# Chapter 11

## Microscopic Analysis of Chromatin Localization and Dynamics in *C. elegans*

## **Christian Lanctôt and Peter Meister**

### Abstract

During development, the genome undergoes drastic reorganization within the nuclear space. To determine 6 tridimensional genome folding, genome-wide techniques (damID/Hi-C) can be applied using cell popu-7 lations, but these have to be calibrated using microscopy and single-cell analysis of gene positioning. 8 Moreover, the dynamic behavior of chromatin has to be assessed on living samples. Combining fast stereo-9 typic development with easy genetics and microscopy, the nematode C. elegans has become a model of choice 10 in recent years to study changes in nuclear organization during cell fate acquisition. Here we present two 11 complementary techniques to evaluate nuclear positioning of genes either by fluorescence in situ hybrid-12 ization in fixed samples or in living worm embryos using the GFP-lacI/lacO chromatin-tagging system. 13

Key wordsNuclear organization, C. elegans, Fluorescence in situ hybridization, Chromatin tagging,14GFP-lacI/lacO, Microscopy15

### 1 Introduction

Nuclear architecture has been studied in a wide range of models, 17 including lower eukaryotes such as S. cerevisiae [1] and inverte-18 brates such as D. melanogaster. In recent years, Caenorhabditis ele-19 gans has also been used, in particular to investigate the relationships 20 between nuclear architecture and cellular differentiation [2-5]. 21 Genome folding has been analyzed either in fixed worms and 22 embryos using fluorescence in situ hybridization (FISH) or in live 23 samples with the *lacO*/GFP-lacI system [2, 3, 5, 6]. Both approaches 24 are highly complementary as each overcomes the limits of the other. 25 The *lacO*/GFP-lacI system has the advantage of allowing the in vivo 26 observation of chromatin position and dynamics. However, the 27 creation of tagged chromatin loci is somehow tedious, and no 28 more than a few loci can be observed simultaneously. In contrast, 29 the main advantage of 3D FISH resides in the ability to detect mul-30 tiple genomic segments in the same nucleus and, in combination 31 with immunolabeling, to relate their positioning to various nuclear 32

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33	compartments. Here we present recently developed methods and
34	reagents for both techniques and discuss the caveats, advantages,
35	and problems associated with them.

36	2	Materials	
37 38	2.1	3D FISH	10× NT buffer: 0.5 M Tris–HCl pH 7.5, 50 mM MgCl <sub>2</sub> , 0.5 mg/ mL BSA. Make 0.1 mL aliquots and store at -20 °C.
39 40 41			100 mM $\beta$ -mercaptoethanol: 3 $\mu$ L of 14.3 M $\beta$ -mercaptoethanol in 0.4 mL of deionized water. Make 0.1 mL aliquots and store at -20 °C.
42 43 44			10× dNTP mix ( <i>see</i> <b>Note 1</b> ): 0.5 mM dATP, 0.5 mM dGTP, 0.5 mM dCTP, 0.2 mM dTTP. Make 20 μL aliquots and store at -20 °C.
45			1 mM labeled dUTP (see Note 2).
46			2 U/µL DNAse I.
47			10 U/µL <i>E. coli</i> DNA polymerase I.
48			0.5 M EDTA.
49			16 °C water bath.
50			PCR primers, forward and reverse, 25 $\mu$ M each.
51			2 mM dAGC: 2 mM dATP, 2 mM dGTP, 2 mM dCTP.
52			1.5 mM dTTP.
53			5 U/μL Taq DNA polymerase.
54			$10 \times$ Taq buffer with MgCl <sub>2</sub> .
55			PCR cycler.
56			1.2 % (w/v) agarose in electrophoresis buffer (e.g., TAE $1\times$ ).
57			95 % EtOH, 70 % EtOH (cold).
58			3 M sodium acetate pH 5.2.
59			100 % methanol (–20 °C, in a Coplin jar).
60			$1 \times PBS.$
61			Rubber cement glue.
62			Deionized formamide.
63			Coverslips: $18 \times 18$ mm #1 thickness; $15 \times 15$ mm #1 thickness;
64			$22 \times 22 \text{ mm #0 thickness.}$
65			Polylysine-treated microscope slides
66			Metal block in dry ice, -80 °C.
67			4 % formaldehyde in $1 \times PBS$ .
68			10 mg/mL yeast tRNA stock.
69			$20 \times$ saline sodium citrate (SSC), $2 \times$ SSC, $0.2 \times$ SSC.

Author's Proof		
	Analyzing Nuclear Organization in Nematodes	
	50 μg/mL RNAse A in 2× SSC (freshly diluted from a 10 mg/mL stock).	70 71
	2×SSC/50 % formamide.	72
	Primary antibodies against hapten and secondary antibodies (see Note 3).	73 74
	0.5 % Triton X-100 (v/v) in $1 \times PBS$ (freshly made).	75
	0.1 N and 0.01 N HCl.	76
	$2 \times$ hybridization buffer: 20 % (w/v) dextran sulfate in $4 \times$ SSC. Make 0.5 mL aliquots and store at -20 °C.	77 78
	4× SSCT: 80 μL of Tween-20 (cut pipette tip) in 400 mL of 4× SSC.	79 80
	4× SSCT-BSA: 2 g of bovine serum albumin (fraction V or purer) in 50 mL of 4× SSCT.	81 82
	l μg/mL DAPI in 2×SSC.	83
	Vectashield mounting medium (Vector Labs).	84
2.2 Live Chromatin	Microscope slides.	85
Imaging	Coverslips 20×20 mm, #1.	86
	Agarose.	87
	Laboratory tape.	88
	M9 buffer (KH <sub>2</sub> PO <sub>4</sub> 3 g/L, Na <sub>2</sub> HPO <sub>4</sub> 6 g/L, NaCl 5 g/L, MgSO <sub>4</sub> 1 mM).	89 90
	Fine forceps.	91
	Scalpel blade.	92
	Mouth pipette (see Note 12).	93
	Hourglass.	94
	Molecular biology reagents.	95
	Gateway® BP Clonase® II mix.	96
	Gateway® LR Clonase® II mix.	97
	Competent DH5α.	98
	pDONR221 lacO/Cb unc-119 middle vector.	99
	pDONRP4P1R.	100
	pDONRP2RP3.	101
	MosSCI plasmids ( <i>see</i> https://sites.google.com/site/ jorgensenmossci/mossci-reagents) [7–9].	102 103
	C. elegans strains:	104
	GW396/AV696 with fluorescent lacI expression: GW396 [ <i>baf-1p</i> driven somatic expression], AV696 [ <i>pie-1p</i> -driven germline and early embryo expression]	105 106
	HT1593 $\mu mc-110$ mutant	107
	1111 <i>370 WW 117</i> mulant.	108

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Methods

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109	Mos insertion strain at locus of interest.
110	Imaging device:
111	Laser scanning confocal microscope/spinning disk confocal
112	microscope.

3D FISH The success of 3D FISH experiments depends crucially on the 3.1 114 labeling of the DNA probe, i.e., the probe should be labeled uni-115 formly. Probes should be highly specific to the target sequence and 116 small in size to allow for efficient penetration in the sample. In this 117 chapter, we describe the two methods of choice for probe synthesis, 118 namely, nick translation and PCR labeling. Equally important are 119 the pretreatment of the sample and the detection of labeled probes. 120

121 3.1.1 Probe Synthesis The choice and synthesis of a target-specific probe is arguably the most crucial step in any 3D FISH experiment. In the case of single-122 copy targets, the probe should be of sufficient length to give a clear 123 signal. For probes against the C. elegans genome, we routinely use 124 fosmids that harbor 30-40 kb of sequence around the chosen 125 genomic target. Individual fosmid clones are identified in WormBase 126 (http://www.wormbase.org) and ordered from Source BioScience 127 (http://www.lifesciences.sourcebioscience.com/clone-products/ 128 genomic-dna-clones/c-elegans-fosmid-library-.aspx). Fosmid DNA 129 is labeled by nick translation. In the case of arrayed sequences (e.g., 130 rDNA), much shorter genomic fragments (0.3–0.5 kb) can serve as 131 template for probe synthesis, which is most conveniently done by 132 PCR on genomic DNA. Both methods rely on the incorporation of 133 a modified deoxynucleotide during DNA synthesis to generate 134 labeled probes. The label is either a hapten that can be immunode-135 tected after hybridization (digoxygenin [DIG] and dinitrophenol 136 [DNP]) or a fluorophore. In our hands, biotinylated probes give 137 unacceptably high background in C. elegans embryos, and, for this 138 reason, we do not recommend using them. 139

140Nick TranslationAs the name implies, the nick translation technique relies on141DNAse I to introduce single-strand breaks in the template DNA142and on the combined 5'-3' exonuclease and polymerase activity of143DNA polymerase I to introduce labeled nucleotides starting from144these pseudo priming sites. The purity of the starting material, the145DNAse activity, and the incubation temperature are the most criti-146cal parameters:

1. Prepare the following mix, in this order.

### Analyzing Nuclear Organization in Nematodes

DNA	l μg	t1.1
10× NT buffer	5 μL	t1.2
100 mM β-mercaptoethanol	5 μL	t1.3
10× dNTP mix	5 μL	t1.4
1 mM labeled dUTP	$1 \ \mu L \ (see \ Note \ 4), \ final \ concentration \ (20 \ \mu M)$	t1.5 t1.6
ddH <sub>2</sub> O	to 48.5 µL	t1.7
Mix by pipetting up and down. Put tube o	on ice. Add the following	t1.8
DNAse I (diluted 1:15 in $ddH_2O$ )	$1 \; \mu L \; (\textit{see Note 5})$	t1.9
10 U/µL E. coli DNA polymerase I	0.5 µL	t1.10 148

- Mix well by pipetting up and down after the addition of 149 enzymes. Do not make air bubbles. Incubate for 90 min at 150 16 °C.
- 3. Transfer the reaction on ice and analyze 1/10 of the reaction 152 (5 μL) on a 1.2 % agarose gel (*see* Note 6). Include 1 kb and 153 100 bp ladders.
- 4. If the probe size is satisfactory (smear between 300 and 155 600 bp), proceed to the next step. If not, add 1 μL of diluted 156 DNAse I to the reaction, return the tube at 16 °C, and incubate further for 30 min at 16 °C. Repeat steps 3 and 4.
- 5. Add 2  $\mu$ L of 0.5 M EDTA to stop the reaction. Mix well.
- 6. Optional. Clean the labeled probe using ion-exchange mini 160 spin columns (e.g., Qiagen or Zymo Research). Elute in 50  $\mu$ L 161 (see Note 7). 162

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7. Store the labeled probe at -20 °C.

### 1. Prepare the following PCR reaction:

Template	200 ng of genomic DNA	t2.1
Forward primer	1 μL	t2.2
Reverse primer	1 μL	t2.3
2 mM dAGC	5 μL	t2.4
1.5 mM dT	$4~\mu L~(\text{final concentration}~120~\mu M)$	t2.5
1 mM labeled dUTP	$3~\mu L~(final~concentration~60~\mu M)$	t2.6
10× Taq buffer	5 μL	t2.7
ddH <sub>2</sub> O	to 49.5 μL	t2.8 165

 Program the following PCR reaction: [94 °C, 3']; [94 °C, 30"; 166 melting temperature, 30"; 72 °C 45"] 25×; [72 °C, 10'] 167

PCR Labeling

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168 169 170		3. Heat the PCR cycler to mix for 1 min at this to polymerase (2.5 U). Mi	$^{\circ}$ 94 °C. Incubate the probe synthesis emperature. Add 0.5 $\mu$ L of Taq DNA ix by pipetting.
171		4. Run the PCR program.	
172		5. Analyze $1/20$ of the PCF	R product $(2.5 \ \mu L)$ on a 1.2 % agarose gel.
173 174 175		6. <i>Optional</i> . Clean the lal spin columns (e.g., Qiag (see Note 7).	beled probe using ion-exchange mini gen or Zymo Research). Elute in 50 $\mu$ L
176 177		7. Measure the probe cond at –20 °C.	centration by spectrophotometry. Store
178	3.1.2 3D FISH	A few points should be me	ntioned before detailing our 3D FISH
179		protocol. First, since the go	al of the technique is to determine the
180		spatial localization of genon	nic segments, it is necessary to use fixa-
181		nucleus i e to limit as mu	in preserve the native structure of the
182		agents. In the following pro	tocol a brief incubation in cold metha-
184		nol is used to wet embryo	s after freeze cracking of the eggshell
185		before fixing them in cold for	ormaldehyde Second we have found it
186		important to perform the h	vbridization in homemade glass cham-
187		bers, so as not to compress	unduly the relatively thick C. elegans
188		samples ( $\sim 20-30 \mu m$ ). Thir	d, as mentioned in Subheading 1, 3D
189		FISH can be combined with	h the immunolabeling of cellular com-
190		ponents. To do so, we pe	rform the incubation with antibodies
191		after the Triton X-100 per	meabilization step, use a Cy3-labeled
192		secondary antibody, and fix	the immunocomplexes with formalde-
193		hyde before proceeding wit	h the FISH protocol. Finally, it should
194		be noted that unlike for hy	bridization to mammalian DNA, it is
195		not necessary to quench the	repetitive sequences when performing
196		FISH on C. elegans DNA	because these make up a much lower
197		proportion of the genome	in this organism (the most abundant
198		repeat element in C. elegan	us, the 439 bp long CE000087, covers
199	Drenevation	only 1.32 % of the genome	[10]).
200	of the Probe Mix	1. Mux the following in an	Eppendori lube, in this order.
+3 1		NT probe 1	vul (typically corresponding
t3.1		NI probe 1	to 400 ng of probe)
t3.3 t3.4		NT probe 2	$x\mu L$ (typically corresponding to 400 ng of probe)
t3.5			
t3.6		10 mg/mL tRNA	$1 \ \mu L \ (see \ Note \ 8)$
t3.7		Deionized water	to 100 μL
t3.8		3 M NaOAC pH 5.2	10 µL
t3 9		95 % EtOH (cold)	400 µL

### Author's Proof Analyzing Nuclear Organization in Nematodes 2. Vortex vigorously. Incubate at -20 °C for at least 2 h. 3. Centrifuge at $13,000 \times g$ for 20 min at 4 °C. 4. Decant supernatant. Wash pellet with 0.5 mL of cold 70 % EtOH. Air-dry on the bench for 2–3 min. 5. Add 20 $\mu$ L of deionized formamide to the pellet (the final probe concentration is 20 ng/ $\mu$ L each). Incubate at 37 °C for 20 min. Pipette up and down to resuspend pellet. Fixation and Pretreatment 1. Isolate embryos according to standard protocols, e.g., by the "bleaching" method. of Embryos 2. Deposit 12 µL of concentrated embryos in the center of a polylysine-treated slide. Cover gently with an $18 \times 18$ mm coverslip. 3. Working at low magnification under the stereomicroscope, adsorb liquid using a filter paper until the embryos are slightly compressed between the slide and the coverslip. 4. Place the slide on a metal block in dry ice. Incubate at -80 °C for at least 60 min. 5. Using a razor blade, pop up the coverslip (i.e., "cracking" the eggshell). Proceed immediately to next step. 6. Immediately fix embryos in cold 100 % methanol, 2 min at -20 °C. 7. Rinse slide(s) 1 min in $1 \times PBS$ at 4 °C (see Note 9). 8. Transfer the slide(s) to precooled (4 °C) 4 % formaldehyde in $1 \times PBS$ . Fix for 10 min at room temperature. 9. Wash slides in $1 \times PBS$ , two times for 2 min at room temperature. 10. Incubate for 5 min in 0.5 % Triton X-100 in $1 \times PBS$ . 11. Wash two times for 2 min in $1 \times PBS$ . 12. Rinse once in 0.01 N HCl. 13. Incubate for 2 min in 0.1 N HCl at room temperature. 14. Wash once with $1 \times PBS$ , 3 min at room temperature. Wash once with $2 \times$ SSC, 3 min at room temperature. 15. Treat with 50 $\mu$ g/mL RNAse A in 2× SSC, 45 min at 37 °C. Perform this step by overlaying the sample with 0.5 mL of the RNAse solution and incubating in a humidified chamber. 16. Wash once with $2 \times SSC$ , 2 min at room temperature. 17. Incubate in $2 \times SSC/50$ % formamide for at least 2 h at room temperature. 1. Dilute the probe to a concentration of 2–5 ng/ $\mu$ L in 100 % Hybridization and Post-hybridization Washes deionized formamide. Add an equal volume of 2× hybridization buffer. 25 $\mu$ L of probe solution is needed per sample (*see* Note 10).

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**Fig. 1** Making homemade hybridization chambers. (a) Using a diamond pen, cut a  $\sim$ 1 mm strip from the side of a 22 mm  $\times$  22 mm coverslip (*thickness #0*). (b) Apply nail polish on one side of the glass strip. (c) Immediately glue the glass strip to the side of a 15 mm  $\times$  15 mm coverslip (thickness #1). (d) Leave to dry and gently break off the protruding pieces of glass

243 244	2. Mix well by pipetting and vortexing. Spin 20 s in a tabletop centrifuge. Keep the probe at room temperature.
245 246 247	3. Take a slide out of the 2× SSC/50 % formamide solution. Gently wipe off excess liquid on either side of the sample using a soft tissue.
248 249 250	4. Prepare a glass chamber: glue with nail polish two small strips of glass of thickness #0 (width of ~1 mm) to the opposite sides of a 15 × 15mm coverslip ( <i>see</i> Fig. 1).
251	5. Pipette 25 $\mu$ L of probe solution on a glass chamber.
252 253 254	6. Using forceps, gently deposit the chamber on the slide, probe facing towards the sample. Let the probe solution spread to the edges of the chamber.
255 256	7. Seal the chamber with rubber cement glue. Let dry completely at room temperature.
257	8. Repeat <b>steps 3</b> –7 for each slide.
258	9. Pre-hybridize the slides overnight at 37 °C overnight.
259 260	10. Denature the probe and target DNA simultaneously by placing the slide for 5 min on a heating block set at 76 °C.
261	11. Incubate at 37 °C for 2–3 days.
262 263	<ol> <li>Fill four Coplin jars with 2× SSC. Preheat three of those at 37 °C.</li> <li>Fill two other Coplin jars with 0.2× SSC and preheat at 55 °C.</li> </ol>
264 265	<ol> <li>Gently remove the rubber cement glue around the chamber.</li> <li>Do not remove chamber.</li> </ol>
266 267 268	14. Place the slide(s) in a Coplin jar filled with 2× SSC at room temperature. The chamber should fall off. If it does not, pull it gently alongside the slide.

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15. Wash for $3 \times 5$ min in $2 \times 5$ SC at $37$ °C.	269
16. Wash for $2 \times 5$ min in 0.2× SSC at 55 °C.	270
1. If only fluorophore-labeled probes were used, go directly to step 17.	271 272
<ol> <li>Set aside a 1 mL aliquot of 4× SSCT-BSA for the dilution of antibodies.</li> </ol>	273 274
3. Transfer the slide to 4× SSCT-BSA. Incubate for 20 min at room temperature.	275 276
<ol> <li>Dilute the primary antibodies in 4× SSCT-BSA. Use only those antibodies that are needed (<i>see</i> Note 3).</li> </ol>	277 278
5. Remove the slide from the blocking solution. Gently wipe off excess liquid on either side of the sample.	279 280
6. Pipette 60 $\mu$ L of primary antibody solution close to the sample. Overlay with an $18 \times 18$ mm coverslip and make sure that the antibody solution covers the sample.	281 282 283
7. Incubate for 2 h at room temperature in a humid chamber, protected from light.	284 285
8. Place the slide in a Coplin jar filled with 4× SSCT at room temperature.	286 287
9. Wash for $3 \times 5$ min in $4 \times$ SSCT at room temperature.	288
10. Dilute the secondary antibodies in 4× SSCT-BSA. Use only those antibodies that are needed.	289 290
11. Centrifuge the secondary antibody solution for 2 min at $13,000 \times g$ . Transfer the supernatant to another microtube.	291 292
12. Remove a slide from the washing solution. Gently wipe off excess liquid on either side of the sample.	293 294
13. Pipette 60 $\mu$ L of secondary antibody solution close to the sample. Overlay with an 18 × 18 mm and make sure that the antibody solution covers the sample.	295 296 297
14. Incubate for 1 h at room temperature in a humid chamber, protected from light.	298 299
15. Place the slide in a Coplin jar filled with 4× SSCT at room temperature.	300 301
16. Wash for $2 \times 5$ min in $4 \times$ SSCT at room temperature.	302
17. Wash for $1 \times 5$ min in $4 \times$ SSC at room temperature.	303
<ol> <li>Gently pipette 100 μL of DAPI solution (final concentration is 1 μg/mL) on the sample. Incubate for 2 min at room tempera- ture, protected from light.</li> </ol>	304 305 306
19. Rinse in 2× SSC and mount with Vectashield. Seal with nail polish.	307 308

Detection

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The protocols presented here have been applied successfully to perform 3D DNA FISH on embryos up to the 150- to 200-cell stage, after which time the signal decreases from the outside to the inside of the embryo due to poorer probe and antibody penetration. Samples are usually imaged on a laser scanning confocal microscope using a 63× plan apochromat oil objective (numerical aperture of 1.4), with optical sections taken at intervals of 300-700 nm. We have observed that some fosmid probes (about 1 in 5) give a signal that consists of several dots of variable intensities throughout the nucleus, especially when direct-labeled with fluorophores. The reason for this high background remains unknown. In any case, the simplest solution to this problem is to replace the bad fosmid with a neighboring or even overlapping one. Typical 3D DNA FISH results are shown in Fig. 2. In this experiment, we used the centrosomes as extranuclear reference points in order to be able to compare gene positioning in the same blastomere from



**Fig. 2** Analysis of gene positioning in early *C. elegans* embryo using immuno-DNA FISH. Raw data (maximal projections, **a**–**e**) and 3D reconstruction (**f**) of a 4-cell stage embryo that was hybridized to the following probes. (**b**) Fosmid WRM0634bG05 (chr. X) labeled with DIG and detected with mouse anti-DIG and anti-mouse FITC. (**c**) Fosmid WRM0619aE04 (chr. V) labeled with Cy3, TAC-1 protein labeled with mouse anti-TAC-1 and anti-mouse Cy3 prior to FISH. (**d**) Fosmid WRM0628cE09 (chr. II) labeled with DNP and detected with rabbit anti-DNP and anti-rabbit Texas Red. (**e**) Fosmid WRM0623bE05 (chr. III) labeled with Atto647N. Remnants of polar bodies (*asterisks*) hybridize poorly due to high level of DNA condensation. Centrosomes are labeled weakly but clearly (*white arrows*). Bleeding of strong Cy3 signals in the Texas Red channel is sometimes observed (*circled* in (**d**))



different embryos. As expected, the background is somewhat 325 higher when using hapten-labeled probe (DIG and DNP), due to 326 the use of primary/secondary antibodies in the detection protocol, 327 but signals are nonetheless clear. 328

3.2 Live Chromatin In *C. elegans*, a number of laboratories have successfully validated 329 Imaging the *lacO*/lacI system for tagging loci in vivo [2, 3, 5, 11-13]. The 330 technique is a two-step process. The first step involves the expres-331 sion of a fusion between a DNA-binding protein, a fluorescent 332 protein, and a nuclear localization signal. Second, lacO repeats are 333 inserted into the genome either as high-copy injected plasmid 334 arrays, as low-copy bombarded transgenes, or as a single-copy 335 insertion. Strains expressing different lacI fusions can be obtained 336 from the laboratories which created them (see below). We there-337 fore focus on the creation of *lacO* insertions, in particular at a given 338 locus using a modified MosSCI transposon-mediated homologous 339 recombination procedure [7]. In the last part, we describe how to 340 mount live embryos for microscopy and the imaging setups adapted 341 for C. elegans. 342

> The DNA-binding Lac repressor is expressed as a fusion with green 343 fluorescent protein (GFP). Expression levels of the protein have to 344 be kept low, as overexpression elevates the background fluores-345 cence, enhances nonspecific binding, and can cause slow-growth/ 346 sick animals. Two strains expressing GFP-lacI from integrated 347 arrays have been published to date. The first one is based on 348 injected integrated arrays, where GFP-lacI is expressed from the 349 housekeeping promoter *baf-1* (GW396 [2]). The second is based 350 on a bombarded construct in where GFP-lacI is transcriptionally 351 regulated by a *pie-1* promoter (AV696 [5]). In the strain GW396, 352 GFP-lacI is visible from about the 20-cell stage to adulthood due 353 to germline silencing (note that this array has a *vit-5::GFP* intesti-354 nal marker expressed from the late L4/adulthood, which hinders 355 GFP-lacI observation). In the AV696 strain, fluorescent signal 356 decreases rapidly at the beginning of embryogenesis, but is useful 357 to locate chromatin in the germline or in early embryos. The two 358 types of expression constructs cannot be used in conjunction due 359 to germline silencing in *trans* of the *pie-1* promoter construct [14]. 360 Both constructs do not contain *lacO* sites in the plasmids used to 361 create them and therefore do not create a GFP-lacI spot. 362

3.2.2 Creation of lacO-Tagged Transgenes and Insertion of lacO into the Genome

3.2.1 Strains for

lacl-GFP

Expression of GFP-lacl/

To allow visualization of a given transgene or locus, a number of 363 binding sites for the bacterial repressor have to be integrated into 364 the genome. Most of the *C. elegans* published arrays obtained by 365 gonadal injection have been created using plasmids that contain a 366 single *lacO* site (the 17 bp consensus sequence recognized by lacI; 367 e.g., all the Fire library plasmids contain this sequence as a single 368 copy). Due to the high-plasmid-copy number in the injected arrays 369 370

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(several hundreds of copies), they are readily detected by GFP-lacI and create a visible spot, usually at the nuclear periphery linked to their silencing [2, 4]. Smaller transgenes (in the range of 10–50 copies) bound by lacI can be created by co-bombarding plasmids of interest with *lacO* repeats (256 repeat, pSR1 [2, 15]). Although the *lacO* repeats do not get integrated each time, this method leads to about 50 % co-integration rate, with transgenes visible when GFP-lacI is expressed in *trans*. These low-copy transgenes are not subject to silencing and display usually a random localization in early embryos [2, 4]. Finally, single targeted genomic insertions of *lacO* arrays can be achieved using MosSCI and derivatives [6, 7]. As this method is new, a detailed description is given below.

As few as 24 lacI binding sites (*lacO*) are sufficient to allow the formation of a visible spot, although the signal-to-noise ratio depends on the expression level of the fluorescently tagged binding protein lacI. We use 256 *lacO* repeats (about 12 kb in size); however, the actual number of sites which get integrated is likely to be lower. Plasmids with binding site repeats are intrinsically recombinogenic in bacteria. It is therefore highly recommended to use recA strains (DH5 $\alpha$ /XL1 blue) or recB recJ (SURE) for amplification/cloning. When defreezing these strains, it is good practice to isolate single colonies and test the length of *lacO* repeats as the repeat stretch has a tendency to shrink in size.

To insert lacO repeats at a site of interest, one needs a Mos insertion located next to the site of interest. Due to the resolution of optical microscopes, "near" means in the next 20-40 kb [16]. Many Mos insertions (about 13,000) have been created by the NemaGENETAG project (http://elegans.imbb.forth.gr/nemagenetag/). These are available as invalidated or validated insertions. All available insertions can be browsed at http://pbil.univ-lyon1. fr/segalat/data/index.php and ordered at http://ums3421.univlyon1.fr/spip.php?article14. This is the most cost-effective manner to create *lacO*-tagged loci, as compared to ZFN/TALEN methods (Transcription Activator-Like Effector Nuclease [17]), in which the nuclease is engineered to create a double-strand break at a given sequence, providing a template for homologous recombination. MosSCI however requires a Mos insertion to be available in the region of interest. However, given that 20 kb genomic distance cannot be resolved using light microscopy, several kilobases can separate the region of interest and the Mos insertion.

Insertion of the *lacO* repeats is achieved using Mos Single-Copy Insertion (MosSCI [7]) and requires homology between the insertion site and the plasmid that serves as a template for homologous recombination. To efficiently achieve the creation of templates with homology surrounding *lacO* repeats, we designed a triple-plasmid Gateway system with a middle *lacO/Cb unc-119* plasmid. This is used to create a plasmid where the *lacO* repeats are flanked with homology on both sides.

[AU1]

Author's	Proof
[AU3]	

1.	Prepare the <i>C. elegans</i> recipient strain by crossing it to an <i>unc-119(ed3)</i> mutant and selecting for unc worms homozygous for the <i>Mos</i> insertion of interest. At this point, it is good to cross in the transgene for the expression of GFP-lacI to have a visual confirmation of <i>lacO</i> integration later ( <i>see</i> above for strain names).	418 419 420 421 422 423
2.	Design primers to amplify left and right sequences located 5' and 3' of the <i>Mos</i> insertion site (Fig. 3a). A size of 1.5 kb is sufficient for efficient recombination [18]. Primers are designed for Gateway BP cloning. Use following 5' extensions for the following primers from a to d (in $5' \rightarrow 3'$ direction):	424 425 426 427 428 429
	<ul> <li>(a) GGGG ACA ACT TTG TAT AGA AAA GTT G—locus- specific sequence</li> <li>(b) GGGG AG TGG TTT TTT GTA GAA AGT TG_l</li> </ul>	430 431
	(b) GGGG AC TGC TTT TTT GTA CAA ACT TG—locus- specific sequence (reverse complement)	432 433
	(c) GGGG ACA GCT TTC TTG TAC AAA GTG G—locus- specific sequence	434 435
	(d) GGGG AC AAC TTT GTA TAA TAA AGT TG—locus- specific sequence (reverse complement)	436 437
	To minimize amplification errors, we use Phusion polymerase (NEB) according to the manufacturer's instruction with the following cycling conditions: [98 °C, 30"]; [98 °C, 10"; 45 °C, 30"; 72 °C 30"/kb] 5×; [98 °C, 10"; 50 °C, 30"; 72 °C, 30"/kb] 20×; 72 °C, 10' 12 °C ∞. Test amplicon length and perform BP cloning according to the manufacturer's manual (Invitrogen), except that it is preferable to transform and plate at least half of the cloning reac-	438 439 440 441 442 443 444 445
C	tion to get a sufficient number of colonies. In our hands, BP cloning has been highly efficient, with at least seven out of eight colonies tested positive for the desired insertion.	446 447 448
3.	Perform the Gateway LR reaction with the 5', middle [see note above about <i>lacO</i> repeats plasmid], and 3' clones (Fig. 3b). Transform at least half of the LR reaction to get a sufficient number of colonies. Test the plasmids by PCR/	449 450 451 452
	miniprep.	453

- 4. Test the length of *lacO* sites on the destination vector. These 454 should be about 10 kb in size to make a visible spot in vivo. 455 Normally the *lacO* repeats length does not change drastically 456 during the LR recombination, as recombination uses the 457 Clonase mix and transformation is carried in DH5α strains. 458
- 5. Carry on MosSCI according to the original protocol [7, 8]
  (Fig. 3c). In our hands, integration at certain loci was more difficult to achieve than at others. The indirect, heat-shock 461 procedure was preferred in such cases.



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a Amplify Mos insertion flanking regions for Gateway cloning



**Fig. 3** Method for insertion of *lacO* repeats at a given *Mos* insertion site. (a) Regions flanking the *Mos* insertions are amplified by PCR using primers a, b, c, and d with Gateway-specific overhangs (*red*, *see* sequence in the text). Amplified homology stretch is recombined in the Gateway 5' and 3' vectors using BP. (b) The template for recombination is created by LR recombination of the vectors created in (a) with the middle *lacO/Cb unc-119*. (c) *Mos*-mediated single-copy integration is carried out in the strain carrying the *Mos* insertion of interest in an *unc-119(ed3)* background, which leads to the integration of the *lacO/Cb unc-119* template at the *Mos* locus. (d) Insertion is tested by PCR using primers located in the *lacO/Cb unc-119* insertion and outside of the homology stretch

463	6. Check insertion by PCR using primers outside of the homology
164	region and in the inserted transgene (Fig. 3d). Also, check for
465	the presence of a visible spot under the microscope. During
466	the homologous recombination procedure, some <i>lacO</i> repeats
467	might get lost, which therefore makes the GFP-lacI spot difficult
468	or impossible to see.

### Analyzing Nuclear Organization in Nematodes



**Fig. 4** Imaging live embryos carrying *lacO* inserts. (**a**) Preparation of the agarose pad, using three microscope slides. The outer two slides are taped to the bench using thick laboratory tape. Additional layers of tape will determine the thickness of the agarose pad. A drop of molten 2 % agarose is placed in the center of the middle slide and immediately covered with a perpendicular slide, creating a flat pad. (**b**) Embryos dissected in an hourglass are transferred to the agarose pad using a mouth pipette. Excess buffer is removed with the pipette, and embryos are grouped with an eyelash (bar = 20 µm). After covering the embryos with a coverslip, the space around the pad is filled with injection oil, and the coverslip is sealed to the slide using Vaseline/lanoline/ paraffin or Vaseline alone. (**c**) Axial z projection of a stack of optical slices of an embryo expressing GFP-lacl from an integrated large array with *lacO* (GW76, *gwls4[baf-1::GFP-lacl; myo-3::RFP]X* [2]). Two large spots corresponding to the homolog chromosomes are visible in each nucleus (bar = 5 µm). (**d**) Axial z projection of a stack of optical slices of an embryo expressing GFP-lacl from an integrated large array with *lacO* and an insertion of *lacO* repeats created using the technique described in Fig. 1 (GW392, *gwls39[baf-1::GFP-lacl; vit-5::GFP]III gwSi13[lacO/Cb unc-119@ttTi9115]V* [4]). The GFP-lacl-expressing array is not visible as it carries no *lacO* sites. The single-copy insertion forms very small spots (*arrows*)

3.2.3 Preparing Worms and Embryos for Imaging

Author's Proof

Preparing Agarose Pads for Microscopy 1. Prepare the molding bench (Fig. 4a).

Put three microscope slides next to each other on a flat 470 surface. Stick both outer slides with tape to the surface, putting thick laboratory tape on top of the slides. The thickness of 472 the pad will depend on the thickness of the tape glued on the 473 outer slides. Put two to three additional layers of tape on the 474 slides to increase the thickness of the pad. 475

2. Prepare 2 % agarose in water and melt at 95 °C. Agarose can 476 be aliquoted in 0.5-1 mL tubes for future use and stored at 477 room temperature. Remelting of aliquoted agarose is achieved 478 by putting the solution at 95 °C for 5 min followed by brief 479 vortexing. The molten agarose can be kept longer at 70 °C 480 instead of 95 °C. At high temperatures, agarose hydrolyzes 481 and does not harden anymore. To image (moving) worms 482 instead of embryos, add 0.5 µL 10 % NaN<sub>3</sub> to the agarose to 483

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484 485 486			inhibit movement. Alternatively, anesthetizing worms with 400 mM ethanol has been recently shown to deliver similar results [19].
487 488 489 490 491 492		3.	Prepare the agarose pad (Fig. 4a). Put a drop $(50-100 \ \mu L)$ of molten agarose on a slide placed in the middle of the two taped slides. Immediately place a second slide, perpendicularly to the first one, creating a flat surface. This can be kept for several hours if the upper slide is not removed (otherwise the pad dries out and becomes unusable).
493 494 495 496 497 498	Preparing Embryos for Live Imaging	1.	Pick gravid adults with fine forceps, and move them to an hourglass filled with M9 ( <i>see</i> <b>Note 11</b> ). Try to avoid taking too much bacteria as <i>E. coli</i> will stick to embryos once dissected and usually has high autofluorescence. If needed, move the adults with a platinum loop to bacteria-free plates prior to putting them in the M9.
499 500 501		2.	Cut adults in two in the middle of the worm (approximately where the vulva is). Squeeze delicately the head and tail parts to release the embryos from the adult carcasses.
502		3.	Using a mouth pipette, aspirate the embryos (see Note 12).
503		4.	Transfer the embryos to the agarose pad (Fig. 4b). Make sure
504			to hold the tip of the mouth pipette vertically during the trans-
505			fer as failure to do so will lead to dripping of embryos out of
506			the capillary. Once over the agarose pad, blow in the mouth
507			pipette to expel the embryos, allow them to settle on the
508			mouth pipette. The M9 layer should be at a minimum height
510			so that embryos do not move anymore when an evelash is put
511			on the pad.
512		5.	Group embryos together using an eyelash tip (a flexible eyelash
513			fixed on a tip/toothpick with nail polish). This is very helpful
514 515			for finding the embryos under the microscope and minimizes the search time at high magnification.
516		6. 1	Put a coverslip on top of the pad/embryos, making sure to
517		í	avoid trapping too many bubbles, in particular close to the
518		(	embryos as they diffract light. The coverslip should be laid at an
519		i	angle on one side of the pad and then slowly lowered on the
520		(	embryos, pushing away bubbles.
521		7. ]	For longer acquisition times, it is recommended to minimize
522		(	evaporation (which would flatten the pad, damage embryos,
523		í	and shift focus). We use injection oil (mineral oil, Sigma M5904)
524		1	In moder water exchange between the pad and the exterior.
525		-	embryos can develop and batch if kept overnight in such a
0∠0 527			chamber. Put enough oil to fill the gap between the slide and
528		1	the coverslip around the agarose pad
J20			the covership around the agarose pad.



3.2.4 Imaging Living Embryos 8. In order to avoid movement of embryos and sliding of the coverslip/pad, glue the coverslip to the slide using a 1:1:1 mix- ture of Vaseline/lanoline/paraffin (on two sides only to avoid impairing gas exchanges). Alternatively, Vaseline can be used alone but is less solid. Both types of glue have to be melted. 533

As for any live imaging, it is good practice to evaluate whether the 534 development of the worms was not disturbed by the laser expo-535 sure. This is especially important when studying chromatin dynam-536 ics, as animals will experience repeated illumination over an 537 extended period of time. Evaluating whether the movements 538 observed correspond to normal behavior or to DNA repair-linked 539 activities is important. An easy way to check this is to compare 540 illuminated and control embryos at approximately the same stage 541 at time of fluorescent imaging and follow their development over 542 time using DIC imaging. 543

Imaging *lacO*-tagged chromatin from embryos is challenging: 544 embryos are relatively thick (about 20 µm) and diffract light; the 545 lacO/GFP-lacI spot can be very small and highly dynamic; more-546 over, the fluorescent signal is low as overexpression of lacI is dele-547 terious for the animals and C. elegans embryos are highly sensitive 548 to focused light. Several imaging systems were assayed and the best 549 system for this application was a spinning disk confocal micro-550 scope. Confocality is essential as for single-copy lacO repeats, the 551 observed structure is small (Fig. 4d); out-of-focus haze would 552 make it invisible. Two systems can achieve confocal images: point 553 scanning and spinning disk confocal microscopes. In the first sys-554 tem a focused laser beam is moving over the sample at high speed; 555 emitted light is filtered for out-of-focus photons by passing through 556 a pinhole and acquired by a photomultiplier. In these systems, 557 maximum laser power as well as laser damage is concentrated on a 558 single point. An alternative solution which minimizes excitation/ 559 damage is spinning disk confocal devices. In spinning microscopy, 560 the laser beam is split in hundreds of focused sub-beams, rotating 561 on the sample at high speed. The laser power on each point as well 562 as the time spent by the laser on each point of the sample is mini-563 mized but repeated several hundred times per second. Emitted 564 light is filtered through the pinholes, and photons are acquired 565 using a highly sensitive CCD camera. The whole field of view can 566 be acquired in as little as 30 ms; acquisition speed is limited by the 567 frame reading rate of the camera. Using a correct combination of 568 lenses/objective/camera, sampling will meet the Nyquist rate 569 (about 90 nm/pixel). The thickness of the embryos (about 20  $\mu$ m) 570 and the small size of *lacO* transgenes make it very difficult to obtain 571 images of entire embryos while keeping them alive (in the z-axis). 572 Emitted light from objects far from the objective is diffracted by 573 the structures above. Hence, when acquiring image stacks of 574 embryos, we limit our acquisition to the 10 µm closer to the 575

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objective if embryos have to be imaged repeatedly for a longer period of time. Alternatively, if only one time point is to be acquired, we apply high laser power to be able to record light emitted from nuclei located far away from the objective.

Typical images acquired with a spinning disk confocal (TILL Photonics) of embryos carrying two different types of *lacO*-tagged chromatin are shown in Fig. 4. Large arrays obtained by gonadal injection and chromosomal integration (about 300–500 copies of the injected plasmids, 3–5 Mb in size) create large heterochromatic structures, which are located at the nuclear periphery in relation with their epigenetic silencing (Fig. 4c [4]). In Fig. 4d, an embryo with 256 *lacO* repeats integrated as a single copy using the technique described above is shown (Subheading 3.2.2). In each nucleus, the GFP-lacI spot is smaller and emerges from background nuclear fluorescence created by unbound GFP-lacI. Finally, it has to be noted that chromatin inside the nucleus is dynamic; hence precise positioning of the GFP-lacI */lacO* spots necessitates a fast acquisition device. Typically, imaging should not be more than 100 ms per optical slice.

595	4	Notes		
596			1.	Due to the AT richness of the C. elegans genome (66 %), we
597				increase the final concentration of unlabeled dTTP in the nick
598				translation reaction to 20 $\mu$ M. The molar ratio of labeled
599 600				dUTP to unlabeled dTTP is 1:1 (compared to 2:1 or higher in most protocols).
601 602			2.	Our preferred labeled nucleotides for a 4-color 3D DNA FISH experiment are DIG-dUTP, Cv3-dUTP, DNP-dUTP, and
603				Atto647N-dUTP. We have also used successfully FITC-dUTP
604				and Cy5-dUTP. However, labeling with Atto594-dUTP and
605				Atto488-dUTP proved to be inefficient.
606			3.	We use the following primary antibodies at a 1:200 dilution to
607				detect hapten-labeled probes: mouse anti-digoxygenin
608				(Jackson ImmunoResearch) and rabbit anti-DNP (Sigma).
609				The secondary antibodies (1:400 dilution) are highly adsorbed
610				FITC-conjugated goat anti-mouse and Texas Red-conjugated
611				goat anti-rabbit, both from Jackson ImmunoResearch.
612			4.	While the quantity of labeled dUTP given here works well in
613				most cases, we found it had to be increased to $3 \mu\text{L}$ for efficient
614				labeling with FITC-dUTP.
615			5.	. The optimal dilution of DNAse I has to be determined empiri-
616				cally and will result in the generation of fragments of 300-
617				600 bp in length after a 90 min incubation. The activity is mainly
618				influenced by the purity of the template DNA. Initial experi-
619				ments should include a range of dilutions, from 1:500 to 1:10.

- 6. DIG-labeled fragments migrate slower than unlabeled ones of 620 similar length. DNP- and Cy3-labeled probes are poorly 621 stained by DNA-intercalating agents (e.g., ethidium bromide, 622 SYBR Green). Note that the reaction can be stored at -20 °C 623 at this point. If the reaction needs to be resumed after storage, 624 a fresh aliquot of *E. coli* DNA polymerase I should be added to 625 the reaction. 626
- 7. The cleaning of labeled probes (e.g., on ion-exchange spin columns) is optional. However, we have found that performing this step leads to an improvement in signal-to-noise ratio in experiments involving *C. elegans*, and we recommend doing it.
- 8. We use tRNA as a carrier during probe precipitation instead of 631 ssDNA, which sticks to the eggshell and thus gives undesirable 632 background upon DAPI staining.
  633
- 9. We have found that slides have to be transferred from cold 634 methanol (-20 °C) to cold PBS and then cold formaldehyde 635 (4 °C) in order to minimize the loss of embryos. By doing so, 636 we also found that subsequent incubations and washes could 637 be performed in Coplin jars without significant loss of embryos 638 from the slide. 639
- 10. The final probe concentration is  $1-2.5 \text{ ng/}\mu\text{L}$ . In our hands, 640 this range of concentration gives excellent results. The *C. ele-* 641 *gans* probes are repeat poor and can therefore be used at a 642 lower concentration than the ones that are used in FISH 643 experiments on mammalian samples. 644
- 11. In our hands, GFP fluorescence is always higher when worms have been grown at temperature above 20 °C. This is likely due to GFP variants optimized for mammalian expression at 37 °C, as we also observed that fluorescence is higher after heat shock at 34 °C.
- 12. Mouth pipettes are made using capillaries. (a) A 10  $\mu$ L pipette 650 is heated in the flame of an ethanol burner. (b) Once the glass is 651 soft, remove it from the flame and quickly pull apart the ends. 652 (c) Break the ends apart to create a pipette with an end with a 653 diameter of ~40  $\mu$ m. (d) Place the pipette in a mouth pipette 654 aspirator (Sigma A5177 or building plans are available upon 655 request). 656

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# Author Queries

Chapter No.: 11 0001992714

Queries	Details Required	Author's Response
AU1	Please check if "Transcription Activator-Like Effector Nuclease" should be lowercased.	
AU2	Please check if "Mos Single-Copy Insertion" should be lowercased.	
AU3	Please check if edit to sentence starting "A size of" is okay.	

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