines the telomeric organization inside the volume of the nucleus. One crucial property that we analyzed was the distribution of the telomeres inside the nuclear volume. We first segmented the nucleus and found the centre of each telomere. We then found the smallest convex set of polygons that contains all the telomeres (Fig. 1). This was done by using the Quickhull algorithm [27]. In most cases, we found that the volume contained by the telomeres resembles either a sphere or a flattened sphere (disk). It can be described as an ellipsoid with two similar radii (a=b) and a different third one (c, Fig. 2). Such a shape is called a spheroid. The level of flattened of the volume occupied by the telomeres can, therefore, be described by the ratio of the two radii that are different, a / (a + c) or e = a/c. The larger the ratio, the more oblate (or disk-like) is the shape of the volume occupied by the telomeres, while e < 1 means that the volume is spheroidal.

The optical resolution and signal-to-noise ratio are presented in Fig. 3. The images of two neighboring telomeres that are 1200 nm and 400 nm apart, and the corresponding intensity along the line connecting the pair, indicates the smallest telomere distance that can still...
Figure 1
The distribution of the telomeres in the nucleus volume is found by fitting a convex set of polygons that contains all the telomeres. This volume usually looks like either a sphere or a disk and can be described as an ellipsoid.

Figure 2
In general, the ellipsoid's main axes along the x/y/z direction do not coincide with the microscope-slide plane and optical axes xyz. Our program finds an ellipsoid that contains all the telomeres and the size of its main axes a,b,c. In most of the cases the x/y axes of the ellipsoid are similar, i.e., x≈b. Therefore, the ratio a/c is a good measure of the flatness level of the ellipsoid and of the telomere organization inside the nucleus.

Figure 3
Demonstration of the signal-to-noise and spatial resolution of our measurements. The fluorescence intensity is bright (typical signal-to-noise ratio of 10:1). Two pairs of telomeres are shown, 1200 nm apart (top), which can be easily separated, and 400 nm apart (bottom). The insets show the actual images.

be unambiguously distinguished (approximately 200 nm).

It is expected that 83 telomeres will be observed in the interphase nucleus for normal mouse cells (92 for a normal somatic human cell), however, in our measurements we were usually able to identify approximately 40 separated telomere regions in each mouse cell (50 in human cells). Similar results have been described before [23,28]. This is probably due to neighboring telomeres that are closer than the optical resolution (see Fig. 3), but it does not affect the analysis of the telomere distribution in the nucleus as long as the hybridization efficiency is high. This was verified by two-dimensional measurements of all the telomeres in a metaphase spread (using the same probe), where at least 90% of the telomeres are unambiguously observed (Fig. 4).

We first described the major observation of primary BALB/c mouse B lymphocytes that were studied along the cell cycle. These studies were followed by the analysis of immortalized cells. The lymphocytes were sorted according to their DNA content for the determination of the G0/G1, S or G2/M phases (see Methods).
By analyzing cell-cycle sorted primary mouse lymphocytes we found that the 3D telomere organization changes during the cell cycle. Telomeres are widely distributed throughout the nucleus in the G0/G1 and S phases with a calculated a/c ratio of 0.9 ± 0.4, which means a spherical-like volume of distribution. However, during G2, telomeres are not observed throughout the whole nucleus. Their 3D organization changes, with all the telomeres assuming a central structure that we call the telomeric disk, which has never been reported before. In this ordered structure, all the telomeres align in the centre of the nucleus as cells progress into the late G2 phase. The a/c ratio they assume is 6.0 ± 2.0, which means a very flat disk (almost a coin shape).

Typical lymphocytes from different phases are shown in Fig. 5. The a/c ratio of these cells in the G0/G1, S and G2/M phases is 0.8, 0.8 and 6, respectively, and clearly shows the correlation of the a/c ratio with the telomere distribution and the organization of the telomeric disk that we found in the G2 phase. The elongation of the telomeres along the Z axis (the optical axis) relative to the XY plane has the same ratio as the point spread function of our system and results from the power optical resolution along the optical axis. However, this has a very small effect on the shape of the whole nucleus.

Similar results have been observed in primary human lymphocytes, primary human fibroblasts and in normal human epithelial tissue (see additional file for more data). This suggests that chromosomes assume a very precise order that par-aligns them prior to the onset of mitosis. In order to ascertain that the telomeric disk was not the result of a distorted nucleus, our analysis programme
Figure 6
BrDU-positive cells were live sorted and synchronized in the S phase. They were harvested from a culture at time intervals of 3.5–9 hours. The cells were then fixed for 3D analysis. For each time point we have measured: 1. the fraction of nuclei with a telomeric disk; 2. the fraction of cells in mitosis; and 3. the fraction of cells with interphase nuclei but without a telomeric disk. Ninety percent of the cells formed a telomeric disk 3.5 hours after BrDU incorporation and were therefore interpreted as cells in the late G2 phase (black line and circles). Cells entering mitosis (dashed line and squares) peaked at 7.5 hours (65%) and cells in G1 (dotted line and triangles) peaked after 8.5 hours (57%). The increase in the number of metaphases at 8.5 hours cannot be explained and probably lies within the limits of experimental errors.

These results reveal that the telomeric disks are formed in the late G2 phase. As cells progress from G2 to M, chromosomes organize into metaphases and, therefore, the number of cells in interphase with a telomeric disk decreases. Because there is no other state of transition between telomeric disk and mitosis, we conclude that the telomeric disk is the 3D telomeric organization assumed in late G2. Thus, it is also the final stage of the interphase nucleus that permits the organization of the genetic material prior to its entry into the M phase and prior to chromosome segregation. Cells in late G2 with a telomeric disk have additional characteristic features: 1) they exhibit a larger overall nuclear volume than their G1 or S phase counterparts (this increase in size was also confirmed by fluorescent activated cell sorter [FACS] analysis); and 2) they begin to show signs of early reorganization of the chromatin into partially condensed areas (as visualized using the DAPI stained image).

At the end of the M phase, we observe cells that enter into the G1 confirmation of telomeres, with a wide spatial distribution of telomeres throughout a smaller nucleus. In conclusion, this data indicates that the telomeric disk is a novel structure within the interphase nucleus in late G2 that has not been previously described. Its existence points to the fundamental importance of ordered nuclear organization at the end of G2. The telomeric disk probably assures the proper organization of chromosomes prior to mitosis and their organized segregation during mitosis. Together with information that has been previously published on telomeric dynamics [36,28], it is tempting to speculate that telomeres take an active part in the process of chromosome organization into a unique structure, the telomeric disk, during G2. This alignment of telomeres and chromosomes would facilitate the proper subsequent organization of the chromosomes into an equatorial plane during cell division. This process may be driven by the telomeres themselves (that are free of the nuclear matrix) or through the nuclear matrix. The telomeric disk may also allow for a late G2 checkpoint.

compared the telomere distribution volume and shape with that of the 4',6-Diamidino-2-phenylindole (DAPI) – stained nuclei and verified that the nucleus itself still had a spherical-like volume. We rarely found distorted nuclei and excluded these cells from the analysis. The nucleus shown in G2 is not fully spherical. Such a shape is expected, because when the telomeres form a disk, it pulls the chromosomes and forces them to be closer to the disk, which results in an oblate shape as well.

To further study the phase transition timing along the cell cycle we used the synchronous bromodeoxyuridine (BrdU) sorting method. The cell population was pulse-labelled with BrdU in the S phase and flow sorted. Cells were placed back into culture and sub-populations harvested at 3.5, 4, 5, 6, 7, 8, 8.5, 9 and 10 hours after labelling and sorting. The cells were then fixed for 3D analysis. A minimum of 20 cells from each of these sub-populations were measured, analyzed and divided into the following three categories: 1) nuclei with a telomeric disk; 2) cells in mitosis; 3) cells in interphase without telomeric disk and mitotic figures (evaluated as G1 cells). The cell fractions as a function of time are shown in Fig. 6. Most cells (90%) form a telomeric disk 3.5 hours after BrdU incorporation. These cells are, therefore, interpreted as cells in the G2 phase. The fraction of metaphase cells peaks at 7.5 hours (65%) and the cell fraction of interphase cells that does not have a telomeric disk (and is interpreted as being in the G1 phase) peaks at 8.5 hours (57%).
Further work on the subject can also be performed in vivo, as has been shown by Aisenberg et al. [25]. In such a way the full dynamic process can be observed, which is complementary to the single time-points that are shown in our work.

We have continued to observe the distribution of telomeres in cancer cells. Typical 3D images constructed from normal nuclei and from a Burkitt lymphoma cell line (RAJI), as well as from primary mouse plasmacytoma (PCT) and primary human head and neck squamous cell carcinoma (HNSCC) stage IV (Fig. 7), show that telomeres form aggregates and thus a partially altered telomeric disk. Such telomeric aggregates are characterized by both a larger volume and larger integrated intensity than their normal non-overlapping and non-aggregated counterparts. They are not observed in normal cells. Similar results for altered telomeric organization have also been found in human neuroblastoma and colon carcinoma tumor cell lines.

In line with these concepts, oncogenic activation remodels this nuclear order and sets the stage for genomic instability as we have recently measured for conditional c-Myc deregulation. We have found that deregulated expression of c-Myc alters the 3D nuclear organization of chromosomes and telomeres, and makes genomic rearrangements topologically feasible (Chuang et al., in preparation).

Conclusions

In summary, we have shown that 3D optical imaging followed by the analysis of telomeres in the interphase is an important tool for basic research and cancer biology. We have found cell-cycle dependence of the telomere organization in the nucleus, where telomeres align into a telomeric disk during the late G2 phase. Such an organization has never before been reported.

Telomeric aggregates are found in tumor cells and, therefore, an alteration of the telomeric disk is seen. Transient telomeric aggregations potentially cause irreversible chromosomal rearrangements.

The above findings indicate that it is now possible to examine the presence of telomeric aggregates suggestive of genomic instability in individual interphase nuclei and tissue, without the need to examine metaphases. Such new directions of monitoring genomic instability could potentially have an impact on cancer biology, genetics, diagnostic innovations and surveillance of treatment response in medicine.

**Figure 7**

Normal: A normal blood cell; RAJI: A Burkitt lymphoma cell line; PCT: A primary mouse plasmacytoma cell; HNSCC: A primary human head and neck squamous cell carcinoma (stage IV). The distribution of telomeres in cancer cells compared with a normal cell. Images are shown in Fig. 5. Aggregates of telomeres are formed and the telomere disk that appears in the G2 phase is distorted.

**Methods**

**Cells**

Mouse primary cells were directly isolated from BALB/c mice and stimulated with lipopolysaccharide to enter into the cell cycle [29]. Primary mouse fetal liver cells were also directly isolated from BALB/c mice. Mice were studied according to the protocols approved by Canadian Council Animal Care. Immortalized mouse pro B lymphocytes have been described elsewhere [39]. Human primary cells were obtained from healthy donors. Head and neck squamous cell carcinoma and control tissue were obtained from a patient at CancerCare Manitoba upon ethics approval and informed consent.

**Fixation techniques**

Pro B lymphocytes [30] were fixed in four ways: i) following cytospin preparations, cells were fixed in 3.7% formaldehyde (1xPBS/50 mM MgCl2); ii) cells were allowed to
grow on glass slides and were fixed in 1% formaldehyde (3D fixation); iii) cells were fixed in suspension with 2.7% formaldehyde (3D fixation); and iv) cells were fixed in methanol-acetic acid (3:1) according to standard protocols [29]. Tissue was fixed following cryosection (5 μm sections were used) in 1% formaldehyde (1xPBS/50 mM MgCl₂). All hybridizations shown in this report were carried out after 3D fixation.

**Fluorescent activated cell sorter (FACS) analysis**

For FACS analysis, primary lymphocytes were fixed in 70% cold ethanol and stained with propidium iodide (1 μg/mL) following RNase (20 μg/mL) digestion. The stained cells were analysed for DNA content by flow cytometry in a EPICS Altra cytometer (Beckman Coulter). Cell cycle fractions were quantified with WinCycle software (Phoenix Flow Systems, San Diego, CA).

**Cell sorting**

Cells were stained with Hoechst 33342 (Molecular Probes) at a final concentration of 1 μg/mL for 90 minutes at 37°C and 5% of carbon dioxide (CO₂). Cells were sorted according to their DNA content (G₀G₁, S, and G₂M phases) with a EPICS Altra cytometer (Beckman Coulter) equipped with a UV laser (Coherent, excitation at 350 nm) and a 480 nm band-pass filter.

**BrDU labelling**

Pro B lymphocytes were labelled in vivo with 10 μM of BrDU (5-Bromo-2-deoxyuridine, SIGMA-ALDRICH, Lyyen, France) for one hour at 37°C in humidified atmosphere (5% CO₂). BrDU was then detected with 5 μL/1x10⁶ cells of anti-BrDU-PITC (fluorescein isothiocyanate) antibody (TEBU, Le Perray-en-Yvelines, France) at identical conditions for 30 minutes. Thereafter, all BrDU (i.e. FITC)-positive cells were live sorted, placed into culture for different times and harvested at 3.5, 4, 5, 6, 7, 8, 8.5, 9 and 10 hours after labeling and sorting. The cells were then fixed for 3D analysis. For each time point we have measured: 1. the fraction of nuclei with a telomeric disk; 2. the fraction of cells in mitosis; and 3. the fraction of cells in interphase nuclei without telomeric disk and mitotic figures that were evaluated as G₀ and S phase cells.

**Telomere FISH using Cy5-labeled PNA probes**

Telomere FISH was performed as described [31] using a Cy5-labelled PNA probe (Dako, Glostrup, Denmark). Telomere hybridizations were specific as shown by metaphase hybridizations and the correct number of the telomeric signals observed at the ends of chromosomes prepared from primary cells (Fig 4).

**3D image acquisition**

Unless stated otherwise, 20-30 cells were analyzed by 3D imaging from each cell type and phase type. Part of the measurements were done with a confocal microscope (Leica AOR-BSP) and most of them with a conventional Axiosplan 2 (Zeiss) with a cooled AxioCam HR CCD followed by deconvolution [30]. DAPI, FITC and Cy3 filters (Zeiss) were used in combination with Planapo 63x/1.4 oil (Zeiss). Axiovision 3.1 software with a deconvolution module and rendering module were used (Zeiss). Both methods gave similar results.

80-100 sections were acquired for each 3D nucleus, typically with 200 × 200 pixels per section with a ~100 × 100 nm nominal imaging area per pixel (steps of 200 nm along Z). The point-spread function of our system has a full width at half max of approximately 200 nm in the plane and 400 nm along the optical axis.

**3D analysis of telomeres**

In order to analyze the telomere distribution in the nucleus, we developed a special 3D image analysis programme. The main algorithmic part is described below. The programme (TelView) is based on the Matlab computer language (The MathWorks, Natick, MA, USA) and some of the image processing algorithms are based on the Dipimage library (developed at the Quantitative Imaging Group, Delft Institute of Technology, Delft, The Netherlands) [32].

The programme segments the nucleus volume by a derivative-based algorithm using a morphological top and bottom-hat algorithm [33]. The volume, intensity and centre of gravity are calculated for each spot. The programme then finds a principle plane in the nucleus (x'y') that is the closest to all the telomeres (Fig 2). This is especially important when a tissue section is analyzed, because this plane should not necessarily be parallel to the microscope slide plane.

The telomeric distribution inside the nucleus is described by fitting an ellipsoid to the volume occupied by the telomeres (three different main axes, Fig 2). The distributions were found to be either oblate or spherical (i.e. the two principle axes along the main x'y' plane of the sphere are similar). It is, therefore, convenient to describe the distribution volume as a sphere (i.e. an ellipsoid having two axes of equal length). As such, it is simpler to describe the sphere's degree of variation from a perfect sphere by the ratio a/b, where a and b are the similar semi-axes and c is the third one. Such a description reflects the degree to which the telomere's volume is oblate.

**Authors' contributions**

TCYC performed the data analysis, wrote the discussion, performed the hybridizations, took all head and neck cancer samples from tumor collections during surgery to preparation of frozen sections, hybridization, analyses
and patient records. Smo performed some of the hybridizations and analyses while visiting SM’s lab. VG developed the 3D analysis methods, 3D algorithms and the program that analyzes the nuclei, the program was written in its current version and acts as an accompanying author. AVCC organized the data, tables, and G2 phase data. ITY, BV and RD took part in the development of the 3D analysis methods and algorithms, and the program that analyzes the nuclei. VM did the BrdU-labeling experiments and G2 study in France and in SM’s lab. MP performed the G2 analysis in SM’s lab. WB performed the metaphase telomere FISH. PK directed the programme of head and neck surgery and provided some of the samples that were used for the study. TF supervised VM and MP in France. PB supervised Smo in Heidelberg. SM planned and carried out the project that was performed in SM’s lab, and was supervisor for TCYG Smo (while visiting SM’s lab). AVCC, MB, MP and VM, performed part of the experiments.

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References

2.2 c-Myc induces chromosomal rearrangements through telomere and chromosome remodeling in the interphase nucleus.

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Rationale of investigation

c-Myc is a proto-oncogene found in more than 70% of human cancers. Telomere aggregation and dysfunction are frequently observed in tumor cells as described in my paper “The three-dimensional organization of telomeres in the nucleus of mammalian cells” (Chuang et al 2004). One of the mechanisms of telomere dysfunction is mediated by c-Myc deregulation.

Objectives

To investigate the effect of conditional over-expression of c-Myc on telomeres, quantitative 3-D analysis of telomeres is carried out via deconvolution microscopy and the Teloview software. Upon deregulation of c-Myc, spectral karyotyping was carried out to assess chromosomal aberrations.

Hypothesis

Deregulation of c-Myc first leads to telomeric dysfunction including aggregation and fusion, which will lead to breakage-bridge-fusion process that causes chromosomal aberration.

Summary of findings

Conditional c-Myc induction disrupts 3D telomere organization by forming cycles of telomeric aggregates in interphase nuclei. These cycles lead to the onset of genomic instability as demonstrated by the breakage-bridge-fusion
(BBF) cycles that results in nonreciprocal translocations and chromosomal rearrangement. In addition, telomere fusions were observed on metaphases.

**My contribution**

I conceived the project idea in collaboration with Dr. Sabine Mai and Dr. Alice Chuang to investigate *c-Myc* deregulation as a mechanism of telomere dysfunction. I performed 3-D telomere FISH, 3-D image acquisition and analysis of images. The work contributed to figures 1, 2, and 3. I was involved in manuscript preparation, writing, revision and submission to various publishers.
c-Myc induces chromosomal rearrangements through telomere and chromosome remodeling in the interphase nucleus


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In previous work, we showed that telonomers of normal cells are organized within the 3D space of the interphase nucleus in a nonoverlapping and cell cycle-dependent manner. This order is distorted in tumor cell nuclei where telonomers are found in close association forming aggregates of various numbers and sizes. Here we show that c-Myc overexpression induces telonomer aggregations in the interphase nucleus. Directly proportional to the duration of c-Myc deregulation, we observe up to five cycles of telonomer aggregate formation in interphase nuclei. These cycles reflect the onset and propagation of breakage-bridge-fusion cycles that are initiated by end-to-end telomeric fusions of chromosomes. Subsequent to initial chromosomal breakages, new fusions follow and the breakage-bridge-fusion cycles continue. During this time, noncopyclic translocations are generated. c-Myc-dependent remodeling of the organization of telonomers thus precedes the onset of genomic instability and subsequently leads to chromosomal rearrangements. Our findings reveal that c-Myc possesses the ability to structurally modify chromosomes through telomeric fusions, thereby reorganizing the genetic information.

Materials and Methods

Cells and Conditional Myc Activation. Culture conditions have been described for Ba/F3 (25) and ProB (26) cells. The plasmacytoma cell line MOPC410D was a gift of J. M. Futran (National Institutes of Health, Bethesda). Cell viability was determined by hemocytometer counting or trypan blue. The primary mouse plasmacytoma DCPC21 was isolated from a BALB/c mouse (17). v-abl/myc-induced plasmacytomas (30) and primary lymphocytes were collected from BALB/c mice (Central Animal Care unit 02-089).

To activate MyeR (59) in Ba/F3 or ProB cells, 106 cells per ml were treated with 100 nM 4-hydroxynonenal (4HT). Cells were split 24 h before 4HT treatment. Non-4HT treated control cells were cultivated in ethanol, which is used to dissolve 4HT (25, 26, 39). Two different MyeR activation schemes were performed. First, analyses of c-Myc-induced changes in 3D telomere organization were carried out after a single addition of 4HT that was left in the culture medium until its biological effects subsided (10–42). Nuclei were examined every 24 h over a 10-day period. A second time course was performed every 9 h to 12 h (Fig. 1). To enable a time-dependent analysis of Myc activation, 4HT was given for 2 or 12 h and was removed. Alternatively, 4HT was added every 12 h or was given once but left in the culture. MyeR activation was determined by fluorescent immunohistochemistry.

Immunohistochemistry (IFC). Fluorescent IFC of Myc protein was performed as described in ref. 48 by using a polyclonal anti-cMyc antibody (262, Santa Cruz Biotechnology) and a goat anti-rabbit IgG FITC antibody, each at a dilution of 1:100. Analysis was

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performed by using a Zeiss Axiophot 2 microscope. Images were acquired with a Cooke CCD Seniscam Camera.

Cell Death. Apoptotic bodies for control and MycER-activated cells were assessed by two independent observers who scored 300 DAPI-stained nuclei per point in the presence or absence of MycER activation.

Telomere FISH. Ba/F3, P3, and plasmacytoma cells were collected (200 x 10^6 for 10 min) and resuspended in PBS containing 3.7% formaldehyde (Fuka) and incubated for 20 min. The labeled telomere FISH protocol was performed (9, 44) by using Cy3- or FITC-labeled PNA probes (DAKO). Three independent experiments were performed. At least 30 nuclei and 30 metaphases were examined per time point. Imaging of metaphases after telomere FISH was performed by using Zeiss Axioplan 2 with a cooled AxioCam HR B&W, DAPI, Cy3, or FITC filters in combination with Planoapo 63x/1.4 oil objective lens. Images were acquired by using Axiovision 3.1 (Zeiss) in multichannel mode. Because of the presence of multiple variables, the general linear model procedure was used. To assess average aggregation among different groups, a two-way ANOVA test was performed for normality and robustness of the data. For details of all tests performed, see Supporting Materials and Methods, which is published as supporting information on the PNAS web site.

3D Image Acquisition. At least 30 nuclei were analyzed for each time point. Axiovision 3.1 with deconvolution module and rendering module were used. For every fluorochrome, the 3D image consists of a stack of 100 images with a sampling distance of 200 nm along the z and 107 nm in the w direction. The constrained iterative algorithm option was used (45).

3D Image Analysis for Telomeres. Telomere measurements were done with TELOVUE (9, 46). By choosing a simple threshold for the telomere, a binary image is found. Based on that, the center of gravity of intensities is calculated for every object resulting in a set of coordinates (x, y, z) denoted by crosses on the screen. The integrated intensity of each telomere is calculated because it is proportional to the telomere length (47). The integration region is determined by growing a sphere on top of the found coordinate. After every step of growth (iteration), the sum under this volume (the telomere) is subtracted by the sum just surrounding it (background level). When the process of the growth of the sphere does not contribute to an integrated intensity increase, the algorithm stops and the integrated intensity of the telomere with an automatic background correction is obtained.

Chromosome Painting and Measurements of Chromosomal Overlap(s) in Interphase Nuclei. Chromosome painting was carried out as described in ref. 48 by using paints for mouse chromosomes 5 (Cy3), 13 (FITC), 7 (Cy5), 10 (FITC), and 17 (FITC) from Applied Spectral Imaging (Vista, CA). 3D image acquisition of painted nuclei was performed as described above. Measurements of chromosomal overlaps were performed after 3D image acquisition and constrained iterative deconvolution serving as follows: (i) based on the DAPI counterstain image, we determined the 3D boundary of the nuclear volume. Data outside that volume were ignored. (ii) For each one of the chromosomes, we determined an intensity threshold and referred only to those that were above the threshold that belonged to the specific chromosomes. The total volume occupied by each one of the chromosomes is measured (V1 and V3, Figs. 1-3). The volume occupied by both chromosome pairs is measured, V13, by dividing this value by V1 and by V3, the level of overlap relative to the total volume of each chromosome pair was measured, V13/ V1, V13/V3 (for details, see Fig. 8 which is published as supporting information on the PNAS web site). Spectral Karyotyping (SKY). Mouse SKY was performed using a SKY system (Applied Spectral Imaging) (37). Twenty metaphases were examined per time point. Significant values for chromosomal rearrangements were determined after MycER activation. Mean total chromosome numbers and numbers of each chromosome observed for control and Myc-activated cells were computed over time by two-way ANOVA. In addition, statistical analyses were performed for the occurrence of translocations, inversions, and isochromosomes in the experimental period of 120 h. P values of <0.05 were considered significant. Only the frequency procedure was used, followed by Fisher's exact test. The P value of the overall study was <0.0001.

Supporting Information. For additional information, see Figs. 9-12, Movies 1-3, and Tables 2-4, which are published as supporting information on the PNAS web site.

Results

The 3D Organization of Telomeres before c-Myc Activation. We examined whether c-Myc deregulation affected the 3D organization of telomeres in the interphase nucleus. To this end, we analyzed the effect of conditional c-Myc expression in two independent immortalized mouse B lymphocyte lines, Ba/F3 (50) and P3 (50), stably transinfected with MycER (38). For both cell lines, we first evaluated the 3D organization of telomeres in nuclei of non-MycER-activated cells by using primary BaL/c B lymphocytes as a control. Consistent with our previous studies (50), telomeres of normal primary BaL/c B nuclei showed nonoverlapping telomere position in interphase nuclei of primary and immortalized B lymphocytes without overlap in telomere positions. (a) Primary B cell nuclei. (b) Nucleus of normal BaL/c B cell. (c) Nucleus of immortalized BaL/c B cell. Telomeres are shown in red in nuclei in blue. 3D = 3D fluorescence; TDS, 3D side view.
Fig. 1. c-Myc deregulation induces TAs in interphase nuclei of PrEB and Bu/F3 cells (a, b, and c). (a) PrEB cells show normal overlapping telomeres (red arrow). (b) PrEB cells show normal overlapping telomeres (green arrow). (c) PrEB cells show normal overlapping telomeres (green arrow). (d) MycER-activated PrEB cells with TAs (green arrow). (e) PrEB cells show normal overlapping telomeres (green arrow). (f) MycER-activated Bu/F3 cells show the formation of TAs (green arrow).

Fig. 4. c-Myc-induced telomeric aggregates appear in cycles. (A) Conditional c-Myc induction causes TA formation. (Aa) Negative control: non-MycER-activated PrEB nuclei with overlapping 3D telomeric nuclear regions. (B) MycER TAs of various sizes and numbers are present after conditional c-Myc expression at any given time point of TA formation. Telomeres are shown in green, TA by red arrows. (Bb) Positive control. Prostate carcinoma cell line, MDaPC4 (G4), with constitutive c-Myc expression due to T215; shows TAs. Similar results were obtained with primary prostate cancer cells (data not shown). (Bc) c-Myc-induced cycles of TAs. Fold increase in TAs over control levels during a period of 120 h. During this period, c-Myc had been upregulated for different lengths of time (see Fig. 3). Black, 0 h; red, 12 h; and green, 48 h. All results were performed in triplicate. (Bd) c-Myc expression in TAs. c-Myc expression in TAs who had been exposed to 4HT (100 ng/ml) for 120 h. c-Myc expression was detected by anti-c-Myc antibody (red arrows; green arrow, 24 h). Gray, control cells. (Be) c-Myc-induced cycles of TAs. Fold increase in TAs over control levels during a period of 120 h. During this period, c-Myc had been upregulated for different lengths of time (see Fig. 3). (Bf) c-Myc expression in TAs. c-Myc expression in TAs who had been exposed to 4HT (100 ng/ml) for 120 h. c-Myc expression was detected by anti-c-Myc antibody (red arrows; green arrow, 24 h). Gray, control cells. (Bg) c-Myc-induced cycles of TAs. Fold increase in TAs over control levels during a period of 120 h. During this period, c-Myc had been upregulated for different lengths of time (see Fig. 3). (Bh) c-Myc expression in TAs. c-Myc expression in TAs who had been exposed to 4HT (100 ng/ml) for 120 h. c-Myc expression was detected by anti-c-Myc antibody (red arrows; green arrow, 24 h). Gray, control cells. (Bi) c-Myc-induced cycles of TAs. Fold increase in TAs over control levels during a period of 120 h. During this period, c-Myc had been upregulated for different lengths of time (see Fig. 3). (Bj) c-Myc expression in TAs. c-Myc expression in TAs who had been exposed to 4HT (100 ng/ml) for 120 h. c-Myc expression was detected by anti-c-Myc antibody (red arrows; green arrow, 24 h). Gray, control cells. (Bk) c-Myc-induced cycles of TAs. Fold increase in TAs over control levels during a period of 120 h. During this period, c-Myc had been upregulated for different lengths of time (see Fig. 3). (Bl) c-Myc expression in TAs. c-Myc expression in TAs who had been exposed to 4HT (100 ng/ml) for 120 h. c-Myc expression was detected by anti-c-Myc antibody (red arrows; green arrow, 24 h). Gray, control cells. (Bm) c-Myc-induced cycles of TAs. Fold increase in TAs over control levels during a period of 120 h. During this period, c-Myc had been upregulated for different lengths of time (see Fig. 3). (Bn) c-Myc expression in TAs. c-Myc expression in TAs who had been exposed to 4HT (100 ng/ml) for 120 h. c-Myc expression was detected by anti-c-Myc antibody (red arrows; green arrow, 24 h). Gray, control cells. (Bo) c-Myc-induced cycles of TAs. Fold increase in TAs over control levels during a period of 120 h. During this period, c-Myc had been upregulated for different lengths of time (see Fig. 3). (Bp) c-Myc expression in TAs. c-Myc expression in TAs who had been exposed to 4HT (100 ng/ml) for 120 h. c-Myc expression was detected by anti-c-Myc antibody (red arrows; green arrow, 24 h). Gray, control cells.
Table 1. Apoptosis levels in non-MycER and MycER-activated PreB cells

<table>
<thead>
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<th>Time, h</th>
<th>Controls</th>
<th>MycER-activated</th>
<th>Fold Increase</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>3.0</td>
<td>3.0</td>
<td>1.0</td>
</tr>
<tr>
<td>12</td>
<td>2.0</td>
<td>4.0</td>
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<td>24</td>
<td>6.0</td>
<td>12.0</td>
<td>2.0</td>
</tr>
<tr>
<td>30</td>
<td>4.0</td>
<td>10.0</td>
<td>2.5</td>
</tr>
<tr>
<td>42</td>
<td>2.0</td>
<td>8.0</td>
<td>4.0</td>
</tr>
<tr>
<td>48</td>
<td>5.0</td>
<td>10.0</td>
<td>2.0</td>
</tr>
<tr>
<td>60</td>
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<td>11.0</td>
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<td>1.7</td>
</tr>
<tr>
<td>96</td>
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<td>3.0</td>
<td>1.0</td>
</tr>
<tr>
<td>102</td>
<td>2.0</td>
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</tr>
</tbody>
</table>

The nature of c-Myc-induced 3D structural changes in interphase nuclei of conditionally Myc-expressing cells was as follows: at peaks of TA formation and thereafter, a significant increase in end-to-end chromosomal fusions over control levels was observed. This result was followed by a significant increase in broken chromosomes and non-reciprocal translocations (Figs. 5d and 6 and Table 2). In conclusion, TA cycles activate BFB cycles, namely the fusions of two chromosomes, consequently, the formation of dicentric and their subsequent breakage in anaphase (Fig. 5). These cycles are induced by conditional Myc deregulation and lead to the onset of genomic instability, demonstrated by the chromosomal rearrangements resulting from these BFB cycles (Figs. 5 and 6 and Table 2).

Next, we investigated whether cells with TAs died during the course of the experiments. If this possibility was the case, we would expect a correlation of cell death in Myc-activated cells at the peak of TA formation or shortly thereafter. The level of apoptosis was 2-fold higher in Myc-activated cells than in control cells (Table 1). There was no preference for apoptotic cell death for any specific time point during the 120 h. We concluded that BFB cycles, not apoptosis, contributed to the cycles of TA formation.

3D organization of Chromosomes in c-Myc Activated Interphase Nuclei. TAs and the initiation of BFB cycles with subsequent chromosomal rearrangements prompted us to investigate whether chromosomes were affected in their 3D nuclear positions during MycER activation. To this end, we examined the overlap of specific chromosomes over the 120 h period. SKY of MycER-activated PreB cells suggested chromosomal rearrangements involving chromosomes 7, 13, and 17. Additional rearrangements were found but

Fig. 5. Molecular cyrogenetic evidence of BFB cycles in MycER-activated PreB cells. (A) SKY analysis reveals telomeric fusions and chromosome breakage. (a) Upper) Metaphase, raw image (Left); metaphase, classified image (Center); and metaphase, inverted DAPI image (Right). (b) Lower) Spectral karyotype. End-to-end fusion of chromosomes 18 and 4 (red arrow) and fusion of chromosome 1 with a broken piece of chromosome 1 (green arrow) are shown. One broken chromosome 1 is isolated. Note additional broken chromosomes 1, 2, 3, and 7. (c) End-to-end fusions of chromosomes revealed by telomere FISH. (d) Upper) Centromeric fusion (see arrow and insert). (e) Lower) Telomeric fusion (see arrow and insert). (f) Upper) Chromosome bridges. (g) Upper) Short exposure of DAPI-stained nuclei (100 ms). (lower) Longer exposure (900 ms) of same image makes anaphase bridge visible (white arrow) but not metaphase. (h) p20 illustrating chromosome fusions (red arrow) and non-reciprocal translocations (white arrow). Broken chromosomes are also present (chromosomes 4, 6, 12, and 17).
did not reach significant levels (data not shown). We examined three combinations of chromosomes over a 96-h period. This period covered all peaks of TA formation (Fig. 4B). As shown in Fig. 7, we observed a change in overlap between chromosomes 5 (red) and 13 (green) over the time course (Figs. 7 A and B). Both chromosomes were found in close vicinity as the cells entered into the first TA cycle. Chromosomes 10 (green) and 7 (red) also showed increases in the percentage of overlap (Fig. 7 A and B), as did chromosomes 7 (red) and 17 (green) (Fig. 7 A and B). Representative 3D movies are shown in Movies 1–3.

Discussion

**c-Myc Induces Telomeric Aggregates, Fusions, and BBF Cycles.** Previous studies have shown that c-Myc triggers a complex network of genomic instability at the level of single genes (14, 15, 19) and whole chromosomes (16–18) (for review, see ref. 15). In addition, c-Myc induces illegitimate replication initiation (19, 20), chromosomal rearrangements (18), DNA breakage, alterations of DNA repair (21–23), and a low level of point mutations (24, 25). A previously uncharacterized mechanism underlying c-Myc-dependent genomic instability at the chromosomal level directly affects the integrity of the telomeres and was revealed in this study.

The clear periodicity of the TA cycles that was found with four different Myc-activating treatments suggested a biological relevant Myc-dependent process. Theoretically, cycles of Myc-induced TAs could reflect (i) nuclear remodeling with the transient association and subsequent dissociation of telomeres; (ii) end-to-end chromosomal fusions that initiate BBF cycles (52, 53); (iii) c-Myc induced cell death; and (iv) a combination of all of the above. Our data are consistent with BBF cycles and exclude apoptosis as a direct contributor to the TA cycles. Apoptosis occurred at equal levels throughout the study and consistently reached about twice the levels seen in the control cells. The loss of cells was compensated by a 2-fold increase in proliferation in Myc-ER-activated PreB (19). These data also indicate that there is genetic separation of genomic instability and apoptosis as reported in ref. 54. Whether telomere associations and dissociations (33) contributed to the TA cycles is presently unknown.

Direct evidence of BBF cycles in the periodicity of TAs came from a detailed analysis of chromosomal fusions, breakage, and rearrangements observed over the time course of five TA cycles. We demonstrated the occurrence of end-to-end fusions that generated dicentric chromosomes and breaks during anaphase, leaving one chromosome or chromatid with a piece from another chromosome or chromatid. The resulting telomere-free ends continue to undergo fusions with other chromosomes, a cycle of events termed BBF cycle (52, 53). Experimental data suggest these events from fusions to breakages and nonreciprocal translocations. The periodicity of the TA cycles is consistent with a ~12-h population doubling time of the PreB cells (30). Each peak of TAs is consistent with the repeated formation of TAs. Time points after the peak are in agreement with the breakage of dicentric chromosomes. Telomere-free ends initiate new BBF cycle(s) until no more telomere-free chromosomal end(s) persist.

**From Telomeres to Chromosomal Rearrangements: A New Pathway of c-Myc-Dependent Genomic Instability.** Muller (52) and McChesney (55) first described BBF cycles, a mechanism of chromosomal end-to-end fusion that contributes to the onset of genomic instability. BBF cycles contribute to deletions, gene amplification, nonreciprocal translocation, and overall genetic changes that are associated with tumorigenesis (56–60).
Our study showed that c-Myc is the key factor that initiates genomic instability through BDF cycles. Such BDF cycles in telomerase-negative immortalized mouse PBX cells (unpublished data) with long telomeres are distinct from BDF cycles reported for critically short telomeres (51, 52). Some TAs (but not necessarily all) represent fusions, as evident by the analysis of metaphase chromosomes. TAs and end-to-end fusions depend on time and levels of c-Myc activation. Analysis of frequencies of both events showed that they are closely linked. As the fusions initiate BDF cycles, the frequencies of breakage and nonreciprocal translocations increase over time.

A previously uncharacterized pathway of c-Myc-dependent genomic changes thus starts at the telomeric ends of the chromosomes. Both TAs and IBF cycles are the manifestation of deregulated Myc expression, leading to chromosomal rearrangements and telomere misbehavior to genomic instability.

Local chromosomal movement increases chromosomal overlap in the nucleus. This temporal change in local positioning may permit the direct contact of chromosomal ends and facilitate recombinations and, for instance, such movements were observed after c-Myc deregulation and suggested an impact of the oncprotein on local nuclear positioning of chromosomes. Chromosome movements were previously studied and found by others as well (53–60). Several regulatory pathways involving oncogene deregulation may affect the 3D nuclear organization. Oncoproteins, including c-Myc, can alter the 3D nuclear organization and the organization of chromatin (70–72). They also affect the nuclear matrix. High mobility group protein 1(Y) (HMGB1) is a c-Myc-dependent nuclear matrix protein (73) with increased expression during neoplasia (2). The analysis of myc-binding sites in the human genome suggests that c-Myc binds to genes encoding nucleoskeletal components (74). Furthermore, constitutive c-Myc expression was shown to be associated with the down-regulation of the telomere repeat binding protein TRF2 (10), a protein required for telomere capping and genome stability (75). Myc is also involved in the regulation of DNA repair (22, 23) and has been shown to induce DNA breakage (21). This taken together, many different c-Myc-dependent mechanisms could potentially affect the nuclear organization and, as shown here, converge at the telomeres.

We thank Dr. Michael Mowat for critical reading of this manuscript, Mary Chang for statistical analysis, and Cheryl Taylor-Kashian and Lamour Whitt for their help. The work was supported by the Canada Foundation for Innovation, the Canadian Institutes of Health Research, CanadCare Manitoba (S.M.), Fondation de France (P.T.) and French Minister of Foreign Affairs (T.F.), Sander-Stiftung, and Deutsche Krebsforschung (F.B.). The Physics for Technology program of the Foundation for Fundamental Research in Matter, the Delft Inter-Faculty Research Center Life Tech, Cytonet, and the Delft Research program Life Science and Technology (Delft, The Netherlands).
III. DISCUSSION

3.1 Telomere and Its 3D Nuclear Organization in Normal and Tumor Cells

Recent advancements in molecular imaging allow us to study the temporal and spatial organization of telomeres in the interphase nucleus of mammalian cells in 3D space. Weierich et al., 2003 and Chuang et al., 2004 have shown that telomeres are organized in a dynamic cell cycle- and tissue-dependent manner in normal cells (Chuang et al., 2004; Weierich et al., 2003). The 3D telomere distribution in the nucleus can be determined by the ratio of two different nuclei, or a/c ratio (Chuang et al., 2004; Vermolen et al., 2005). When the cells are in G0/G1 and S phases of the cell cycle, the telomeres distribute throughout the entire nuclear space and give a small a/c ratio. When the cell is in G2 phase, telomeres align in the center of the nucleus and form a telomeric disk (Chuang et al., 2004) and give a large a/c ratio due to the disk-like organization of telomeres. Chuang et al. described this by using the primary mouse lymphocyte as an example. The a/c ratios for telomeric position in G0/G1, S, and G2 are 1.4±0.1, 1.5±0.2, and 14±2 respectively. Dynamic telomere organization has been visualized by live cell imaging of human osteosarcoma (U2OS), human cervical carcinoma (HeLa), and mouse MS5 cells. Both short and long ranges of movements were observed over a period of 20 minutes (Molenaar et al., 2003). In addition to cell cycle dependency, dynamic telomere movement also depends on the shape of the nucleus (Chuang et al.,
2004; Ermler et al., 2004). In vivo mobility studies by Bronstein et al. have shown that telomere dynamics govern the short-term anomalous diffusion while telomere binding governs the long-term diffusion which altogether contribute to 3D telomeric organization in the nucleus. Therefore we conclude that telomere organization within the nucleus is dynamic, moving in cell cycle- and cell type-dependent manner (Bronstein et al., 2009; Chuang et al., 2004).

Recent work presented by De Vos et al. confirmed our findings of telomere organization (De Vos et al., 2009). Microterritories occupied by several telomeric ends are visualized using the controlled light exposure microscopy (CLEM). Such close proximity of telomere neighborhood organization can facilitate recombination at the subtelomeric regions which can lead to genomic instability. Additional work can be done in the future with high-resolution microscopy methods to better delineate the chromosomal ends.

There are two major phenotypes of telomeric dysfunction in tumor cells. The first phenotype is critically short telomeres (DePinho and Polyak, 2004). The second phenotype is formation of telomeric aggregates (TAs) independent of telomere size or telomerase activity (Chuang et al., 2004; Louis et al., 2005). It was shown that each telomere of a normal cell occupies its specific 3D space within the nucleus without overlapping or forming aggregates with other telomeres (Chuang et al., 2004). In contrast, different sizes and numbers of TAs are observed in tumor cells (Chuang et al., 2004; Louis et al., 2005). Both phenotypes can lead to breakage-bridge-fusion (BBF) that contribute to genomic
instability (Artandi et al., 2000; DePinho and Polyak, 2004; Murnane and Sabatier, 2004).

3.2 Telomere Remodeling and Chromosomal Aberrations in the 3D Interphase Nucleus

Dynamic alterations of 3D nuclear structures can potentiate genomic instability. When telomeres form aggregates in the 3D nucleus of tumor cells, some of them fuse together and form dicentric chromosomes. When these end-to-end fused chromosomes divide, they form the anaphase bridge first and then break apart, leaving one chromosome with a terminal deletion and another one with a translocated piece. Both chromosomes have telomere free ends, which represent double-stranded DNA break, can fuse with other chromosomes. This BBF cycle was first described by Muller and McClintock (McClintock, 1941; McClintock, 1942; Müller, 1938). Louis et al. showed conditional c-Myc deregulation can induce cycles of telomeric aggregates in interphase nuclei. The resulting BBF cycles lead to chromosomal rearrangements and the onset of genomic instability (Louis et al., 2005).

3.3 Mechanism of Altered Telomeric Nuclear Organization

In the study by Louis et al conditionally de-regulated c-Myc oncogene leads to generation of cycles of TAs over 144 hours that was past the initial c-Myc deregulation (Louis et al., 2005). This suggests that the oncoprotein is no longer required for the downstream effects of subsequent cycles of TA formation.
The time of c-Myc deregulation was proportional to the number of TA cycles observed. As a result of c-Myc deregulation, TA formation precedes chromosomal end-to-end fusions and the onset of chromosomal instability. Fusion of telomeric ends can be confirmed by the presence of inter-nuclear bridge and dicentric chromosomes as cells progress through the cell cycle.

c-Myc-induced TA formation is independent of critical shortening of telomeric end and telomerase activity. However, formation of TA has been observed in critically shortened telomeres in Hodgkin lymphoma cells (Knecht et al., 2010).

It has been shown that Myc box II mutant is unable to induce TA formation (Caporali et al., 2007). This indicates the Myc box II is a conserved element within the N-terminus of c-Myc is needed for all known functions of c-Myc (Stone et al., 1987). Only full length c-Myc but not myc box II mutant Myc is able to induce tumor formation in SCID mice (Fest et al., 2005).

3.4 Applications of 3D Telomere and Chromosome Organization in Medicine

3.4.1 Mechanism of Disease

Formation of TAs have been described in various tumor cell lines and primary tumors, including human Burkitt lymphoma, neuroblastoma and colon carcinoma and primary mouse plasmacytoma and primary human head and neck squamous cell carcinoma and glioblastoma (Chuang et al., 2004; Gadji et al.,
2010; Mai and Garini, 2006), It is shown that the telomere dysfunction is involved in the transition of the Hodgkin cells (H cells) to the Reed-Sternberg cells (RS cells) (Knecht et al., 2009). In addition, significant telomere shortening and formation of TAs were observed in RS cells in EBV positive Hodgkin’s lymphoma (Knecht et al., 2010). These telomere-poor or telomere-free “ghost” nuclei are formed in end-stage tumor cells which are incapable of dividing. Additional work to elucidate the structural organization of chromosomes in the RS cells will be of interest in understanding the nature of progression of Hodgkin’s lymphoma.

3.4.2 Cancer Biomarkers

The topic of cancer biomarkers is broad and involves multiple disciplines of science and technology. The 3D positioning patterns of telomeres and chromosomes can be used as diagnostic and prognostic biomarkers in the near future. Specific 3D alterations in the relative arrangement of chromosome territories may be specific to a disease entity. As telomere dysfunction occurs prior to chromosomal re-arrangement (Louis et al., 2005), telomeric aggregates can be an attractive biomarker for early detection.

In the paper presented by Gadji et al, the 3D telomeric profile that was characteristic of each group correlated with short-term, intermediate and long-term survival and time to progression in a patient cohort of 11 individuals (Gadji et al., 2010). The three telomeric signatures including the total number of telomeres and the nuclear distribution and presence of TAs were evaluated using
the Teloview program and the SpotScan system. This study indicates a correlation between clinical outcome parameters and 3D telomere organization. In addition, the new SpotScan system provides a high throughput means to assess TAs. This may be important in time-critical situations such as assessing intra-operative frozen sections performed during oncologic surgery. The 3D nuclear structure thus can be of prognostic, predictive value and possibly provide pharmacodynamic biomarkers, to provide information on natural course of disease, treatment response and optimal dose of experimental therapeutic agents, respectively.

3.4.3 Molecular Marker in Other Disease Processes

The first observation of TAs in 3D space was made in tumor cells. However, additional studies have confirmed that TAs are present in other conditions and diseases in different tissue types and body fluids. Examples include the formation of TAs with various numbers and sizes during senescence of human mesenchymal stem cells (Raz et al., 2008) and were found in the placenta of pregnant women with pre-eclampsia (Sukenik-Halevy et al., 2009), amniocytes from pregnant women carrying trisomy 21 fetuses (Hadi et al., 2009) and leukocytes from chronic hepatitis C patients (Amiel et al., 2009). The potential mechanisms for TA formation in different diseases may include oxidative stress, inflammation and infection. Therefore we propose that TAs can be used as a new diagnostic and prognostic tool for different disease entities. TAs may represent a new class of biomarker, namely the optical biomarker.
Future studies to unravel the nature of disease and the process of the disease development and disease progression can be performed from a different perspective via observation of aberrant 3D organization of telomere and chromosomal organizations. The latter will in turn allow us to understand “cancer as a disease of DNA organization and cell structure” (Pienta et al., 1989) and other disease processes in humans.
IV. References


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