Chromatin remodeling in *Drosophila* preblastodermic embryo extract

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Chromatin remodeling is essential for nuclear reprogramming. However, the factors and mechanisms involved in this remodeling are still poorly understood and current experimental approaches to study it are not best suited for molecular and genetic analyses. Here we report on the use of *Drosophila* preblastodermic embryo extracts (DREX) in chromatin remodeling experiments. Our results show that incubation of somatic nuclei in DREX induces changes in chromatin organization similar to those associated with nuclear reprogramming, such as rapid binding of the germline specific linker histone dBigH1 variant to somatic chromatin, heterochromatin reorganization, changes in the epigenetic state of chromatin, and nuclear lamin disassembly. These results raise the possibility of using the powerful tools of *Drosophila* genetics for the analysis of chromatin changes associated with this essential process.

Results

Incubation of somatic nuclei in DREX induces dBigH1 incorporation. An early event in reprogramming of somatic nuclei transplanted into oocytes is the binding to chromatin of the oocyte specific linker histones H1. In this regard, incubation of somatic nuclei prepared from *Drosophila* S2 cells in DREX, which is enriched in dBigH1, resulted in its incorporation to chromatin. Immunofluorescence analyses (IF) showed a clear association of dBigH1 with S2 nuclei after incubation in DREX (Fig. S1a). Notice that no dBigH1 was detected in S2 nuclei prior to incubation. Furthermore, fractionation into soluble nuclear and chromatin bound material, detected the presence of dBigH1 in the chromatin bound fraction (Fig. 1a). dBigH1 binding was detected as early

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as 1′ after incubation in DREX and increased progressively during incubation (Fig. 1a). ChIP-qPCR analysis confirmed these results since, after incubation in DREX, significant dBigH1 occupancy was detected at multiple genomic sites, both single-copy and repetitive (Fig. 1b), suggesting that dBigH1 binding occurred across chromatin.

In nuclear transfer (NT) experiments, binding of the oocyte specific H1s usually results in displacement of the corresponding somatic H1s 8,19–21,30,31. In this regard, no significant reduction in the amount of chromatin bound dH1 was detected after incubation in DREX for up to 2 h (Fig. 1c). In addition, ChIP-qPCR experiments showed that dH1 occupancy was not impaired upon dBigH1 binding (Fig. 1d). Furthermore, IF experiments detected only a weak negative correlation between dBigH1 and dH1 content in DREX-treated nuclei (Fig. S1b). Altogether, these results suggest that dBigH1 binding occurs without significant dH1 displacement. Notice, however, that after 1′ of incubation dH1 content significantly decreased though dBigH1 binding was only weak (Fig. 1c).

Incubation in DREX induces histone acetylation. Nuclear reprogramming is usually accompanied by changes in the epigenetic state of chromatin 32–35. In this regard, we observed that incubation in DREX resulted in a significant increase of global H3Ac (Fig. 2a). ChIP-qPCR experiments confirmed that H3Ac levels increased at multiple loci after incubation in DREX for up to 2 h (Fig. 2b). Increased H3Ac observed upon incubation in DREX was not associated with dBigH1 binding since a similar increase was also observed when dBigH1 binding was strongly impaired by the addition of αdBigH1 antibodies to DREX (Fig. S2). Increased histone acetylation suggests that incubation in DREX induces transition to a more active chromatin conformation. Thus, we also analyzed whether incubation in DREX affected the levels of H3K4me3, a modification accumulating at promoters of active genes. In this regard, although incubation in DREX did not significantly affect global H3K4me3 levels (Fig. 2c), we detected increased H3K4me3 levels at promoters of several genes (Fig. 2d). This increase was higher at promoters of developmentally regulated genes, highly expressed during early embryogenesis but silent in S2 cells, than at ubiquitously expressed genes (Fig. 2d). We noticed that global H3K4me3 showed a tendency to increase at short incubation times (p-value = 0.1309 at 1′) (Fig. 2c). Similarly, although global levels of the chromatin bound active RNap II forms were not significantly affected (Fig. S3a and S3b), they showed a tendency to increase at short incubation times, in particular those of the promoter-proximal Ilo^ext^ form (Fig. S3a) (p-value = 0.1779 at 20′).
H3K27me3 also showed a tendency to increase (Fig. S3c). Altogether, these results suggest that incubation in DREX alters the epigenetic landscape of somatic chromatin.

**DREX induces heterochromatin reorganization and nuclear lamin disassembly.** Next, we analyzed whether incubation in DREX affected heterochromatin organization. For this purpose, we performed IF experiments using α-HP1a antibodies, which mark heterochromatin. In somatic S2 cells, HP1a is distributed in distinct heterochromatin foci (Fig. 3a, top and Fig. S4a). Upon incubation in DREX, the α-HP1a immunostaining pattern was altered, showing a more uniform staining along the nuclei with slight accumulation on the periphery (Fig. 3a, bottom and Fig. S4b). After 2 h of incubation, only ~15% of nuclei preserve the normal HP1a foci pattern. Loss of HP1a foci was not the consequence of a global reduction in HP1a content since global HP1a levels were not significantly reduced upon incubation in DREX for 1 h (Fig. 3c), but, on the contrary, they tend to increase at long incubation times (p-value = 0.0879 at 2 h). We also observed that incubation in DREX induced the formation of aberrant HP1a foci extruding from the nuclear surface in ~15% of nuclei (Fig. 3b). After 2 h of incubation, only ~15% of nuclei preserve the normal HP1a foci pattern. Loss of HP1a foci was not the consequence of a global reduction in HP1a content since global HP1a levels were not significantly reduced upon incubation in DREX for 1 h (Fig. 3c), but, on the contrary, they tend to increase at long incubation times (p-value = 0.0879 at 2 h). We also observed that incubation in DREX induced the formation of aberrant HP1a foci extruding from the nuclear surface in ~30% of nuclei (Fig. 3b). Similar results were obtained when immunostaining was performed with α-H3K9me3 antibodies, which also mark heterochromatin (Fig. S5). Also in this case, a uniform α-H3K9me3 pattern was observed upon incubation in DREX (Fig. S5a) and ~30% of nuclei showed extruded H3K9me3 foci (Fig. S5b). It has been reported that somatic nuclei incubated in *Xenopus* egg extracts eventually disassemble and undergo apoptosis 36. However, the major heterochromatin reorganization observed upon incubation in DREX did not reflect general chromatin destabilization and/or...
apoptosis since release of histones H3 and dH1 was not detected upon incubation in DREX for as long as 24 h (Fig. 3d, bottom). In contrast, chromatin disassembly was observed in control nuclei after 6 h of incubation in the absence of DREX (Fig. 3d, top). Furthermore, ChIP-qPCR experiments showed that incubation in DREX did not significantly reduce H3K9me2 occupancy at several heterochromatic elements, including different types of transposable elements (TE) and satellite DNAs (Fig. 3e), suggesting that heterochromatin stability was not significantly compromised.

Incubation in DREX also induced nuclear lamin disassembly as judged by IF (Fig. 4a). After 2 h of incubation, ~50% of nuclei showed no detectable α-lamin reactivity at the nuclear envelope (NE), compared to only 2% in control nuclei (Fig. 4c). Interestingly, α-lamin reactivity was generally not detected in nuclei showing diffuse α-H3K9me3 immunostaining, while it was intense in nuclei with preserved α-H3K9me3 foci (Fig. 4b). After 2 h of incubation in DREX, all nuclei with α-H3K9me3 foci showed α-lamin reactivity, while this proportion was reduced to ~45% in nuclei lacking usual α-H3K9me3 foci (Fig. 4c). Altogether, these results suggest a correlation between heterochromatin reorganization and nuclear lamin disassembly.

Discussion
Here we report that incubation of somatic nuclei in DREX induces changes in chromatin organization similar to those associated with nuclear reprogramming. On one hand, we observed rapid incorporation of the Drosophila germline specific linker histone dBigH1 into the somatic nuclei. NT experiments performed in Xenopus and mammals showed that incorporation of the oocyte specific linker histone variants B4 and H1oo into the donor nuclei is an early event in nuclear reprogramming8,19–22,30,31. B4 binding precedes loading of oocyte RNApol II and expression of a dominant negative B4 form significantly inhibits transcription of many reprogrammed genes19. Along the same lines, expression of H1oo in mouse ESCs impairs differentiation37 although it does not improve iPSC formation38. How oocyte specific H1s might contribute to nuclear reprogramming remains not well understood. Oocyte specific H1s are less positively charged than their somatic counterparts and, therefore, their
interaction with DNA is weaker and condense chromatin less than somatic H1s, rendering it more accessible to chromatin modifiers, remodelers and transcription factors. In this regard, Xenopus B4 is more mobile than somatic H1s and B4-containing chromatin is more accessible to remodeling factors. B4 binds pervasively across chromatin of the donor nuclei and, concomitantly, somatic H1s are released, suggesting competition of somatic H1s by the oocyte specific variants. However, this competition does not appear to play an important role in reprogramming since overexpression of somatic H1s does not interfere with B4 binding and subsequent activation of pluripotency genes. Moreover, in mouse fibroblasts, binding of H1oo is detected after NT, while release of somatic H1s occurs later at 30' after NT. Similarly, somatic H1s replacement can last hours in NT experiments with bovine cells. Finally, our results indicate that, upon incubation in DREX, dBigH1 binds along chromatin without affecting somatic dH1 occupancy. In fact, dH1 occupancy is significantly reduced only at short incubation times when dBigH1 binding is very low.

Our results also show that DREX induces changes in the epigenetic landscape of chromatin, which are in agreement with the global epigenetic remodeling of chromatin observed during reprogramming of somatic cells to iPSCs. In particular, we observed increased global H3Ac that is maintained throughout the incubation time course. Increased histone acetylation is observed in fully reprogrammed iPSCs and ESC chromatin is hyper-acetylated compared to differentiated cells. We also observed that H3K4me3 levels increased more intensively at promoters of developmentally regulated genes that are silent in S2 cells but highly expressed in early embryogenesis, suggesting their reactivation. Interestingly, pluripotency-related and developmentally regulated genes are known to acquire H3K4me3 at promoters during nuclear reprogramming. Finally, though not statistically significant, global levels of H3K4me3 and the chromatin bound promoter-proximal active RNApol IIo form tend to increase at short incubation times. In this regard, NT experiments in Xenopus showed loading of oocyte basal transcription factors and RNApol II leading to genome-wide transcriptional reprogramming and selective activation of pluripotency genes. Notably, our results showed that increased histone acetylation induced by DREX does not require binding of dBigH1, suggesting that, at least in part, the epigenetic changes occurring during reprogramming do not depend only on the activities of the oocyte specific H1s.

Incubation in DREX also induces profound changes in chromatin/nuclear organization. On one hand, at short incubation times, DREX induces heterochromatin reorganization since HP1a/H3K9me3 foci disassemble. A decrease in the number of HP1a foci has also been reported during reprogramming to iPSCs. In this regard, chromatin of pluripotent cells is largely decondensed and heterochromatin is organized in larger and

Figure 4. Incubation in DREX induces nuclear lamin disassembly. (a) IF with αlamin (white) of somatic S2 nuclei after incubation in DREX or in control conditions for 2 hours, and prior to incubation (0 h). DNA was stained with DAPI (blue). Scale bars are 20 µm. (b) As in (a), but staining also with αH3K9me3 (red). Arrows indicate nuclei showing diffuse αH3K9me3 staining. Scale bars are 7 µm. (c) The proportions of nuclei positive for lamin or not are presented after incubation in DREX or in control conditions for 2 hours. After incubation in DREX, the proportions of lamin positive/negative nuclei are also presented for nuclei showing H3K9me3 foci or not. (N = 202 for DREX; N = 184 for control).
fewer domains that become smaller, more abundant and hypercondensed as cells differentiate41,47–50. Interestingly, incubation in DREX did not decrease H3K9me2 occupancy at multiple heterochromatic elements, suggesting that DREX affects condensation but not the actual heterochromatin content of somatic nuclei. Oocyte specific H1s might be one of the factors contributing to heterochromatin decondensation since, in humans, H1oo is required for decondensation of sperm chromatin51. At long incubation times, HP1α foci reform and extrude from nuclei. Interestingly, extrusion of heterochromatic sequences was also reported in somatic plant cells undergoing meiosis52. Finally, we also observed that DREX induces disassembly of nuclear lamin, a nuclear envelope component of differentiated cells that is absent in ESCs53. Similar results were reported earlier using a Drosophila oocyte cell-free extract54. Nuclear lamin disassembly is considered a marker of reprogrammed cells, since it is detected at the nuclear envelope in partial iPSCs, but not in fully reprogrammed iPSCs55. Interestingly, nuclear lamin disassembly strongly correlates with heterochromatin reorganization, which might account for the heterochromatin extrusion observed after long-term exposure to DREX.

In summary, our results show that DREX induces several changes associates with gain of pluripotency, such as binding of the germline specific linker histone dBigH1, epigenetic remodeling, heterochromatin reorganization and nuclear lamin disassembly. However, it is highly unlikely that DREX induces full reprogramming of somatic nuclei. Nevertheless, the use of DREX offers the possibility of applying the powerful genetics techniques developed in Drosophila to the analysis of factors and mechanisms involved in chromatin remodeling during this essential process.

Materials and Methods

Antibodies. Rabbit αdBigH1 antibodies are described in49 (1:5000 (WB), 1:4000 (IF)). Rabbit αH1 antibodies were kindly provided by Dr. J. Kadonaga and are described in48 (1:20000 (WB), 1:4000 (IF)). Rat αHP1α antibodies are described in56 (1:10000 (WB), 1:400 (IF)). The rest of antibodies were commercially available: rabbit αH4 (Abcam ab10158, 1:15000 (WB)), rabbit αH3K4me3 (Abcam ab8580, 1:2000 (WB)), rabbit αH3Ac (Mill 06-599, 1:20000 (WB)), mouse αH3K9me2 (Abcam ab1220, ChIP 2 μl), rabbit αH3K9me3 (Mill 07-442, 1:75 (IF)), rabbit αPol II Ser2P (Abcam ab5095, 1:5000 (WB)), rabbit αPol II Ser5P (Abcam ab5131, 1:5000 (WB)), and mouse αlamin (DSHB ADL67.10, 1:1000 (WB)). Jackson and ThermoFisher commercial secondary antibodies were used for immunofluorescence, while IRDye 680LT and 800CW conjugates (LI-COR) and horseradish peroxidase-coupled antibody (Jackson) were used for WB detection.

Incubation in DREX and cellular fractionation. DREX was prepared in Exb50 buffer (10 mM HEPES pH 7.6, 50 mM KCl, 1.5 mM MgCl₂, 0.5 mM EGTA pH 8, 10 mM β-glycerophosphate, 10% Glycerol), as described in57. S2 cells were maintained in Schneider medium at 25°C. Nuclei were isolated using Dounce homogenizer and Buffer A containing: 0.23 M sucrose, 15 mM Tris pH 7.4, 60 mM KCl, 0.25 mM MgCl₂, 15 mM NaCl, 0.15 M spermine, 0.5 M spermidine, 0.2 mM PMSF, 14 mM β-MetOH. Purified nuclei were incubated in DREX for the indicated times. In control experiments, nuclei were incubated under the same experimental conditions in Exb50 buffer. After incubation nuclear fractionation was performed. Nuclear pellet was washed in Exb50 buffer and then lysed for 30 min in 10 mM HEPES pH 7.9, 3 mM EDTA, 0.2 mM EGTA, 1 mM DTT, Protease Inhibitor Cocktail. Centrifugation was carried out at 3000 g for 5 min resulting in supernatant (soluble nuclear fraction) and in pellet (chromatin).

Immunostaining experiments. For immunostaining experiments nuclei incubated in DREX were washed, resuspended in PBS and placed on concanavalin slides for 30 min. Fixation was performed for 15 min in 4% paraformaldehyde, followed by washing in PBS. Nuclei were permeabilized in PBS with 0.3% Triton, and blocked in: PBS, 0.3% Triton, 2% BSA. Primary antibody incubation was performed over-night in PBS-T/BSA and appropriate secondary antibody incubation for 1 h. Slides were mounted in Mowiol (Calbiochem-Novabiochem) containing 0.2 ng/ml DAPI (Sigma), visualized in a Leica TCS SPE confocal microscope, and analyzed using FIJI software.

ChIP experiments. For preparing chromatin for ChIP, crosslinking of DREX-incubated and control nuclei was performed in 1.8% formaldehyde for 10 min at room temperature. Glycin (125 mM) was added to stop the reaction. Nuclei were washed with PBS, Wash Buffer A (10 mM HEPES pH 7.9, 100 mM EDTA, 0.5 mM EGTA and 0.25% Triton X-100) and Wash Buffer B (10 mM HEPES pH 7.9, 100 mM NaCl, 1 mM EDTA, 0.5 mM EGTA, 0.01% Triton X-100). The pellet was then resuspended in TE (10 mM Tris-HCl pH 8, 1 mM EDTA) and 1% SDS was added, followed by centrifugation at 3300 g/10 min, at 4°C. TE wash was performed. Then TE buffer with 0.1% SDS and 1 mM PMSF was added. Sonication was performed in 15-ml tubes in Bioruptor sonicator where 26 sonication cycles of 30 s ON/30 s OFF were performed at high intensity. To check the size of the sonicated DNA part of the sample was checked on an agarose gel after de-crosslinking. In continuation, 1% Triton X-100, 0.1% Deoxycholate and 140 mM NaCl were added. Precioiling of chromatin samples was performed with Protein A sepharose for 1 h, followed by addition of antibody and overnight incubation at 4°C. Incubation was continued for additional 4 h upon Protein A sepharose addition. Five washes with RIPA buffer (140 mM NaCl, 10 mM Tris-HCl pH 8, 1 mM EDTA, 1% Triton X-100, 0.1% SDS, 0.1% Deoxycholate), one wash with LiCl ChIP buffer (250 mM LiCl, 10 mM Tris-HCl pH 8, 1 mM EDTA, 0.5% NP-40 and 0.5% Deoxycholate) and two washes with TE buffer were performed. Samples were RNase-treated. De-crosslinking was performed overnight at 65°C upon addition of 0.1 M NaHCO₃ and 1% SDS. In continuation, samples were treated with Proteinase K and DNA extraction with Phenol–Chloroform followed by EtOH precipitation was performed.

For ChIP-qPCR, triplicates were subjected to real-time PCR using SYBR Green I Master Mix and LightCycler 480 Instrument (Roche). Percentages of immunoprecipitated material were calculated by the ΔΔCt method. Primers used in these experiments are listed in Supplementary Table S1.
Data availability. All data generated or analysed during this study are included in this article (and its Supplementary Information files).

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**Acknowledgements**
We are thankful to Miloš Tatarski for help with the preparation of DREX and to Dr. J. Kadonaga for αdH1 antibodies. We are also thankful to Dr. J. Bernués for helpful discussions and advice. This work was supported by grants from MINECO (BFU2015-65082-P), the Generalitat de Catalunya (SGR2014-204), and the European Community FEDER program. This work was carried out within the framework of the “Centre de Referència en Biotecnologia” of the Generalitat de Catalunya. EŠ acknowledges receipt of a NEWFELPRO Fellowship of the Croatian Government and Ministry of Science and Education within EU FP7 Programme framework.

**Author Contributions**
F.A. and E.Š. designed the research, analyzed the data and wrote the manuscript. E.Š. performed most of the experiments, J.F.-M. prepared the DREX and A.C. performed experiments described in Figure 2d during revision of the manuscript. All authors have participated in data interpretation and discussion. All authors have read and approved the final manuscript.

**Additional Information**
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-018-29129-8.

**Competing Interests:** The authors declare no competing interests.

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