Chapter Title	Visualizing the Spatial Relationship of the Genome with the Nuclear Envelope Using Fluorescence In Situ Hybridization			
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Abstract	Much of the genome is anchored chromatin binding proteins such nucle proteins. Even though there are glob assess what parts of the genome are it is also essential to be able to visua reveal their individual relationships This is executed by fluorescence in situ flattened nuclei (2D-FISH) or 3-dime combination with indirect immunoflu This chapter explains the protocols for indirect immunofluorescence and dis analysis. Due to the nuclear envelope p nucleoskeleton, we also describe ho	hip with the nuclear envelope in cells. at the nuclear periphery, tethered by ear lamins and other integral membrane al assays such as DAM-ID or ChIP to associated with the nuclear envelope, alize regions of the genome in order to with nuclear structures in single cells. u hybridization (FISH) in 2-dimensional ensionally preserved cells (3D-FISH) in corescence to reveal structural proteins. r 2D- and 3D-FISH in combination with scusses options for image capture and proteins being part of the non-extractable w to prepare DNA halos through salt o study genome behavior and association
Keywords (separated by " - ")	•	-FISH - 3D-FISH - Genome organization sitioning - Nuclear envelope - Nuclear

### Chapter 24

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#### Abstract

The genome has a special relationship with the nuclear envelope in cells. Much of the genome is anchored 8 at the nuclear periphery, tethered by chromatin binding proteins such nuclear lamins and other integral 9 membrane proteins. Even though there are global assays such as DAM-ID or ChIP to assess what parts of 10 the genome are associated with the nuclear envelope, it is also essential to be able to visualize regions of 11 the genome in order to reveal their individual relationships with nuclear structures in single cells. This is 12 executed by fluorescence in situ hybridization (FISH) in 2-dimensional flattened nuclei (2D-FISH) or 13 3-dimensionally preserved cells (3D-FISH) in combination with indirect immunofluorescence to reveal 14 structural proteins. This chapter explains the protocols for 2D- and 3D-FISH in combination with indirect 15 immunofluorescence and discusses options for image capture and analysis. Due to the nuclear envelope 16 proteins being part of the non-extractable nucleoskeleton, we also describe how to prepare DNA halos 17 through salt extraction and how they can be used to study genome behavior and association when com-18 bined with 2D-FISH. 19

Key wordsFluorescence in situ hybridization, 2D-FISH, 3D-FISH, Genome organization,20Chromosome territories, Gene positioning, Nuclear envelope, Nuclear lamins21

#### 1 Introduction

Chromosomes and genes are spatially organized within inter-23 phase nuclei, interacting with proteinaceous nuclear structures 24 which anchor and tether chromatin [1, 2]. Global methodolo-25 gies, such as DAM-ID, utilizing the ease of sequencing DNA, can 26 assess the spatial relationship of the whole genome with specific 27 nuclear structures such as nuclear lamins [3-5]. These analyses 28 bring useful information to the field, however as studies are data-29 rich and expensive, they might be excluded as a method of choice 30 if a large number of samples are required (e.g., steps in a differ-31 entiation pathway or large numbers of patient samples). 32

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Furthermore, using cells in mixed populations will result in averaged information and the genome, for example, can be positioned differently in proliferating versus non-proliferating cells, e.g., in a primary cell culture [6]. Real-time imaging in live cells [7, 8] can also be an informative approach to visualize genome interaction with nuclear structures; however, this may present challenges [9]. Therefore, despite global and real-time analyses, there is a necessity to analyze individual cells by fluorescence in situ hybridization (FISH) to reveal specific regions of interest in the genome, and analyze their spatial relationship to specific nuclear structures such as the nuclear envelope [10, 11].

FISH is a cytogenetic technique that utilizes labeled DNA or RNA probes to hybridize to specific DNA sequences on chromosomes, or to RNA [12, 13]. It is possible to delineate whole chromosomes, chromosome arms, chromosome bands, gene loci, specific regions of genes or RNA transcripts [14], and analyze their spatial relationship with other FISH signals or nuclear landmarks such as the geometric center of nuclei, nucleoli, or the nuclear envelope. The nuclear envelope can be revealed by indirect immunofluorescence using antibodies [15–17], but the FISH denaturation process can negatively affect nuclear envelope antigens, especially A-type lamins. The use of formamide for DNA denaturation can also destroy the signal from fluorescently tagged proteins but antibodies that recognize the tag itself can be used [18]. Indirect immunofluorescence can be done prior to the FISH, by fixing the antigen-antibody complexes in place with paraformaldehyde; in this scenario, the nuclear envelope can still be revealed with specific antibodies [18-22]. However, the simplest way of revealing the nuclear edge is by using a DNA stain and where it ends is where the edge of the nucleus is; this is however as long as nuclei have maintained their integrity and shape during the procedure [23–26]. Use of phase contrast microscopy or lipophilic dyes can be useful, in cases where the nuclear envelope is compromised, to reveal the nuclear membranes. Abnormal nuclear structures such as blebs can create issues when defining a nuclear edge since they could be defined as an extension of the nuclear envelope or as a separate entity [27].

For 2D-FISH, whereby nuclei are flattened, a standard epifluorescence microscope with a cooled charge coupled device camera will suffice for visualization and imaging of FISH signals. However, for 3D-FISH, microscopy requires optical sections through the z-axis, either using a confocal laser scanning microscope or other microscope systems with a motorized stage or piezo-driven objectives. The 3D datasets then need to be processed through deconvolution packages to remove out of focus fluorescence. There are new opportunities to visualize FISH signals within interphase nuclei using super-resolution microscopy such as 3D-Structural Illumination Microscopy (3D-SIM) [16] or Visualizing the Spatial Relationship of the Genome...

high-throughput flow imaging [28, 29]. The high power lenses 81 that can be added to imaging flow machines, combined with the 82 extended depth of field, enable software to be developed to per-83 form positional analyses of chromosomes and genes in millions of 84 individual interphase nuclei for one sample in 3D. Individual cells 85 can then be sub-categorized into for example proliferating or not, 86 differentiated or not, transgenic or not, by co-staining with spe-87 cific markers. A further advancement is in high-throughput / 88 high-resolution imaging, an approach called "Deep Imaging": 89 there, individual cells are imaged at high resolution by automated 90 software and FISH signals positioned by automated analysis soft-91 ware built into the system [30, 32]. 92

Analyzing the position of genes, sub-chromosomal regions or 93 whole chromosomes within interphase nuclei requires the capabil-94 ity to be able to perform measurements and approaches that place 95 FISH signals into specific regions of the nucleus. In 3D, this is 96 usually done using the reconstructed stack of optical sections or a 97 gallery of the 2D optical sections to measure to the nearest point 98 at the nuclear edge in any of the x, y, z axes, or by taking the mea-99 surement to the geometric center of the nucleus and expressing 100 this a percentage of the radius. 3D radial analysis packages have 101 been developed [17, 31, 32] to perform measurements between 102 nuclear landmarks and FISH signals, or between FISH signals [33, 103 34]. Freely available software to position genes and chromosomes 104 includes Smart 3D-FISH [35] and NEMO [36]. However, many 105 laboratories use 2D preparations to perform their spatial position-106 ing analyses since similar nuclear positions for genes and chromo-107 somes are found consistently in 3D, confirming that positional 108 analyses can be performed in 2D [37]. In order to position FISH 109 signals in flattened nuclei different groups have developed their 110 own positional analysis scripts similar to the original created by 111 Paul Perry, MRC Human Genetics Unit, Edinburgh [38], whereby 112 the nucleus is segmented into 5 shells of equal area and the inten-113 sity of the FISH signal and DNA signal from the DNA dye DAPI 114 are measured [38–40]. The DNA signal intensity is used to nor-115 malize the FISH signal data. Others do not perform any normal-116 ization for DNA content across the flattened nuclei but use specific 117 measurements to nuclear landmarks such as the nuclear edge and 118 the nuclear center [41]. With such 2D studies, it is imperative that 119 the data be comparable between samples since they do not give 120 absolute position but an averaged probabilistic position. 121

By using FISH and delineation of the nuclear envelope or 122 other nuclear structures, it is possible to determine which parts of 123 the genome are colocalized or spatially close to nuclear structures 124 such as the nuclear lamina, nuclear pores and other nuclear components. This chapter outlines methods for 2D- and 3D-FISH in 126 combination with indirect immunofluorescence, and discusses the 127



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value of 2D vs 3D analyses. We also include methods for the DNA halo extraction protocol combined with 2D-FISH, making it possible to gather data about which regions of the genome are embedded or attached to insoluble nuclear structures (Fig. 1).



**Fig. 1** The composite figure displays images from 2D-FISH protocol to visualize chromosome territories in panel A (*red* and *green*) and gene loci in panel B (*green*). The nucleus is counterstained with DAPI to reveal DNA. This allows the edge of the nucleus to be delineated easily allowing for erosion scripts (*panel A*) to define the edge of the nucleus, as is shown by the cartoon in *panel A. Panel B* displays gene loci FISH signals using a

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#### 2 Materials

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	Careful selection of probe sequence and type is required when ask- ing questions about which parts of the genome are associated with nuclear structures. Chromosome painting probes are commercially available and ready labeled containing <i>Cotl</i> DNA for suppression of repetitive sequences [42] ( <i>see</i> <b>Note 1</b> ). Templates of individual chromosomes, that have been either flow-sorted [43] by size or microdissected are available [44]. Labeling of original template is normally performed by degenerate oligo primer-PCR [45]. Specific centromere and telomere probes are also commercially available as are many gene probes. When performing gene positioning studies we use commercially available bacterial artificial chromosomes (BAC) containing the sequence of interest ( <i>see</i> <b>Note 2</b> ). The BAC DNA is released from the bacteria using a commercial kit for DNA extraction, following measurement of the amount of DNA gener- ated. A nick-translation reaction is performed to label the BAC DNA with either biotin or digoxigenin conjugated nucleotides, performed through commercially available kits. Other methods of labeling include random priming and end labeling through PCR methods. The method for labeling probes described here is the DOR	<ol> <li>133</li> <li>134</li> <li>135</li> <li>136</li> <li>137</li> <li>138</li> <li>139</li> <li>140</li> <li>141</li> <li>142</li> <li>143</li> <li>144</li> <li>145</li> <li>146</li> <li>147</li> <li>148</li> <li>149</li> <li>150</li> <li>151</li> </ol>
<ul> <li>2.1 Two-Dimensional Fluorescence In Situ Hybridization</li> <li>2.1.1 Cell Culture and Fixation</li> </ul>	<ol> <li>DOP-PCR method for whole chromosome painting probes.</li> <li>Specific cell medium supplemented with 10 % (v/v) fetal bovine serum (FBS) and 2 % (v/v) penicillin/streptomycin.</li> <li>Versene (phosphate buffered saline (137 mM NaCl, 2.7 mM, KCl, 10 mM Na<sub>2</sub>HPO<sub>4</sub>, 1.8 mM KH<sub>2</sub>PO<sub>4</sub>; pH 7.4) with 0.2 % (w/v) ethylenediaminetetraacetic acid; EDTA) kept refrigerated at 4 °C, pre-warmed to 37 °C.</li> </ol>	152 153 154 155 156 157 158

**Fig. 1** (continued) BAC probe labeled with biotin and fluorescently tagged streptavidin. One of the gene loci is localized at the nuclear edge, whereas the other maybe at the edge but it is not possible to tell with 2D-FISH and simple measurements. This is why the erosion scripts can be used with a large number of nuclei. Using 3D-FISH the positioning of a FISH signal in the *x*, *y* or *z* coordinates can be revealed by confocal laser scanning microscopy reconstruction or analysis of a gallery of 2D images as shown in the cartoon in *panel D*. Chromosome territories are drawn, that can be seen to touch the nuclear edge in the *x* and *y* planes but also the *z* plane as one of the territories in *red* is at the bottom of the nucleus. *Panel C* was kindly provided by Dr. Marion Cremer, Ludwig-Maximillian University, Munich; it shows the relationship of chromosome *X* territories (*green*) at the nuclear edge with lamin B1 (*red*) in great detail due to the use of super-resolution microscopy (3D-SIM) [16, 51]. *Panels E, F* and *G* display DNA halo images and how to delineate the residual nucleus and DNA halo (*panel E*). *Panel F* displays territories of chromosomes 17 and 18 (*red*) in proliferating primary fibroblasts (Ki67 in *green*) and in senescent fibroblasts (negative for Ki67). Chromosome 18 is seen at some distance from the residual nucleus in the DNA halo in both proliferating and senescent DNA halos. In *panel G* DNA halos are also displayed subjected to 2D-FISH with labeled BAC probes revealing gene loci signals (*red*), again Ki67 is revealed in *green*. Scale bar, 10  $\mu$ m



159 160		3. 0.25 % trypsin produced from a 2.5 % stock solution, which has been diluted (1:10 v/v) in Versene.
161 162		4. Fixative: methanol–acetic acid (3:1 v/v), made fresh and made ice-cold.
163		5. Hypotonic solution: 0.075 M KCl ( <i>see</i> Note 3).
164 165	2.1.2 Slide preparation and Denaturation	1. Glass microscope slides that can withstand high and low temperatures.
166 167		2. 20× sodium citrate saline (SSC, 300 mM sodium citrate hydrate, 3 M NaCl).
168 169		3. Denaturation solution: 70 % (v/v) formamide, 2× SSC, pH 7.0 ( <i>see</i> Note 4).
170		4. Water baths.
171 172		5. Ethanol series comprising of 70, 90, and 100 % $(v/v)$ high quality ethanol at room temperature.
173		6. Glass Coplin jars that can withstand heat.
174		7. Oven that can be set at 70 °C.
175	2.1.3 Degenerate Oligo	1. PCR machine.
176	Primer-PCR	2. $5 \times$ GC Buffer containing 2 mM MgCl <sub>2</sub> when at 1×.
177		<ol> <li>dACGTP mix made up in water at a concentration of 2 mM.</li> </ol>
178		4. dTTP at a concentration of 2 mM.
179 180		5. Forward and reverse DOP primers at a concentration of 20 μM (CCGACTCGAGNNNNNATGGG).
181		6. Taq DNA polymerase.
182		7. PCR grade water.
183		8. Template DNA—chromosome paint.
184		9. Biotin-16-dUTP or digoxigenin-11-dUTP.
185	<u> </u>	10. Agarose.
186		11. Agarose mini gel kit.
187		12. DNA ladder—containing bands at 500 bp and lower.
188		13. TAE buffer.
189	2.1.4 Probe Preparation	1. Labeled total human chromosome or BAC DNA probes.
190	and Hybridization	2. Cotl DNA.
191		3. Herring sperm DNA.
192		4. 3 M sodium acetate.
193		5. 4× SSC.
194		6. High quality ethanol.
195		7. Hybridization mixture: 50 % formamide, 10 % dextran sulfate,
196		2× SSC, and 1 % Tween 20.

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	8. Heating block.	197
	9. 70 % formamide and 2× SSC solution, pH 7.0.	198
	10. Humidified hybridization chamber.	199
	11. Rubber cement.	200
	12. Glass coverslips $(22 \times 22 \text{ mm})$ .	201
	13. Glass Coplin jars (see Notes 5 and 6).	202
	14. Bench top centrifuge.	203
2.1.5 Washing 2D FISH	1. Buffer A: 50 % (v/v) formamide, $2 \times$ SSC; pH 7.0.	204
and Visualization of Probe	2. Buffer B: 0.1× SSC; pH 7.0.	205
	3. Blocking solution: 4 % bovine serum albumin (BSA).	206
	4. Streptavidin conjugated to a fluorochrome.	207
	5. Antibody reacting with digoxigenin conjugated to a fluorochrome.	208 209
	6. $4 \times$ SSC solution containing 0.05 % Tween 20.	210
	7. Vectashield containing 6-diamidino-2-phenylindole (DAPI).	211
2.1.6 Ki67 Antigen	We have found that chromosome positioning changes according to	212
Marker for Proliferation in Primary Cells	whether cells are proliferating, quiescent or senescent $[6, 46-48]$ . It is possible to identify replicative senescent cells within a proliferating	213 214
in rinnary ocno	culture grown in high serum since they are anti-Ki67 negative. It is	214 215
	also possible to work with quiescent cells using Ki67 as a marker in	216
	a young culture containing fewer senescent cells. After 4 days, the Ki67 is no longer visible. It is also possible to observe three different	217
	stages of G1 by Ki67 patterns type Ia, type Ib, and type II [49].	218 219
	1. $1 \times PBS$ .	220
	2. $1 \times PBS$ containing 1 % new born calf serum.	221
6	3. Anti-Ki67 antigen antibody.	222
	4. Fluorochrome-conjugated secondary antibody raised against the appropriate species.	223 224
	5. Glass coverslips.	225
2.1.7 Microscopy and Analysis	1. Epifluorescence microscope equipped with a 100× lens and filter sets to permit the capture of at least fluorochromes emitting	226
and Analysis	photons in the blue, green, and red spectra.	227 228
	2. Cooled charged coupled device camera.	229
	3. Computer and program to run microscope and camera.	230
	4. Hardware and software to visualize captured images.	231
	5. Image analysis to perform measurements from FISH signal to	232
	nearest edge the nucleus or geometric center of the nucleus or	233
	erosion analyses as discussed in the Introduction.	234
	6. Programs for data handling, analysis, and presentation.	235



236 237 238 240 241 242 243 244 245 246 247 248 249 250 251 252 253	<ul> <li>2.2 Three- Dimensional Fluorescence In Situ Hybridization</li> <li>2.2.1 Cell Culture and Fixation</li> </ul>	<ol> <li>Cell medium of choice for specific cell type containing 10 % (v/v) fetal bovine serum (FBS) and 2 % (v/v) penicillin/ streptomycin.</li> <li>Versene (phosphate buffered saline (137 mM NaCl, 2.7 mM, KCl, 10 mM Na<sub>2</sub>HPO<sub>4</sub>, 1.8 mM KH<sub>2</sub>PO<sub>4</sub>; pH 7.4) with 0.2 % (w/v) ethylenediaminetetraacetic acid; EDTA) kept at 4 °C, and pre-warmed to 37 °C.</li> <li>0.25 % trypsin from a 2.5 % stock solution, which has been diluted (1:10 v/v) in Versene.</li> <li>Chambered tissue culture dishes to hold slides.</li> <li>Glass microscope slides that can withstand high temperatures.</li> <li>1× phosphate buffered saline.</li> <li>4 % paraformaldehyde (w/v) (<i>see</i> Note 7).</li> <li>Permeabilization solution: 0.5 % Triton X-100 (v/v) and 0.5 % saponin (w/v) in 1× PBS.</li> <li>20 % glycerol in 1× PBS solution.</li> <li>Liquid nitrogen.</li> <li>Glass Coplin jars.</li> </ol>
254	2.2.2 Probe Preparation and Hybridization	See Subheading 2.1.4 for probe preparation for 2D FISH.
255 256 257 258 259 260 261	2.2.3 Slide Preparation and Denaturation	<ol> <li>Denaturation buffer A: 70 % formamide, 2× SSC, pH 7.0.</li> <li>Denaturation buffer B: 50 % formamide, 2× SSC, pH 7.0.</li> <li>Glass coverslips (22×32 mm).</li> <li>0.1 N HCl.</li> <li>2× SSC.</li> <li>Coplin jars.</li> <li>Rubber cement.</li> </ol>
262 263	2.2.4 Washing and Visualization of Probe in 3D-FISH	See Subheading 2.1.5 for washing and probe visualization for 2D FISH.
264	2.2.5 Using Ki67 as a Proliferation Marker	See Subheading 2.1.6 for using Ki67 antibody.
265 266 267 268 269 270	2.2.6 Microscopy and Analysis	<ol> <li>Fluorescence microscope capable of acquiring optical Z-Stacks with high powered lens such as a confocal laser scanning micro- scope (with at least two lasers), Deltavision, 3D-simulation illumination microscope, 3D-STORM [50].</li> <li>Software for wide-field deconvolution, 3D reconstruction, measurements in 3D.</li> </ol>

2.3 FISH in Combination with Indirect Immunofluorescence Visualizing the Spatial Relationship of the Genome...

The best way to visualize the nuclear envelope is by indirect 271 immunofluorescence using antibodies to nuclear lamins—A-types 272 and B-types, nuclear pores or integral membrane proteins such as 273 emerin, LAP2β, SUN1, SUN 2, MAN1, or other nuclear envelope 274 transmembrane proteins [18] and even nuclear pore proteins. The 275 issue with combining FISH with visualization of proteinaceous 276 structures is that the fixation, and specifically the formamide, can 277 affect the ability of specific antibodies to bind to their antigen post-278 FISH. We have found this especially difficult for nuclear envelope 279 proteins such as A-type lamins, which for mapping associations 280 with genomes and the nuclear envelope are the presumed first 281 choice. It is not possible to just use an integrated GFP fused pro-282 tein since GFP is also affected by the FISH process and is no longer 283 visible post-FISH. 284

There are two ways to get around this problem: either use the 285 primary and fluorescently tagged secondary antibodies and fix 286 them prior to the FISH protocol, or use the primary antibody 287 prior to the FISH protocol, fix them in place with paraformalde-288 hyde and then carry out the FISH protocol, following the hapten 289 visualization step, or concomitant with it, use the fluorescently 290 labeled secondary antibody to reveal the primary antibody bound 291 to the antigen. As mentioned above, GFP fluorescence does not 292 survive the FISH denaturation process but it possible to use an 293 antibody to GFP to reveal the exogenous protein. 294

The 2D-FISH protocol could, given that it uses acetic acid, 295 alter structure of proteins. Therefore, it would be necessary to per-296 form a pilot experiment to determine if the antigen of choice at the 297 nuclear envelope survives fixation and the FISH denaturation. 298 With respect to analyzing genome interaction with the nuclear 299 envelope, it is more appropriate to use 3D preserved cells or nuclei. 300

1. Reagents for 2D or 3D FISH (see Subheadings 2.1 and 2.2). 301

- 2. Primary antibodies to specific nuclear envelope antigen, Ki67 302 or GFP. 303
- 3. Secondary antibodies reacting with species primary antibodies 304 are made in 4 % paraformaldehyde. 305
- 4.  $1 \times$  phosphate buffered saline. 306 5. Glass Coplin jars. 307
- 6. New born calf serum. 308 309
- 7. Mounting medium containing a DNA counterstain.
- 8. Humid incubation chamber.
- 9. Glass coverslips. 311

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2.4 DNA Halo Although this chapter concerns visualizing the interaction of the 312 genome with the nuclear envelope we have decided to include a Preparations 313 protocol to look at DNA halos which is a technique we have been 314



	using in our laboratory to give a further insight into the behavior
	and binding of the genome to the nucleoskeleton (also known as
	the nuclear matrix). Many of the nuclear envelope proteins and
	complexes at the nuclear edge are part of the nuclear matrix and indeed, a number of nuclear envelope proteins are also found deep
	within the nucleoplasm as part of the unextractable nucleoskele-
	ton, such as the nuclear lamins.
322 323	1. Temperature resistant glass microscope slides or poly-L-lysine coated slides.
324	2. Chambered tissue culture dishes.
325	3. Glass Coplin jars.
326	4. CSK buffer: 10 mM Pipes pH 7.8; 100 mM NaCl, 0.3 M
327	sucrose, 3 mM MgCl <sub>2</sub> , 0.5 % Triton X-100.
328	5. Phosphate buffered saline $10\times$ , $5\times$ , $2\times$ , and $1\times$ .
329	6. Extraction buffer: 2 M NaCl, 10 mM Pipes pH 6.8, 10 mM
330	EDTA, 0.1 % digitonin, (see Note 8) 0.05 mM spermine,
331	0.125 mM spermidine.
332	7. Ethanol series 10, 30, 70, and 95 % made up in water
333 334	8. See Subheadings 2.1.4 and 2.1.5 for reagents required for 2D-FISH
335	9. Epifluorescence microscope capable of taking 2D images.
336	10. CCD camera with hardware and software to control micro-
337	scope and camera.
338	11. Image analysis software.
339 <b>3 Methods</b>	
<ul> <li>340 3.1 Two-Dimensional</li> <li>341 Fluorescence In Situ</li> </ul>	1. Culture adherent cells for at least 1 week after recovering from liquid nitrogen storage ( <i>see</i> <b>Note 9</b> for suspension cells).
Hybridization	
	2. Remove medium and wash the cells once with Versene.
343 <i>3.1.1 Cell Culture</i> 344 and Fixation	3. Harvest cells by incubation 0.25 % trypsin in Versene; neutral- ize the trypsin by adding at least an equal volume of medium.
345 346	4. Centrifuge at $400 \times g$ for 5 min; remove the supernatant. Resuspend cells in a small amount of remaining medium.
347	5. Treat harvested cells with a hypotonic solution (0.075 M KCl)
348 349	for 15 min at room temperature to swell the cells ( <i>see</i> Note 10).
350 351	6. Centrifuge the cell suspension at $300 \times g$ for 5 min using a bench top centrifuge.
352 353	7. To fix the cells, remove the supernatant and resuspend the cell pellet in a small amount of buffer left to avoid cell clumping.

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		8. Add ice-cold methanol: acetic acid (3:1 v/v) drop-wise to the cells with gentle shaking.	354 355
		9. Incubate cells at 4 °C or on ice for at least 1 h, up to 18 h ( <i>see</i> Note 11).	356 357
		10. Centrifuge cells at $300 \times g$ at 4 °C for 5 min and repeat the fixing steps 8, 9 and 10 four to five times, but only leaving the cells in the fix solution on ice for 5–15 min until more than 90 % of the cells have lost their cytoplasm. Use phase contrast with X40 lens to view the slides prior to use. If there are still cells with cytoplasm, keep repeating the fixation steps until there are fewer than 10 %.	358 359 360 361 362 363 364
		11. Store fixed cell suspensions at $-20$ °C until required.	365
	3.1.2 Slide Preparation and Denaturation	1. Drop the cells onto humid or damp slides from a height, air dry the slides and then age the cells at 70 °C for 1 h or for 2 days at room temperature.	366 367 368
		2. Pass the aged slides through an ethanol row of 70, 90 and 100 % ethanol for 5 min in each solution, followed by air drying.	369 370 371
[AU1]		3. Pre-warm the slides at 70 °C for 5 min.	372
		4. Incubate in denaturating solution (70 % (v/v) formamide, 2× SSC, pH 7.0 at 70 °C for 2 min.	373 374
		5. Immediately plunge the slides into ice-cold 70 % ethanol for 5 min and then again pass through the ethanol row (90 and 100 % ethanol).	375 376 377
		6. Air-dry the slides and keep them warm on a warm plate until hybridization with the probe.	378 379
	3.1.3 DOP-PCR Protocol	1. To make 50 $\mu$ l of secondary template from the primary template, add together in a small Eppendorf tube that fits the PCR machine to be used: 10 $\mu$ l DOP-PCR buffer, 5 $\mu$ l 2 mM dACGTP, 5 $\mu$ l 2 mM dTTP, 5 $\mu$ l 20 $\mu$ M DOP primers, 23 $\mu$ l PCR grade water, 1 $\mu$ l of chromosome paint template and 1 U of <i>Taq</i> polymerase (to be added last).	380 381 382 383 384 385
		2. Place into PCR machine.	386
		<ul> <li>3. For amplification of primary product into secondary template use the following temperature and times: 1 cycle of 95 °C for 3 min followed by 98 °C for 20 s; 31 cycles of 62 °C for 1 min followed by 72 °C for 30 s; 1 cycle of 72 °C for 5 min; 4 °C hold.</li> </ul>	387 388 389 390 391
		4. To check the amplification and length of the secondary template, run 3 $\mu$ l of the PCR product in a 1 % agarose gel with markers that give you the ability to assess smears of PCR product between 200 and 500 bp.	392 393 394 395

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396 397 398		5.	Use the unlabeled secondary template DOP-PCR product as the next template DNA and add 5 $\mu$ l to a plastic PCR tube the correct size for your machine chambers.
399 400 401 402		6.	To the secondary template, add 10 ml of DOP-PCR buffer, 5 ml dACGTP (2 mM), 2 $\mu$ l dTTP (2 mM) and 10 $\mu$ l of biotin- 16-dUTP or digoxigenin-11-dUTP, 5 $\mu$ l of DOP-PCR primers, and 1 unit of <i>Taq</i> polymerase.
403 404 405 406 407		7.	For amplification of primary product into secondary template use the following temperature and times: 1 cycle of 95 °C for 3 min followed by 98 °C for 20 s; 34 cycles of 62 °C for 1 min followed by 72 °C for 30 s; 1 cycle of 72 °C for 5 min; 4 °C hold.
408 409		8.	Using a small aliquot check that there are PCR bands between 200 and 500 bp in an agarose gel.
410 411 412 413 414	3.1.4 Probe Denaturation and Hybridization	1.	Prepare the probe mixture by ethanol precipitation of the labeled chromosome paint (8 $\mu$ l per slide) with the addition of <i>Cot</i> -1 DNA (7 $\mu$ l per slide), herring sperm (3 $\mu$ l per slide), 1/20th volume of 3 M sodium acetate, and 2× volume of high quality ethanol (ice-cold).
415 416		2.	Incubate the mixture at $-80$ °C for at least 30 min; centrifuge at full speed in a standard microfuge for 30 min at 4 °C.
417 418		3.	Wash the DNA pellet with ice-cold 70 % ethanol and centrifuge at full speed in a standard microfuge for 15 min at 4 °C.
419		4.	Dry the pellets at 50 °C on a hot block (see Note 11).
420 421			Dissolve the probes in 12 $\mu$ l (per slide) of hybridization mix overnight at room temperature.
422 423 424		6.	Denature the probes at 75 °C for 5 min; allow the <i>Cot</i> -1 DNA to anneal to the repetitive sequences at 37 °C for at least 10 min but no longer than 2 h.
425 426		7.	Incubate slides at 70 °C in 70 % formamide, 2× SSC, pH 7.0 for 2 min.
427 428		8.	Slides are plunged into ice-cold 70 % ethanol and then placed in 90 and 100 % ethanol for 5 min each at room temperature.
429 430		9.	Apply 12 $\mu$ l of the probe to the slide and cover with pre- warmed 22 × 22 mm coverslip and seal with rubber cement.
431 432		10.	Allow the slides to hybridize with the probe in an humidified hybridization chamber at $37 ^{\circ}$ C for 2 days.
433 434 435	3.1.5 Washing 2D-FISH Slides		Post hybridization, gently remove the rubber cement and wash the slides three times for 5 min each in buffer A (50 % (v/v) formamide, $2 \times$ SSC, pH 7.0) preheated to 45 °C.
436 437 438		2.	Wash the slides in buffer B ( $0.1 \times$ SSC, pH 7.0) pre-warmed at 60 °C but used in the 45 °C water bath, three times for 5 min each before transferring to 4× SSC at room temperature.

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3. Incubate the slides with a 100 $\mu$ l blocking solution (4 % bovine serum albumin; BSA, w/v) for 10 min at room temperature.	439 440
4. Incubate the slides in 100 $\mu$ l of fluorescently tagged streptavi- din or anti-digoxigenin conjugated to a fluorochrome diluted in 1 % BSA for 1 h at room temperature.	441 442 443
5. Wash the slides in a 4× SSC solution containing 0.5 % Tween 20 in the dark at 42 °C with three changes of the solution every 5 min.	444 445 446
6. Mount the slides in mounting medium containing DAPI.	447
1. Once cells have been washed, hapten visualized and slides checked positive for the FISH signals, soak the coverslips in $1 \times PBS$ for 30 min at room temperature.	448 449 450
2. Dilute primary Ki67 antibody in $1 \times$ PBS containing 1 % new born calf serum and add 100 µl to the slide and cover with a $22 \times 40$ mm coverslip.	451 452 453
3. Incubate for 1 h at room temperature or 30 min at 37 °C or overnight at 4 °C.	454 455
4. Wash in $1 \times PBS$ for 15 min with 3 changes.	456
5. Place 100 µl of appropriate secondary antibody on the slides, cover with a 22×40 mm coverslip and incubate 1 h at room temperature, or 30 min at 37 °C, or at 4 °C for 4 h.	457 458 459
<ol> <li>Mount in suitable mounting medium containing DAPI (see Note 12).</li> </ol>	460 461
1. Examine the slides using 100× oil immersion lens.	462
2. Cells can be imaged according to their Ki67 (proliferative status).	463 464
3. Select nuclei randomly by following a rectangular scan pattern, and capture gray-scale images of these nuclei.	465 466
4. Capture at least 50–60 images per slide and convert into TIFF or PICT format.	467 468
5. Pass the images through an erosion analysis script (see Introduction). These scripts are devised to divide each cap- tured nuclei in five concentric shells of equal area, the first shell starting from the periphery of the nucleus going to the interior of the nucleus (fifth shell). The script measures the pixel inten- sity of DAPI and the chromosome probe in these five shells and puts the data obtained into a table.	469 470 471 472 473 474 475
6. Normalize the probe signal by dividing the percentage of the probe by the percentage of DAPI signal in each shell. Thus, the normalized proportion of probe is calculated in all five shells for at least 50 nuclei.	476 477 478 479
	<ol> <li>Incubate the slides with a 100 μl blocking solution (4 % bovine serum albumin; BSA, w/v) for 10 min at room temperature.</li> <li>Incubate the slides in 100 μl of fluorescently tagged streptavidin or anti-digoxigenin conjugated to a fluorochrome diluted in 1 % BSA for 1 h at room temperature.</li> <li>Wash the slides in a 4× SSC solution containing 0.5 % Tween 20 in the dark at 42 °C with three changes of the solution every 5 min.</li> <li>Mount the slides in mounting medium containing DAPI.</li> <li>Once cells have been washed, hapten visualized and slides checked positive for the FISH signals, soak the coverslips in 1× PBS for 30 min at room temperature.</li> <li>Dilute primary Ki67 antibody in 1× PBS containing 1 % new born calf serum and add 100 μl to the slide and cover with a 22×40 mm coverslip.</li> <li>Incubate for 1 h at room temperature or 30 min at 37 °C or overnight at 4 °C.</li> <li>Wash in 1× PBS for 15 min with 3 changes.</li> <li>Place 100 μl of appropriate secondary antibody on the slides, cover with a 22×40 mm coverslip and incubate 1 h at room temperature, or 30 min at 37 °C, or at 4 °C for 4 h.</li> <li>Mount in suitable mounting medium containing DAPI (<i>see</i> Note 12).</li> <li>Examine the slides using 100× oil immersion lens.</li> <li>Cells can be imaged according to their Ki67 (proliferative status).</li> <li>Select nuclei randomly by following a rectangular scan pattern, and capture gray-scale images of these nuclei.</li> <li>Capture at least 50–60 images per slide and convert into TIFF or PICT format.</li> <li>Pass the images through an erosion analysis script (see Introduction). These scripts are devised to divide each captured nuclei in five concentric shells of equal area, the first shell starting from the periphery of the nucleus going to the interior of the nucleus (fifth shell). The script measures the pixel intensity of DAPI and the chromosome probe in these five shells and puts the data obtained into a table.<!--</td--></li></ol>

480 481		7.	Plot histograms using these data, with standard error bars $(+/-SEM)$ .
482		8	Alternatively, make measurements to the nearest edge or geo-
483		0.	metric center of the nucleus to position genes/chromosomes.
484			These types of analyses can be meaningful in 2D especially
485			when studying peripherally located FISH signals.
486	3.2 Three-	1.	Culture adherent cells on sterile glass slides placed in a cham-
487	Dimensional		bered tissue culture dish for at least 2 days at 37 °C, 5 % $\rm CO_2$
488	Fluorescence In Situ		at a starting density of $1 \times 10^5$ cells per slide.
489	Hybridization	2.	Wash slides three times in $1 \times PBS$ .
490	(3D-FISH)	3.	Fix in 4 % paraformaldehyde (w/v) for 10 min at room
491	3.2.1 Cell Culture		temperature.
492	and Fixation		Wash slides three times in $1 \times PBS$ .
493		5.	Permeabilize cells with 0.5 $\%$ Triton X-100 (v/v) and 0.5 $\%$
494			saponin (w/v) in 1× PBS solution for 20 min at room
495			temperature.
496			Wash slides three times in $1 \times PBS$ .
497		7.	Incubate slides in a solution of 20 % glycerol in 1× PBS solu-
498			tion for at least 30 min at 4 °C
499		8.	Snap-freeze slides in liquid nitrogen for 15–30 s.
500		9.	Repeat step 8–9 another 4–5 times without the 30 min incuba-
501			tion in glycerol.
502		10.	Place slides in boxes and store at -80 °C until required.
503	3.2.2 Probe Preparation	See	Subheading 3.1.4
	and Hybridization		
504	3.2.3 Slide Preparation	1.	Thaw slides taken out of $-80$ °C storage at room temperature.
505	and Denaturation	2.	Repeat the freeze-thaw process in liquid nitrogen for another
506			4-5 times, soaking the slides in 20 % glycerol between each
507			freeze-thaw if this has not been done prior to freezing.
508		3.	Wash excess glycerol from the slides using three changes of $1\times$
509			PBS for 10 min each, followed by depurination in 0.1 N HCl
510			for 5 min at room temperature with shaking (see Note 13).
511		4.	Wash off excess acid with $2 \times SSC$ for 15 min with three changes
512			of the buffer and then incubate slides in 50 % formamide, $2 \times$
513			SSC, pH 7.0 solution overnight.
514		5.	Denature the slides by incubation in denaturation buffer A
515			(70 % formamide, 2× SSC, pH 7.0) pre-warmed at 73 °C for
516			precisely 3 min.
517		6.	Rapidly transfer the slides to denaturation buffer B (50 $\%$ for-
518			mamide, 2× SSC, pH 7.0) pre-warmed at 73 °C for 1 min.

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		During this time, add 10 $\mu$ l of previously denatured probe that has been allowed to partially re-anneal at 37 °C to a pre- warmed 22 × 22 mm coverslip.	519 520 521
		7. Once the slide is ready after denaturation, remove excess dena- turing solution with a tissue without disturbing the cells.	522 523
		8. Present the slide immediately to the probe on the coverslip.	524
		9. Seal the coverslip on the slide using rubber cement and leave to hybridize on the slide in a pre-warmed humidified chamber at 37 °C for 2 days.	525 526 527
	3.2.4 Washing 3-D FISH	Same as for 2D-FISH; see Subheading 3.1.5.	528
	3.2.5 Using Ki67 as a Marker for Proliferation	Same as for 2D-FISH; <i>see</i> Subheading 3.1.6.	529
	3.2.6 Microscopy and Analysis	1. Capture the images of nuclei using a microscope capable of capturing in three dimensions.	530 531
		2. Collect stacks of optical sections with an axial distance of 0.2 μm for high resolution from random nuclei ( <i>see</i> <b>Note 14</b> )	532 533
		<ol> <li>For confocal images you could obtain stacks of 8-bit gray-scale</li> <li>2D images with eight averages from each optical image.</li> </ol>	534 535
		4. Assess the positioning of chromosomes in relation to the nuclear edge by performing measurements using image analysis packages as described in the introduction, whereby the distance between the geometric center of each chromosome territory/gene signal and the nearest nuclear edge can be measured.	536 537 538 539 540
		5. Perform measurements for at least 20 nuclei for each sample.	541
		6. Plot frequency distribution curves with the distance between the center of chromosome territory and the nearest nuclear periphery on the x-axis and the frequency on the y-axis.	542 543 544
	3.3 3D-FISH	1. Place slides with fixed cells into a Coplin jar full of $1 \times PBS$ .	545
	Combined	2. Place slides in an humidified chamber	546
	with Indirect Immunofluorescence	3. Pipette 20–100 $\mu$ l of diluted primary antibody (in PBS 1 % new born calf serum or similar) and cover with a glass coverslip.	547 548
		4. Incubate at room temperature for 1 h, at 37 $^{\circ}$ C for 30 min or overnight at 4 $^{\circ}$ C.	549 550
		<ol> <li>Remove coverslips gently and place slides in glass Coplin jar in 1× PBS. Place on a rocker at a gentle speed changing buffer three times for a total wash time of 15 min.</li> </ol>	551 552 553
		6. Carefully wipe away PBS from the back of slides and replace in humidified chamber and repeat steps 3–5 but with the appropriate fluorochrome-conjugated secondary antibody. Incubate at room temperature for 1 h, 37 °C for 30 min or 4 °C for no longer than 4 h.	554 555 556 557 558

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559 560 561 562 563 564			After the final wash, incubate slides in 4 % paraformaldehyde made up in $1 \times PBS$ for 10 min at room temperature. Place slides in $2 \times SSC$ to prevent drying out before proceeding to the 3D FISH protocol ( <i>see</i> Note 15) for description of doing primary antibody staining before FISH and secondary antibody after FISH.
565 566 567	3.4 DNA Halo Preparations	1.	Either grow adherent cells directly onto glass microscope slides in culture or allow suspension cells to settle and attach to poly- L-lysine coated slides.
568		2.	Rinse slides twice in $1 \times PBS$ .
569		3.	Add CSK buffer for 15 min.
570		4.	Replace CSK buffer with extraction buffer and leave for 4 min.
571 572		5.	Rinse slides consecutively in $10\times$ , $5\times$ , $2\times$ , and $1\times$ PBS for 1 min each.
573 574		6.	Take slides through an ethanol series comprising 10, 30, 70, and 95 % ethanol.
575 576		7.	Air-dry slides and bake for 2 h at 70 °C. The slides are now ready to be subjected to 2D-FISH.
577 578	3.4.1 Measuring the Ratio of the Residual	1.	At 100× magnification, select at least 50 images of extracted cell nuclei per sample at random.
579 580	Nucleus to the Maximum Extent of the DNA Halo	2.	Capture of 8-bit gray-scale 2D images of each DNA halo in TIFF format using a high resolution digital camera.
581		3.	Open the image in NIH ImageJ software or similar.
582 583		4.	Configure ImageJ to analyze area (Analyse > Set Measurements, tick the area checkbox).
584 585		5.	Using the adjust threshold tool (Image>Adjust>Threshold), move the top slider to cover only the residual nucleus.
586 587		6.	Measure the area using analyze particles (Analyze > Analyse Particles) and record the measurement.
588 589 590		7.	Using the adjust threshold tool again, move the top slider to cover the entire area of the DNA halo, noting that the DNA halo will likely be very pale at its greatest extent.
591		8.	Repeat step 6 and open the next image.
592 593 594		9.	The ratio of the residual nucleus to the maximum extent of the DNA halo is calculated by dividing the total area of the DNA halo by the area of the residual nucleus.
595 596		10.	Plot histograms using these data and standard error bars representing $+/-$ SEM.



3.4.2 Analyzing Positioning of Whole Chromosomes Within a DNA Halo

- Visualizing the Spatial Relationship of the Genome...
- Using an image analysis program the images are cropped and 597 split into individual color channels, i.e., red, blue, and green. 598
   Save each channel separately. 599
- Identify and select the residual nucleus in an imaging software program to perform density threshold and binary functions to reveal not only the edge of the nucleus but also the center.
- 4. The distance from the nuclear center to each furthest chromosome territory edge (ctE) can then be determined.
   604
- 5. These distances are divided by the distance from the nuclear 605 center to each respective nuclear edge (NE). 606

#### 4 Notes

1. Sometimes commercial paints scrimp on the Cot-1 DNA and<br/>it may be necessary to add more Cot1 to avoid repetitive ele-<br/>ments in the genome from being hybridized to.608610

607

- The BAC gives you your gene of interest but be aware that 611 BACs may contain other sequence adjacent to the gene of 612 interest.
   613
- 3. This concentration of KCl in the hypotonic solution maybe
  too high for some cells; in some instances, this may need to be
  reduced to 0.05 M KCl (e.g., with snail cells).
- 4. Formamide is a teratogen and its use should be forbidden to 617 pregnant women. Use in a fumehood otherwise. 618
- 5. Use glass Coplin jars that can withstand high heat, i.e., >75 °C: 619 otherwise they crack. 620
- 6. To avoid large amounts of formamide in the denaturing step,
  it is possible to use a heating block and place the probe and
  sample together and co-denature them. Alternatively it is possible to purchase a programmable temperature controlled slide
  processing system that will perform the denaturing steps and
  the hybridization step as well.
- 7. The Cremer laboratory recommends 2 % paraformaldehyde as 627 it preserved 3D structure better when using super-resolution 628 microscopy (53).
- Digitonin is a dangerous reagent and requires a careful and 630 comprehensive risk assessment.
- 9. Suspension cells can be used for any FISH methodologies. 632
  They need to be placed on a glass microscope slide. This can be 633
  done by cytospinning but this disturbs three-dimensionality. 634
  Cells can also be placed on slides coated with poly-L-lysine and 635
  allowed to stick. These cells can then be subjected to 3D 636
  protocols. 637

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638 639	10. It is very important that cells are fully suspended and triturated so that they are separated and not clumped together.
640	11. The DNA must not be over-dried as it will not be possible to
641	resuspend it in the hybridization mix. It is best to keep an eye
642	on the probe as it dries and take it off the heating block when
643	the liquid has evaporated but the DNA looks "glassy."
644	12. At this stage it is also possible to improve the FISH process by
645	using an RNAseA treatment for 30 min. Only do this if you are
646	not detecting RNA obviously but also bear in mind that RNA
647	is part of the nuclear structure and a component of chromo-
648	somes themselves.
649	13. For protocols specifically for super-resolution microscopy [18].
650	14. The Ki67 antibody can disassociate from its antigen after a few
651	hours. 4 % paraformaldehyde (5 min at room temperature) can
652	be used to fix the antibody in place after washing off excess
653	antibody.
654	15. It is also possible to perform the primary antibody step first
055	prior to the FISH procedure and fix it into place with 4 % para-
655	
656	formaldehyde for 10 min at room temperature. The diluted
656	formaldehyde for 10 min at room temperature. The diluted
656 657	formaldehyde for 10 min at room temperature. The diluted secondary antibody should be mixed with the streptavidin or

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#### 668 References

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- Bourne G, Moir C, Bikkul U et al (2013)
  Interphase chromosome behavior in normal and diseased cells. In: Yurov Y (ed)
  Human interphase chromosomes: the biomedical aspects. Springer, New York,
  pp 9–33
- 675 2. Foster HA, Bridger JM (2005) The genome
  676 and the nucleus: a marriage made by evolu677 tion. Chromosoma 114:212–229
- 878 3. Németh A, Conesa A, Santoyo-Lopez J et al
  (2010) Initial genomics of the human nucleo1080 lus. PLoS Genet 6:1–11
- 4. Ikegami K, Egelhofer TA, Strome S, Lieb JD 681 (2010) Caenorhabditis elegans chromosome 682 arms are anchored to the nuclear membrane via discontinuous association with LEM-2. 684 Genome Biol 11:1–20 685
- 5. Guelen L, Pagie L, Brasset E et al (2008)
  Domain organization of human chromosomes revealed by mapping of nuclear lamina interactions. Nature 453:948–951
  688
- Bridger JM, Arican-Goktas HD, Foster HA 690 et al (2014) The non-random repositioning of 691 whole chromosomes and individual gene loci 692

in interphase nuclei and its relevance in disease, infection, aging, and cancer. Adv Exp
Med Biol 773:263–279

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- 696 7. Chubb JR, Boyle S, Perry P, Bickmore WA
  697 (2002) Chromatin motion is constrained by
  698 association with nuclear compartments in
  699 human cells. Curr Biol 12:439–445
- 8. Zink D, Cremer T, Saffrich R et al (1998)
  Structure and dynamics of human interphase
  chromosome territories in vivo. Hum Genet
  102:241–251
- Wallace LM, Moreo A, Clark KR, Harper SQ (2013) Dose-dependent toxicity of humanized renilla reniformis GFP (hrGFP) limits its utility as a reporter gene in mouse muscle. molecular therapy—nucleic acids 2:1–6.
- 10. Elcock LS, Bridger JM (2010) Exploring the
  relationship between interphase gene positioning, transcriptional regulation and the nuclear
  matrix. Biochem Soc Trans 38:263–267
- 11. Elcock LS, Bridger JM (2008) Exploring the
  effects of a dysfunctional nuclear matrix.
  Biochem Soc Trans 36:1378–1383
- 12. Bridger JM, Kalla C, Wodrich H et al (2005)
  Nuclear RNAs confined to a reticular compartment between chromosome territories. Exp
  Cell Res 302:180–193
- 13. Lampel S, Bridger JM, Zirbel R, Mathieu U,
  Lichter P (1997) Nuclear RNA accumulations
  contain released transcripts and exhibit specific
  distributions with respect to Sm antigen foci.
  DNA Cell Biol 16:1133–1142
- 14. Volpi EV, Bridger JM (2008) FISH glossary:
  an overview of the fluorescence in situ hybridization technique. Biotechniques 45:385–386
- 15. Marshall WF, Dernburg AF, Harmon B, Agard
  DA, Sedat JW (1996) Specific interactions of
  chromatin with the nuclear envelope: positional determination within the nucleus in
  Drosophila melanogaster. Mol Biol Cell 7:
  825–842
- 16. Markaki Y, Smeets D, Fiedler S et al (2012)
  The potential of 3D-FISH and super-resolution structured illumination microscopy for
  studies of 3D nuclear architecture. Bioessays
  34:412–426
- 17. Luo L, Gassman KL, Petell LM et al (2009)
  The nuclear periphery of embryonic stem cells
  is a transcriptional permissive and repressive
  compartment. J Cell Sci 122:3729–3737
- 18. Zuleger N, Boyle S, Kelly DA et al (2013)
  Specific nuclear envelope transmembrane proteins can promote the location of chromosomes to and from the nuclear periphery.
  Genome Biol 14:1–48
- 19. Lund E, Oldenburg AR, Delbarre E et al
   (2013) Lamin A/C-promoter interactions

specify chromatin state-dependent transcription outcomes. Genome Res 23:1580–1589 751

- Bridger JM, Herrmann H, Muenkel C, Lichter 752 P (1998) Identification of an interchromosomal compartment by polymerization of 754 nuclear-targeted vimentin. J Cell Sci 111: 755 1241–1253 756
- Bridger JM, Lichter P (1999) Analysis of 757 mammalian interphase chromosomes by FISH 758 and immunofluorescence. In: Bickmore W 759 (ed) Chromosome structural analysis: a practical approach. Oxford University Press, Oxford, 761 UK, pp 103–123 762
- Solovei I, Cremer M (2010) 3D-FISH on cultured cells combined with immunostaining. Methods Mol Biol 659:117–126
   763
   764
   765
- 23. Hatch E, Hetzer M (2014) Breaching the nuclear envelope in development and disease. J Cell Biol 205:133–141
   768
- 24. Bridger JM, Kill IR (2004) Aging of 769 Hutchinson–Gilford progeria syndrome fibroblasts is characterised by hyperproliferation and increased apoptosis. Exp Gerontol 39: 772 717–724 773
- 25. Goldman RD, Shumaker DK, Erdos MR et al (2004) Accumulation of mutant lamin A
  causes progressive changes in nuclear architecture in Hutchinson–Gilford progeria syndrome. Proc Natl Acad Sci U S A 101: 778
  8963–8968 779
- 26. Shimi T, Pfleghaar K, Kojima S et al (2008)
  The A- and B-type nuclear lamin networks:
  microdomains involved in chromatin organization and transcription. Genes Dev 22:
  3409–3421
  784
- 27. Bercht Pfleghaar K, Taimen P, Butin-Israeli V
  et al (2015) Gene-rich chromosomal regions are preferentially localized in the lamin B deficient nuclear blebs of atypical progeria cells. Nucleus 6:66–76
  785
- 28. Lalmansingh A, Arora K, DeMarco R et al (2013) High-throughput RNA FISH analysis by imaging flow cytometry reveals that pioneer factor Foxal reduces transcriptional stochasticity. PLoS One 8:1–12 794
- 29. Basiji DA, Ortyn WE, Liang L, Venkatachalam
  V, Morrissey P (2007) Cellular image analysis and imaging by flow cytometry. Clin Lab Med
  27:653–670
  798
- 30. Roukos V, Pegoraro G, Voss TC, Misteli T 799 (2015) Cell cycle staging of individual cells by fluorescence microscopy. Nat Protoc 801 10:334–348 802
- 31. Shiels C, Adams NM, Islam SA, Stephens DA, Freemont PS (2007) Quantitative analysis of cell nucleus organisation. PLoS Comput Biol 3(7), e138
  803
  804
  805
  806

- 807 32. Bewersdorf J, Bennett BT, Knight KL (2006)
  808 H2AX chromatin structures and their response
  809 to DNA damage revealed by 4Pi microscopy.
  810 Proceedings of the National Academy of
  811 Sciences of the United States of America
  812 103:18137–18142
- 813 33. Jost K, Haase S, Smeets D et al (2011)
  814 3D-Image analysis platform monitoring relocation of pluripotency genes during reprogramming. Nucleic Acids Res 39:1–8
- 817 34. Bolzer A, Kreth G, Solovei I et al (2005)
  818 Three-dimensional maps of all chromosomes
  819 in human male fibroblast nuclei and prometa820 phase rosettes. PLoS Biol 3:826–842
- 35. Gué M, Messaoudi C, Sun JS, Boudier T
  (2005) Smart 3D-FISH: automation of distance analysis in nuclei of interphase cells by
  image processing. Cytometry A 67:18–26
- 36. Iannuccelli E, Mompart F, Gellin J, LahbibMansais Y, Yerle M, Boudier T (2010) NEMO:
  a tool for analyzing gene and chromosome territory distributions from 3D-FISH experiments. Bioinformatics 26:696–697
- 37. Foster HA, Griffin DK, Bridger JM (2012)
  Interphase chromosome positioning in in vitro
  porcine cells and ex vivo porcine tissues. BMC
  Cell Biol 15:13–30
- 38. Croft JA, Bridger JM, Boyle S et al (1999)
  Differences in the localization and morphology of chromosomes in the human nucleus.
  J Cell Biol 145:1119–1131
- 838 39. Mehta IS, Kulashreshtha M, Chakraborty S,
  839 Kolthur-Seetharam U, Rao BJ (2013)
  840 Chromosome territories reposition during DNA
  841 damage-repair response. Genome Biol 14:1–15
- 842 40. Skinner BM, Robertson LB, Tempest HG et al
  843 (2009) Comparative genomics in chicken and
  844 Pekin duck using FISH mapping and microar845 ray analysis. BMC Genomics 10:1–11
- 846 41. Federico C, Cantarella CD, Di Mare P, Tosi S,
  847 Saccone S (2008) The radial arrangement of the
  848 human chromosome 7 in the lymphocyte cell
  849 nucleus is associated with chromosomal band
  850 gene density. Chromosoma 117:399–410

- 42. Lichter P, Cremer T, Borden J, Manuelidis L,
  Ward DC (1988) Delineation of individual human chromosomes in metaphase and interphase cells by in situ suppression hybridization using recombinant DNA libraries. Hum Genet 80:224–234
  854 856
- 43. Harris P, Boyd E, Ferguson-Smith MA (1985)
  Optimising human chromosome separation for the production of chromosome-specific DNA libraries by flow sorting. Hum Genet 70:59–65
  859
- 44. Meltzer P, Bittner M (2001) Chromosome 862 microdissection. Curr Protoc Hum Genet. 863 Chapter 4:Unit4.8 864
- 45. Telenius H, Carter NP, Bebb CE et al (1992)
  Degenerate oligonucleotide-primed PCR:
  general amplification of target DNA by a single
  degenerate primer. Genomics 13:718–725
  868
- 46. Bridger JM, Boyle S, Kill IR, Bickmore WA (2000) Re-modelling of nuclear architecture in quiescent and senescent human fibroblasts. Curr Biol 10:149–152 872
- 47. Mehta IS, Figgitt M, Clements CS, Kill IR, Bridger JM (2007) Alterations to nuclear architecture and genome behavior in senescent cells. Ann N Y Acad Sci 1100:250–263
  876
- 48. Mehta IS, Amira M, Harvey AJ, Bridger JM
  (2010) Rapid chromosome territory relocation by nuclear motor activity in response to serum removal in primary human fibroblasts. Genome Biol 11:1–23
  880
- 49. Bridger JM, Kill IR, Lichter P (1998) 882 Association of pKi-67 with satellite DNA of 883 the human genome in early G1 cells. 884 Chromosome Res 6:13–24 885
- 50. Schermelleh L, Heintzmann R, Leonhardt 886 H (2010) A guide to super-resolution fluorescence microscopy. J Cell Biol 190: 888 165–175 889
- 51. Markaki Y, Smeets D, Cremer M, Schermelleh 890
  L (2013) Fluorescence in situ hybridization applications for super-resolution 3D structured illumination microscopy. Methods Mol Biol 950:43–64

Author's Proof



# Author Query

Chapter No.: 24 0002647412

Query	Details Required	Author's Response
AU1	Closing parenthesis is missing in the sentence starts with "Incubate in denaturating solution (70 " Please check and clarify.	

horestear