Chromosome positioning in interphase nuclei of hematopoietic stem cell and myeloid precursor

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Abstract

Human myelopoiesis is an intriguing biological process during which multipotent stem cells limit their differentiation potential generating precursors that evolve into terminally differentiated cells. The differentiation process is correlated with differential gene expression and changes in nuclear architecture. In interphase, chromosomes are distinct entities known as chromosome territories and they show a radial localization that could result in a constrain of inter-homologous distance. This element plays a role in genome stability and gene expression. Here, we provide the first experimental evidence of 3D chromosomal arrangement considering two steps of human normal myelopoiesis. Specifically, multicolor 3D-FISH and 3D image analysis revealed that, in both normal human hematopoietic stem cells and myelod precursors CD14-; chromosomal position is correlated with gene density. However, we observed that inter-homologue distances are totally different during differentiation. This could be associated with differential gene expression that we found comparing the two cell types. Our results disclose an unprecedented framework relevant for deciphering the genomic mechanisms at the base of normal human myelopoiesis.

Introduction

The organization of a eukaryotic nucleus reflects its specific expression profile, with dynamic rearrangements and repositioning involved in the regulation of gene expression and differentiation. Chromosomes represent the largest nuclear structural units and, in interphase, are organized in distinct entities known as chromosome territories (CTs) whose radial arrangement is nonrandom.1-3 In general CTs show a gene density driven positioning in spherical nuclei while a chromosome size driven positioning in ellipsoid nuclei.4 Moreover CTs radial localization preference could result in specific interchromosomal arrangement which are diverse in different cell types,3-7 during cell differentiation, development,8-10 and in cancer.11-13 The preferential spatial proximity of genomic loci likely enhances the probability of reciprocal chromosome rearrangement as chromosomal translocations,14,15 typical of pathologies such as Acute Myeloid Leukemia (AML).

On the other end it has been recently observed that homologous chromosomes are more distant than heterologous and a possible explanation is that the separation of homologous chromosomes could allow the maintenance of two copies of mono-allelically expressed genes at a distance, avoiding their co-regulation, whereas other mechanisms would control transient inter-chromosomal contacts required for differential allelic expression.16

Given these evidences, the human myelopoiesis, i.e., the process that leads to differentiation of myelopoietic cell lineages (erythroid, megakaryocytic, granulocytic and mono/macrophage), represents an ideal model to investigate chromosome positioning. Higher order chromatin organization during myelopoiesis has been recently studied considering specific chromatin features,17 while not chromosome position. Moreover several studies have demonstrated a differential gene expression during myeloid transition.18-20

In the present study, we used multicolor 3D-FISH and image analysis to reconstruct the chromosomal positioning during two steps of differentiation: normal human hematopoietic stem cells and derived myeloid precursors CD14-. Moreover we have measured the distance between the edges of homologs in the two cell types and we have observed a significant difference that can be associated with a differential gene expression, which occur during differentiation.

Materials and Methods

Ethics statement

Human CD34+ cells were purified upon donor’s informed written consent from umbilical Cord blood (CB) samples. Mononuclear cells were isolated by Ficoll-Hypaque (Lymphoprep; Nycomed Pharma, Oslo, Norway) gradient separation and washed twice with phosphate-buffered saline, and then CD34+ cells were separated using magnetic cell sorting procedure (EasySep Human CD34+ positive selection kit; StemCell Technologies, Vancouver, BC, Canada). CD34+ cell purity assessed by flow cytometry was ≥95% (Supplementary Figure S1). After immunomagnetic separation, CD34+ cells were seeded in 24-well plates at 5 ×105/mL in Iscove’s modified Dulbecco’s medium (IMDM) (GIBCO, Grand Island, NY, USA) containing 20% Human Serum (Bio-Whittaker, Walkersville, MD, USA), SCF (50 ng/mL), Flt3-ligand

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(Flt3L) (50 ng/mL), TPO (20 ng/mL), IL-6 (10 ng/mL) and IL-3 (10 ng/mL) (all from R&D Systems, Minneapolis, MN, USA) and fixed 24 hours later.

CD14+ myeloid precursors were obtained by in vitro differentiation of CB derived CD34+ cells performed as already described. Briefly, CB CD34+ cells were cultured in IMDM added with 20% FCS (Bio-Whittaker, Walkersville, MD, USA), in the presence of human hematopoietic cytokines: SCF (50 ng/mL), Flt3-ligand (Flt3-l) (50 ng/mL), IL-6 (10 ng/mL) and IL-3 (10 ng/mL) (all from R&D Systems, Minneapolis, MN, USA). After 7 days of culture, hematopoietic cells were analyzed, by flow cytometry, for CD14 antigen expression, estimated at about 15-20% of the entire cell population. Then CD14+ and CD14- cell fractions were obtained by immunomagnetic separation using a cell sorting procedure (EasySep® Human CD14 Positive Selection Kit, StemCell Technologies).21

Cell fixation, and FISH pretreatment

For 3D-FISH experiments, cells were allowed to attach to poly- L-lysine-hydrobromide (Sigma) coated cover slips. Cells were fixed in 4% paraformaldehyde (PFA) in 0.3 × PBS for 10 min. Permeabilization steps included treatment with 0.5% Triton-X100 (15 min), with 20% glycerol in PBS for at least 1 h, repeated freeze/thawing in liquid nitrogen, and incubation in 0.1 N HCl (5 min) and pepsin solution (2.5 mg/mL pepsin in 0.01 N HCl at 37°C for 5-10 min).10 Slides were stored at 4°C in 50% formamide/2 × SSC. This technique is commonly used to preserve the 3D nuclear architecture and to study chromosome territory topology.

Probe labeling, hybridization, and detection

For 3D-FISH, the hybridization and detection protocol described by Solovcr12 was followed. All DNA probes were labeled in the presence of Biotin-dUTP (Roche) or TexasRed- dUTP (Molecular Probes), respectively, and mixed with 10-fold excess of human Cot-1 DNA. In situ hybridization was performed for 48 h, followed by stringency washes for 3 × 5 min in 0.1 × SSC (60°C). Biotinylated probes were detected with Avidin-Alexa 488 (Molecular Probes). Metaphase chromosomes and 3D fixed interphase nuclei were counterstained for 10 min with DAPI (2 μg/mL).

Digital microscopy and image processing

Metaphase FISH images were captured with a Nikon Eclipse 90i system, equipped with Nomarski differential interference contrast (DIC) optics. Samples were photographed with a DS-5Mc Nikon digital camera and the resulting photographs were analyzed by using the Nikon AcU2 software program. Light optical serial sections of nuclei studied in three color 3D-FISH experiments were obtained with a laser scanning confocal microscope (LSM 410; Carl Zeiss MicroImaging) equipped with Ar and He/Ne lasers. For imaging of multicolor 3D-FISH experiments a Leica TCS SP2 laser scanning confocal microscope (Leica Microsystems) with beam splitters tuned for DAPI, Alexa 488, and TexasRed was used. Nuclei were scanned with an axial distance of 200 nm between consecutive light optical sections yielding separate stacks of 8-bit gray-scale images for each fluorescence channel with a pixel size of 60 nm. For each optical section, images were collected sequentially for all fluorochromes used, followed by correction of the axial chromatic shift for each fluorescence channel as described by Walter et al.21 Confocal image stacks were processed with ImageJ software (http://rsb.info.nih.gov/ij) using the deconvolution plugin to enhance resolution.

Quantitative 3D data evaluation and statistical analysis

Quantitative 3D evaluation of light-optical serial sections was performed using voxel-based software algorithms eADS (enhanced Absolute 3D Distances to Surfaces). eADS software was used to determine the shortest distances (in nm) between CT signals and the nuclear border and between the surface of the corresponding CT. The surface of the reference structure (CT, nucleus) was determined by applying a user set threshold: all voxels with intensities below the set threshold are set to zero. The surface of the reference structure contains all voxels with an intensity >0, which are directly adjacent to at least one voxel with an intensity equal to 0.16,24-26 For evaluations of distance measurements inside the nucleus deconvolution and normalization was...
performed to facilitate the comparison of different experiments. To test the data set for significant differences (P<0.05), the Mann-Whitney Rank Sum Test (U-Test) was applied on the results of the quantitative evaluation programs described above.

Chromosome territory volume (V), surface (S), and a roundness factor (RF) were computed. RF is defined as $RF = \frac{\pi V^2}{S^3}$, $0 < RF \leq 1$.

We calculated nuclear radii (Rn) and diameters (Dn) from $Rn = (3V/4\pi)^{1/3}$, where V is the thresholded nuclear pixel volume. We normalized the absolute spatial separations between chromosomes (CT1:CT2) as a fraction of nuclear diameter ((CT1:CT2)/Dn) to account for natural variations in nuclear size, which may influence relative positioning.

Edge-to-edge distances were assessed using the absolute distance to surface (ADS) program. This program allowed for quantification of the shortest distance of given sub-CT units with respect to a boundary structure in 3D. In these evaluations, overlapping homologues result in a null distance. The statistical significance of the inter-homologue and distances was tested using Student’s t-test.

**Analysis of gene expression profiles**

Gene expression profiles of 12 hematopoietic stem cell (CD34+) and 11 progenitors (CD14-) samples were obtained from four different datasets downloaded from gene Expression Omnibus (Supplementary Table S1). Briefly, expression values were generated from Affymetrix HG-U133A raw intensity signals using the HG-U133A custom definition files for human GeneChips (version 15.0.0) based on Entrez gene (http://brainarray.mbi.ucsf.edu/Brainarray/Dataset/CustomCD F/15.0.0/entrezg.asp). Intensity values for a total of 12080 probe sets have been background adjusted, normalized using quantile normalization, and gene expression levels calculated using median polish summarization (Robust Multiarray Analysis, RMA). Differentially expressed chromosomal regions between CD34+ and CD14- samples have been identified using the LAP procedure, with the Significance Analysis of Microarray algorithm (SAM) two-class statistics coded in the PREDA R package.

The percentage of false positive predictions was estimated with 10000 permutations and the q-value (i.e., False Discovery Rate, FDR) threshold was set to 0.5. Functional over-representation analysis has been performed using DAVID (http://david.abcc.ncifcrf.gov/) and Gene Ontology Biological Process categories.

**Results and Discussion**

We first used multicolor 3D-FISH and 3D image analysis to localize chromosome positions with respect to the nuclear border in HSC CD34+ and myeloid precursor CD14- cells. Specifically, we investigated a total of six chromosomes, i.e., chromosome 6, 8, 9, 15, 17, and 21, which are commonly implicated in translocations associated with acute myeloid leukemias, with the aim of evaluating if their nuclear position shows a preferential sub nuclear localization in each of the two cell populations. When considering the average chromosome position with respect to the nuclear border (Table 1) and the distribution of the chromosome territory (in terms of relative DNA content with respect to the normalized distance to the nuclear border; Figure 1A), hematopoietic stem cells and myeloid precursors share the same localization for chromosomes 6, 21, 15, and 17 (P>0.05) while the position of chromosomes 8 and 9 resulted significantly different (P=0.006 and P=0.023, respectively). Chromosome positions are consistent with average gene density, quantified as the ratio between the number of genes and the length in Mb of each chromosome (Table 1). In particular, chromosome 17, which has the higher gene density (21.96 gene/Mb), has the more internal localization in both cell types (HSCs=−1746.33 nm; myeloid precursors =−2312.43 nm). On the contrary, chromosome 8 has the lower average gene density (9.02 gene/Mb) and is very peripheral in both cell types, although its position is closer to the nuclear border in precursors (Figure 1A).

Interestingly, chromosome 9, characterized by an average gene density slightly higher than that of chromosome 8 (10.80 as compared to 9.02), results more internal in both the cell types, and particularly in myeloid precursors. Chromosome distances from the nuclear border allowed reconstructing a plausible map of the chromosome distribution during the differentiation of hematopoietic stem cells (Figure 1B) to myeloid precursors (Figure 1C). Homologous are shown as a single entity. Overall, positional data and the chromosomal positional map indicate that no drastic changes occur during differentiation of HSCs into CD14- myeloid precursors as regards to the chromosomal preferential localization within sub nuclear regions. This evidence is in perfect agreement with the relative small number of chromosomal regions found differentially expressed when comparing the gene expression profiles of CD34+ HSCs and CD14- myeloid precursors. Indeed, the regional analysis of transcriptional levels identified only 19 and 21 chromosomal regions up-regulated in HSCs and precursor cells, respectively (Supplementary Tables S2 and S3). Nevertheless, the functional annotation of differentially expressed regions found in this comparison supports a link between transcriptional differences and changes of chromatin organization during the transition from the stem to the precursor level, as recently highlighted. To investigate if indeed chromatin compaction changes during the transition from stem cells to precursors, we compared chromosome volumes, calculated from the images voxels, and found that the two cell types significantly differ for the

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Table 1. Average chromosome position with respect to the nuclear border in hematopoietic stem cells (CD34+) and myeloid precursors (CD14-). P-values are calculated with a Mann-Whitney rank sum test (U-Test). The number of analyzed nuclei per chromosome per cell type was 30.

| Chromosome | Average gene density (gene/Mb) | Average chromosome position (nm) | P-value |
|------------|-------------------------------|----------------------------------|---------|
|            | HSC (CD34+)                   | Precursors (CD14-)               |         |
| 6          | 12.01                         | −1149.25                         | −1266.07 | 0.098 |
| 8          | 9.02                          | −1154.85                         | −794.01  | 0.006 |
| 9          | 10.80                         | −1174.93                         | −1427.96 | 0.023 |
| 15         | 12.35                         | −1452.98                         | −1899.18 | 0.261 |
| 17         | 21.96                         | −1746.33                         | −2312.43 | 0.245 |
| 21         | 9.35                          | −1573.82                         | −1345.54 | 0.4   |
of gravity which is commonly use to study CT organization may not reflect interactions of these CT at their borders, for this kind of analysis we have performed an edge-to-edge evaluation. We found that the inter-homologue distances were always shorter in precursor, significantly for chromosome 6, 9, and 15 (Figure 2 and Table 3).

Since the localization of the chromosomes is always consistent with gene density correlated distribution, the differences found can be most likely attributed to differential gene expression that can affect chromatin organization and compaction during the transition from the stem to the precursor level.

**Conclusions**

Multicolor 3D-FISH and 3D image analysis of chromosomal positioning during the differentiation of normal human hematopoietic stem cells into derived myeloid precursors indicate that there are no drastic changes in chromosomal preferential localization within sub nuclear regions during this differentiation transition. This evidence is in accordance with results obtained from the comparison of gene expression profiles of HSCs and myeloid precursor, where a relative small number of chromosomal regions were identified as differentially expressed. Nevertheless, the small differences in chromosomal radial positioning between the two cell types are accompanied by different chromosome volumes that could be correlated to variations of chromatin compaction occurring during differentiation. Moreover, stem cells and myeloid precursor present a high difference in homologous distances, which could be associated with differential gene expression.

Overall, these results represent the first experimental evidence of the arrangement of chromosomal position in two steps of normal human myelopoiesis and constitute a working framework relevant to decipher the genomic mechanisms at the base of myeloid differentiation.

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