

## Future Prospects of 3D Human Chromosome Imaging by Serial Block Face Scanning Electron Microscopy

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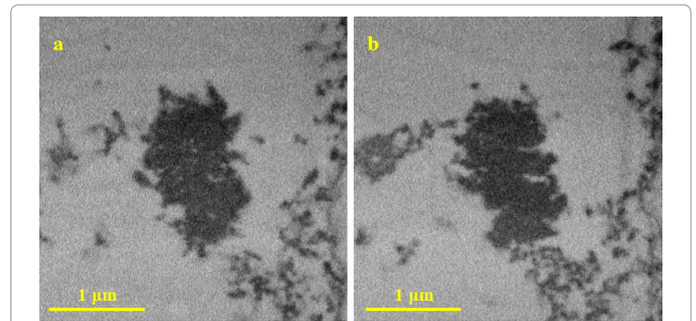
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### Commentary

The higher order structure of human chromosomes remains to be elucidated with the 30 nm mystery still remaining. For internal structure determination, transmission electron microscopy (TEM) cannot be used as the chromosomes are too thick (approx. 1.4 microns) and scanning electron microscopy (SEM) is a surface imaging technique. For this purpose, the three-dimensional (3D) serial block face scanning electron microscopy (SBFSEM) was used on imaging mitotic human chromosome for the first time [1].

The SBFSEM technique [2] uses a diamond knife to cut thin slices of samples embedded into resin after preparing the samples using standard electron microscopy preparation techniques. The serial images by the SBFSEM do not need to be aligned because the system has automated image acquisition [2]. No published protocols were available for chromosome sample preparation for SBFSEM but rather for tissue samples [3] including rat liver to investigate chromatin in interphase nuclei [4]. This study used multiple fixation steps with complex long-time sample preparation procedures [4]. The chromosome sample preparation procedure for SBFSEM by Yusuf et al. [1] is much simpler and shorter. This study used centrifugation after each preparation step causing sample loss therefore the study successfully optimised the sample preparation procedures by starting with large scale cell cultures, synchronization and pooling samples together to get a concentrated starting yield. Further improvements were made after cleaning the chromosome sample by filtration, staining using platinum blue (a DNA specific heavy metal stain); [5] and embedding in a small amount of resin. Full 3D was obtained using the images taken and the typical X shaped chromosome was obtained with no internal structure. Further sample preparation changes from polyamine to methanol acetic acid showed internal pores [1].

SBFSEM has been used down to 20 nm sections showing clear signs of internal structure [1]. Even though porous features or cavities were seen on the chromosome arms, the resolution was limited to 11 × 11 × 20 nm. Charging effects and radiation damage were believed to give the current limit of resolution. Despite the 11 nm nominal lateral resolution, no 30 nm structures were seen. New images with 8.3 × 8.3 nm pixel size show consistent staining in each slice (Figure 1a) and in slices further down in the series (Figure 1b) of the same chromosome (150 nm apart), also having porous information. Complementary methods such as focused ion beam scanning electron microscopy (FIBSEM) have provided 3D information on plant chromosomes (not human) due to sample preparation difficulties [6]. 3D X-ray coherent diffraction imaging (CDI) provided 120 nm resolution with little internal fine structure on a human chromosome [7]. Therefore there is no doubt that the SBFSEM method will be further used in the future for exploring higher order structure of the human genome, hopefully providing higher resolution as the techniques are improving,



**Figure 1:** SBFSEM of a single human mitotic chromosome prepared from a lymphoblastoid cell line (GM18507). 2D sections are stained with platinum blue. (a) shows a section after SBFSEM from a whole chromosome (b) is a section further down the series being 150 nm apart from (a). Pixel size is 8.3 × 8.3 nm.

for example employing new generation back scattered electron (BSE) detectors. Various sample preparation procedures need to be developed that would include decondensing the chromatin by removing divalent cations, elimination of hypotonic buffer and imaging chromosomes by directly slicing cells. Currently no cryogenic stage for the instrument is available therefore the samples have to be imaged at room temperature after embedding in resin, however freeze substitution [8] of chromosomes after high pressure freezing would be a positive way forward. This would reduce the damage rate by preventing the diffusion of free radicals. Cryo-FIB has been used for cell imaging [9] and is yet to be experimented on human chromosomes in a single cell.

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