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1 **Early epigenetic reprogramming in fertilized, cloned and parthenote embryos**

2 Lessly P Sepulveda-Rincon¹, Edgar del Llano Solanas^{1,2}, Elisa Serrano-Revuelta^{1,2}, Lydia
3 Ruddick^{1,2}, Walid E Maalouf¹, Nathalie Beaujean^{2*}

4 ¹Child Health, Obstetrics and Gynaecology, School of Medicine, University of Nottingham,
5 Nottingham, United Kingdom

6 ²INRA, UMR1198 Biologie du Développement et Reproduction, F-78350 Jouy-en-Josas,
7 France

8 *Correspondence to Nathalie Beaujean, INSERM U846, INRA USC1361, Stem Cell and Brain
9 Research Institute, Department of Pluripotent stem cells in Mammals; 18 avenue Doyen Lépine,
10 69675 Bron, France email: nathalie.beaujean@inserm.fr

11 **Footnotes:**

12 Present address of Lydia Ruddick: Birmingham Women's Fertility Centre, Birmingham, UK,
13 B15 2TG

14 Present address of Edgar del Llano Solanas: Institute of Animal Physiology and Genetics;
15 Rumburska 89, 277 21 Libechov; Czech Republic

16

17

18 **Abstract:**

19 Despite ongoing research in a number of species, the efficiency of embryo production by
20 nuclear transfer remains low. Incomplete epigenetic reprogramming of the nucleus introduced
21 in the recipient oocyte is one factor proposed to limit the success of this technique. Nonetheless,
22 knowledge of reprogramming factors has increased -thanks to comparative studies on
23 reprogramming of the paternal genome brought by sperm upon fertilization- and will be
24 reviewed here. Another valuable model of reprogramming is the one obtained in the absence
25 of sperm fertilization through artificial activation - the parthenote- and will also be introduced.
26 Altogether the objective of this review is to have a better understanding on the mechanisms
27 responsible for the resistance to reprogramming; not only because it could improve embryonic
28 development but also as it could benefit therapeutic reprogramming research.

29 **Keywords: Oocyte ; Nuclear transfer ; Embryonic genome activation ; Histones post-**
30 **translational modifications ; DNA methylation**

31

32 **Introduction to Nuclear Reprogramming**

33 The cells of an adult mammal show a striking variation in structure and function, conferred by
34 the differential expression of tightly regulated and specific gene networks. With few exceptions,
35 individual cell types have been shown to retain the entire genetic content of the totipotent
36 embryo. Yet, specific gene expression patterns associated with differentiated cell states are
37 highly stable and conserved after somatic cell division [1]. The process of restricting expression
38 to lineage-appropriate subsets of genes is ongoing throughout development and is now
39 understood to reflect an accumulation of “epigenetic” changes at specific gene loci [2] and [3].
40 The term epigenetics, coined by Conrad Waddington in the 1940s, is now used to refer to “the
41 study of changes in gene function that are mitotically and/or meiotically heritable and that do
42 not entail changes in DNA sequence” [4] and [5]. These changes include the large scale
43 positioning of chromosomes and genes within the nucleus as well as local modifications to
44 DNA and chromatin [6] and [7]. Epigenetic changes affect the accessibility of DNA to the
45 transcription machinery, hence, gene expression [6] and [7]. Local modifications include
46 histone posttranslational modifications (PTMs such as methylation, acetylation,
47 phosphorylation, and so forth), DNA methylation, and remodeling of the chromatin [4] and [8].
48 Moreover, all these local modifications may specifically recruit factors, as in recruitment of
49 bromodomain proteins to acetylated histones and of chromobox family proteins to methylated
50 histones [9].

51 Each differentiated cell type has a specific profile of epigenetic modifications at key loci,
52 resulting in expression of only type-appropriate genes. Deviations from this profile in vivo are
53 frequently associated with disease [10]. It is also increasingly recognized that deviations from
54 normally observed epigenetic patterning can contribute to the altered cell behavior found by
55 cancer cells [11]. On the other hand, alteration of these epigenetic modifications with the aim
56 of conferring a more developmentally plastic cell state is referred as nuclear reprogramming

57 and is attempted experimentally via a number of different techniques [12] and [13]. The first
58 amphibian and mammalian cloned animals were achieved by inserting a donor nucleus into an
59 enucleated recipient oocyte [14]. In this approach (cloning by nuclear transfer [NT]), the oocyte
60 has to reprogram the injected nucleus, trying to mimic reprogramming of maternal and paternal
61 DNA during natural fertilization (Fig. 1) [15]. Mammalian nuclei have also been reprogrammed
62 by transfer to the germinal vesicle of *Xenopus* oocytes [13] and [16] or by the fusion of donor
63 cells with an “embryonic dominant” cell type [12]. These techniques use the natural
64 reprogramming abilities of oocytes, embryos, and embryonic cells, without requiring
65 knowledge of the precise factors required for reprogramming. However, as knowledge of
66 reprogramming factors has increased, alternative techniques involving exposure of cells to
67 specific combinations of transcription factors have grown in popularity. Nowadays, somatic
68 cells can be virally transfected, at least in mouse, with no more than four key transcription
69 factors (Oct4, Sox2, Klf4, and Myc) to induce pluripotency (Fig. 1) [17]. The availability of
70 induced pluripotent stem cells (iPS cells) from different species is also increasing rapidly [18],
71 although the underlying molecular mechanisms remain to be investigated. Specific
72 combinations of transcription factors have also been used to switch directly from one cell type
73 to another, a process known as transdifferentiation [19].

74 The goals of this experimental nuclear reprogramming are twofold. First, to elucidate the roles
75 of different epigenetic marks (and associated protein complexes) in nuclear reorganization at
76 fertilization and during development and, second, to develop applications that benefit to human
77 health. Such applications include the reprogramming of readily accessible cell types such as
78 dermal fibroblasts to produce cell lines (iPS cells) to be used for drug screening or study of
79 disease pathways [20] and [21]. These iPS cell lines could be used to select the most effective
80 treatment for the individual patient or for the production of cells and organs for autologous

81 transplants without the ethical or immunological problems associated with allogeneic
82 transplantation [20] and [21].

83 As a research tool, nuclear reprogramming continues to yield insights into the mechanisms and
84 complexes involved in differential control of gene expression [13]. Despite this, and successful
85 cloning experiments in a wide range of species, the efficiency of all techniques, as measured by
86 proportion of nuclei leading to developmentally plastic cells or healthy adult animals, remains
87 very low. Considering the possible therapeutic benefits of successful nuclear reprogramming,
88 there is a great deal of interest in understanding the mechanisms responsible for this resistance
89 to reprogramming.

90

91 **Reprogramming at Fertilization**

92

93 In mammalian species, the formation of the embryo begins with the fusion of two highly
94 specialized haploid cells (sperm and oocyte) which gives place to a genetically new diploid
95 organism: the zygote (or 1-cell stage embryo) with two haploid “pro” nuclei, the paternal and
96 the maternal one (Fig. 1). The “early mammalian” or “preimplantation” embryo development
97 compresses the time from fertilization until the implantation of the embryo in the mother's
98 uterus. During this period of development, epigenetic reprogramming of the genome inherited
99 from the gametes is crucial [22] and [23]. Indeed, during the formation of gametes, both oocyte
100 and sperm cells are subjected to epigenetic changes that permit the expression of specific genes
101 required for germ cell development. As gamete maturation is near to completion, a
102 reorganization of the genome occurs. Paternal genome becomes highly methylated and compact
103 as histones are replaced by protamines [24] and [25]. On the other hand, the oocyte undergoes
104 a chromatin restructuring from a nonsurrounded nucleolus (open chromatin with few defined
105 chromatin surrounding the nucleolus and transcriptionally active) to a surrounded nucleolus

106 conformation (highly condensed chromatin with clear presence of chromatin around the
107 nucleolus and transcriptionally silent) (Fig. 2) [26] and [27].

108 From fertilization, both the incoming paternal DNA complement and that of the oocyte itself
109 are reprogrammed in a number of steps, resetting chromatin to the embryonic form capable of
110 undergoing further changes required during development [28] and [29]. The defined epigenetic
111 status of the previous gametes' genome must now turn into a whole new epigenome proper of
112 an early embryo with totipotent capacity [23], [30] and [31]. To do so, the paternal and maternal
113 genome undergo global demethylation, and although many studies have led to contrasted results
114 regarding the dynamics and the extend of this demethylation [32], it appears that the
115 demethylation process continues after the first cell cycle in the preimplantation embryo up to
116 the blastocyst stage in many mammals [33], [34], [35], [36] and [37]. At this point, the first cell
117 lineage determination takes place (the formation of the inner cell mass (ICM) and of the
118 trophoctoderm (TE)) and new methylation patterns emerge together with cell differentiation
119 and specialization until the whole organism is formed [35], [38] and [39].

120 In addition to this DNA demethylation occurring after fertilization, it has been shown in mouse
121 that many of the histones replacing the protamines on the paternal genome are already
122 acetylated such as lysines 8 and 12 of histone H4 [40]. Moreover, for a correct development,
123 the paternal pronuclei has to be hyperacetylated with the further acetylation of lysines 5 and 16
124 of H4 and lysines 9, 14, 18, and 27 of histone H3 [31], [41] and [42]. On the other hand, some
125 histone PTMs such as trimethylation of lysine 20 on histone H4 and trimethylation of lysine 9
126 on histone H3 (H3K9me3, Fig. 2) are inherited exclusively from the maternal pronucleus,
127 creating an asymmetry between the two parental genomes in the embryo (it would not be
128 possible to include in this work all known histone PTMs, their fluctuation and their roles; for a
129 complete review of known histone PTMs see [30]). These asymmetries persist for varying
130 lengths of time in the developing embryo. For an example, lysine 4 methylation on histone H3

131 is evenly distributed throughout DNA by the two-cell stage [42], whereas H3K9me3 remains
132 asymmetrically distributed until the four-cell stage [30]. Other modifications are found to differ
133 from the ICM and TE cells, such as H4/H2AS1P which is much frequent in the nucleosomes
134 of TE than ICM cells [43] or the general methylation of H3K27 which is found only in the ICM,
135 whereas in the TE it is only present in the inactivated X chromosome [44].

136 The function of this asymmetry just after fertilization has not yet been fully understood,
137 although it is thought to be required for a proper development. Indeed, embryos are
138 transcriptionally silent until the end of the one-cell stage, when a small number of embryonic
139 genes are transcribed from the paternal genome [45] and [46]. This asymmetrical minor
140 activation is followed by the major embryonic genome activation (EGA) later on, associated
141 with a much more frequent rate of production of transcripts and the number of genes transcribed
142 [46] and [47]. The reprogramming of histone modifications has been proposed to be significant
143 for triggering transcription and EGA, correlating the accumulation of transcriptionally
144 permissive marks on the paternal genome and minor activation and between more widespread
145 reprogramming and EGA [48], [49] and [50]. Among the differences observed in
146 preimplantation embryo between mammalian species, the timing of embryo genome activation
147 is a major one. In mouse embryos, EGA occurs at two-cell stage, whereas in bovine and rabbit
148 embryos it occurs at the eight-cell stage [51] and [52]. Remarkably though, it is believed that
149 the fourth-fifth cell cycle in the bovine embryos is critical for chromatin remodeling and
150 embryos that are unable to modify their chromatin structure for gene activation arrest at this
151 stage. For example, distribution of H3K27me3 has been studied semiquantitatively in bovine
152 embryos, where levels were found to decrease from oocytes to their minimum at eight-cell
153 stage, corresponding with EGA [53] and [54]. The decline in H3K27me3 is independent of cell
154 division, indicating an active removal mechanism, where histone demethylase KDM6B has
155 been implicated as the enzyme catalyzing the removal [55]. Similarly, it appears that sheep

156 oocytes and embryos have a specific Dnmt1 transcript involved in DNA methylation
157 maintenance whose levels decrease when the embryonic genome becomes active at the 8/16-
158 cell stage. Interestingly, reducing Dnmt1(12b) by RNA interference prevents embryo
159 compaction at the morula stage, showing the importance of DNA methylation for embryonic
160 preimplantation development [56].

161 Therefore, it seems that although the dynamics of some epigenetic marks are not conserved
162 between all mammalian species, they are always closely related with the formation of an “open”
163 chromatin state allowing gene expression regulation during preimplantation development.

164

165 **Reprogramming after cloning by Nuclear Transfer**

166 Cloned embryos are the result of the enucleation of an oocyte and transfer of the diploid nucleus
167 from another cell (Fig. 1). After such NT procedure, donor cell nuclei often get an incomplete
168 reprogramming which is thought to lead to abnormal development in clones [15]. In particular,
169 the donor chromatin needs to undergo epigenetic changes and modifications to get an
170 embryonic-like chromatin structure as seen in sheep, mouse, bovine, and rabbit NT embryos
171 [57], [58], [59] and [60]. The timing and manner to achieve this conformation will depend on
172 the type of cell used as donor for NT. Embryonic stem (ES) cells proliferate fast and appear to
173 have a more open chromatin conformation than cumulus cells, which may have a more
174 compacted genomic structure. This property seems to make the chromatin of ES cells more
175 accessible to the cytoplasm of the recipient oocyte and to efficient reprogramming [61].
176 Similarly, we observed that NT of murine iPS cells results in higher rates of blastocysts and
177 live-born cloned mice than embryonic fibroblasts (46% blastocysts and 1.3% liveborn for iPS
178 cells vs. 3.5% and 0% for fibroblasts, respectively) [62]. Altogether, it seems that chromatin of
179 the donor cells often remains too compact.

180 Trimethylation of lysine 9 of histone H3 has been proposed to limit the success of nuclear
181 reprogramming. H3K9me3 is indeed associated with the repression of transcription [63], and
182 its localization has been shown to be strongly correlated with constitutive heterochromatin,
183 where it recruits heterochromatin protein 1 (HP1 β also called chromobox protein homolog 1)
184 [64]. H3K9me3 distribution has also been revealed to significantly expand during the
185 differentiation of human ES cells into fibroblasts, a process which involves spreading of
186 heterochromatin [65]. Consistent with these observations, H3K9me3 has been shown to persist
187 after bovine and mouse NT experiments (Fig. 2) [58], [66] and [67], and H3K9me3 levels in
188 lymphocytes have been correlated with decreased potential for nuclear reprogramming [68].

189 A number of approaches have targeted H3K9me3 to improve nuclear reprogramming. In cell
190 fusion experiments by Antony et al. [69], the transient induction of histone lysine demethylase
191 KDM4D (also known as JMJD2B) in ES cells increased the proportion of cell reprogramming
192 by 30% despite the rapid restoration of H3K9me3 levels thereafter. Similarly, the transient
193 expression of KDM4D caused a twofold increase in the efficiency of reprogramming somatic
194 cells into iPSCs [70]. Recently, it was shown that removal of H3K9me3 by overexpression of
195 KDM4D can restore transcriptional reprogramming in mouse-cloned embryos [71]. Such
196 transient overexpression of KDM4D in cloned embryos has also been proven to efficiently
197 improve reprogramming both in mouse and human cloning experiments, giving much higher
198 rates of blastocysts [71] and [72].

199 Histone acetylation is also very important for appropriate development in preimplantation
200 embryos. Studies regarding histone acetylation patterns in rabbit embryos [73] and bovine
201 embryos [74], produced either by in vitro fertilization or somatic cell NT, have shown
202 significant differences. In vitro fertilized embryos always presented higher histone acetylation
203 compared with their counterpart cloned embryos, underlying once again the compactness of
204 chromatin after NT.

205 The use of histone deacetylase inhibitors (HDACi), as scriptaid (SA) or trichostatin A (TSA),
206 to increase of acetylated histones and helping the chromatin opening in cloned embryos has
207 been reported. The first successful group obtaining full-term developed embryos after NT from
208 somatic cells was the group led by Kishigami et al. [75], although at almost the same time
209 another study was reported demonstrating that TSA could improve clone development [76]. An
210 increase of the blastocyst yield and improvement of embryo quality after TSA treatment has
211 been obtained with various donor cells: fibroblasts, neural stem cells, spleen cells, and cumulus
212 cells [77]. It has also been reported that this drug can help with gene expression regulation. For
213 example, whereas cloned embryos reported a failure in the expression of Oct4—an important
214 factor for pluripotency maintenance—TSA treatment favored Oct4 expression in the correct
215 number of cells at the blastocyst stage [78] and [79].

216 Thereafter, SA was reported to be a novel HDACi with less toxicity than TSA because it had a
217 high efficiency, not lethal even at high concentrations [80]. Moreover, SA treatment could
218 support full-term development of inbred cloned embryos. In fact, it appears that inhibition of
219 HDAC is an important factor of reprogramming [81]. Hence, the use of HDACi has resulted in
220 significant improvements in cloning efficiency of many species including human [82].

221 Moreover, HDACi also favors global chromatin reprogramming and thereby gene expression
222 in several species such as mouse or pig, by acting not only on acetylation of histones but also
223 on H3K9me3 [83] or even DNA methylation [84] and [85]. HDACi improve genome-wide gene
224 expression regulation bringing total gene expression profile of clones to resemble that of
225 fertilized pups [86]. We also found that addition of HDACi during the first cell cycle in cloned
226 mouse embryos could improve nuclear remodeling of pericentromeric heterochromatin that
227 reorganized around nucleolar precursors such as in fertilized embryos [61]. Remarkably, the
228 use of HDACi was also correlated with increased number of ICM cells and correct further
229 development to term [61].

230 Research on somatic cell NT embryos has been very useful in portraying that these epigenetic
231 modifications not only have the ability to alter the expression of genes but also strongly
232 demonstrate how their misregulation can disturb preimplantation embryonic development.
233 Developmental inefficiency of cloned embryos and aberrant chromatin state seem to be tightly
234 linked. The use of HDACi and of histone demethylases transient expression can however
235 promote the formation of an “open” chromatin structure after NT, improving the reorganization
236 of early embryo nucleus and thereby reprogramming.

237

238 **Reprogramming in Parthenotes**

239 Research in early mammalian development is carried out mostly on fertilized embryos.
240 However, there is another way to study embryo development. Parthenogenetic activation is
241 another valuable model to produce embryos in the absence of sperm fertilization through the
242 artificial activation of a metaphase II oocyte (Fig. 1) [87]. In some species (such as various
243 fishes, ants, snakes, or amphibians) parthenogenesis is a common method of asexual
244 reproduction in which an unfertilized oocyte is able to develop into a whole new individual.
245 Nonetheless, in mammals, parthenogenesis does not occur naturally, and if it does, it is only a
246 consequence of erroneous oocyte maturation and embryos never develop to term [87]. In
247 mouse, developmental arrest of parthenotes occurs before Day 10 of gestation but this time
248 varies among species [88].

249 In normal conditions, ovulated oocytes advance from metaphase I to metaphase II and they
250 remain arrested at this stage until they are fertilized by sperm. For the first cell division to occur,
251 a series of events triggered by the entrance of a spermatozoon, known as oocyte activation,
252 must take place. Broadly, the main trigger factor is the phospholipase- ζ brought by the sperm
253 into the oocyte's cytosol [89]. A number of signaling pathways are then activated, which result

254 in a calcium release inside the oocyte. This calcium increase is translated in the activation of
255 Ca^{2+} /calmodulin-dependent protein kinase II which in turn, will inactivate the “cycle blocking”
256 proteins maturation promoting factor and cytostatic factor. The inhibition of these last two,
257 releases the oocyte from its arrest, and activation can be confirmed by the exocytosis of cortical
258 granules, resumption of meiosis, extrusion of the second polar body, and the formation of
259 pronuclei. Without sperm, it is necessary to artificially induce oocyte activation if
260 parthenogenetic embryos are to be obtained in the laboratory. There are different protocols
261 capable of overcoming the arrested state of a metaphase II oocyte which may include
262 temperature alterations, electrical pulses, and changes in osmolarity [87]. Contemporary
263 protocols are mostly based on calcium mobilizing compounds (i.e., ethanol, strontium, or
264 calcium ionophore) to foster the initial calcium release in the cytoplasm: protein kinase
265 inhibitors or protein synthesis inhibitors (i.e., cycloheximide or 6-dimethylaminopurine) to
266 inactivate the maturation promoting factor and/or the cytostatic factor and, finally, a
267 microfilament inhibitor (i.e., cytochalasin B) to avoid the extrusion of the second polar body
268 [90] and [91]. Indeed, avoiding the extrusion of the second polar body is necessary to maintain
269 the diploidy in the future embryo (Fig. 1) [92]. Thus, diploid parthenotes only possess maternal
270 genetic information and will be homozygous. In particular, diploid parthenotes will not present
271 the two sets of maternal and paternal imprinted genes, reason why, mammalian parthenotes
272 never develop completely unless genetically modified or by the production of chimeras with
273 fertilized embryos [93], [94] and [95].

274 Therapeutically, because these embryos are not normally viable for full development,
275 parthenotes are also being studied as a stem cell source as it would carry very few ethical issues
276 [88]. Moreover, parthenotes are an effective tool to evaluate genetic effects on the process of
277 maternal genomic imprinting [94] and [96]. They also offer a means to study the contribution
278 of maternally derived factors, as well as the absence of paternal factors to early development.

279 In NT experiments, oocyte activation is performed after NT to induce the resumption of meiosis
280 in the oocyte's cytoplasm. Comparing cloned embryos and parthenotes can, therefore, be
281 particularly helpful when it comes to study the precise cytoplasmic factors required for
282 reprogramming within the recipient oocyte. Chromatin reorganization has been compared
283 between fertilized embryos, clones, and parthenotes in few studies. Parthenotes seem to have
284 less problems than their counterpart cloned embryos in adopting the proper heterochromatin
285 conformation at very early stages, at least in mouse and rabbit embryos (Fig. 2) [59] and [97].
286 On the other hand, some epigenetic modifications take place more rapidly in parthenotes.
287 Acetylation of histone H4 after formation of the pronuclei has been observed earlier in bovine
288 and mouse parthenotes, probably due to the absence of the paternal genome [41] and [98].
289 Remarkably, we observed in a preliminary study that supplementation of TSA during the first
290 embryonic cycle as in NT experiments resulted in an even more open chromatin structure in
291 term of histone acetylation and in extended survival of mouse parthenotes post implantation
292 (unpublished data). All these observations make parthenotes an interesting model to study
293 reprogramming by the oocyte's cytoplasmic factors, in the absence of any sperm supply.

294

295 **Conclusion and Perspectives**

296 Epigenetics is the area of molecular science which has been dusted off the shelves and gained
297 a newfound interest. In order to have a better comprehension of the complex interrelationships
298 between all the various components of the epigenome and the way that each individual part
299 operates, it has been essential to decipher key elements of the nuclear reprogramming in early
300 embryos. However, understanding the connection between chromatin structure, gene
301 expression, genome organization, creation of the nuclear architecture, and how all these cellular
302 processes come together during embryogenesis still needs further studies. What it also needs to
303 be remembered is that epigenetic changes can arise from external agents such as environmental

304 cues, dietary, stress, and chemical contaminants to mention some examples, which in turn,
305 cause a chain effect to the chromatin modifying agents and their respective genes or gene
306 families affecting normal development and disease through their actions on the epigenome [10].

307 This is particularly important from a clinical point of view. Indeed, the main goal in a fertility
308 clinic is to raise embryos under the best culture conditions after gamete retrieving and in vitro
309 fertilization to afterward transfer the highest quality embryo to the mother's uterus and achieve
310 a successful pregnancy [99]. This is nowadays an effective and common process thanks to all
311 the research and advancements in assisted reproductive technologies which have been based on
312 the knowledge obtained from studies mainly using mouse embryos because of their easy access
313 and manipulation. Therefore, studies in early mammalian embryos (such as mouse or rabbit)
314 and their reprogramming could possibly help to improve embryo culture conditions to promote
315 development of better quality embryos with higher potential for further development, thus
316 increasing the success rates of assisted reproductive technologies [30], [35] and [51].

317 Elucidation of the roles of epigenetic marks in nuclear reprogramming would also benefit
318 human health, especially the reprogramming of iPS cells. In particular, some recent publications
319 suggest that ES cells derived from cloned embryos may be closer to ES cells derived from in
320 vitro derived embryos than iPS cells in terms of epigenome and transcriptome [82] and [100].
321 We hope better understanding of epigenetic remodeling mechanisms will shed some light on
322 cell reprogramming and further application on stem cell therapies.

323

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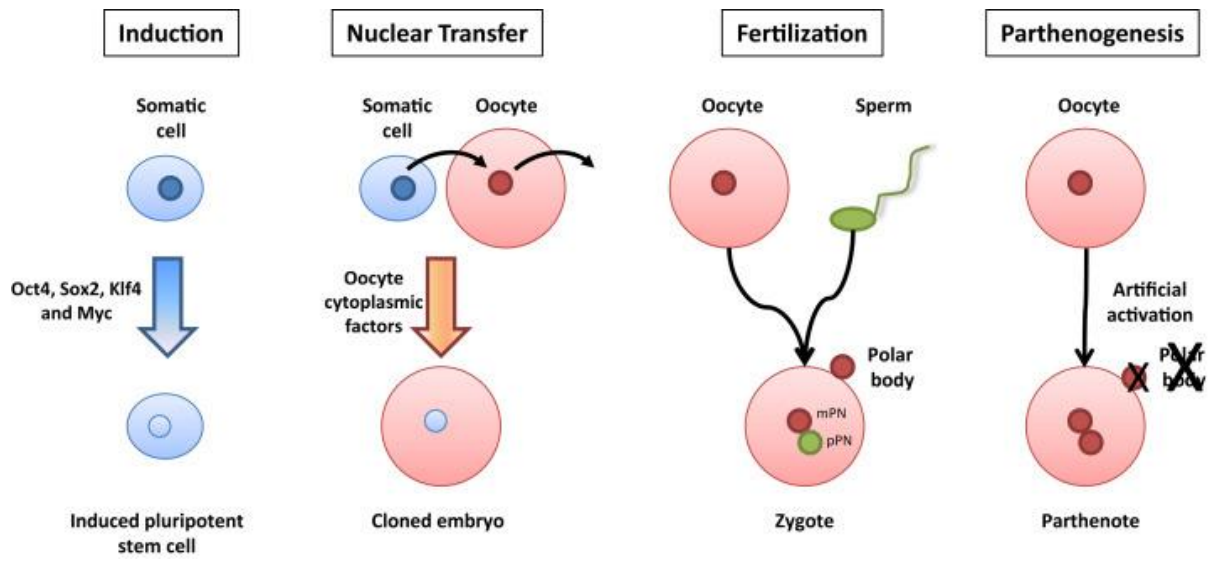
589 **Figure legends**

590 Figure 1: Strategies used to induce nuclear reprogramming include (from left to right):
591 induction by overexpression of embryonic pluripotent transcription factors, nuclear transfer of
592 somatic cell nuclei into enucleated recipient oocytes, fertilization through sperm penetration
593 and parthenogenesis by artificial activation.

594 Figure 2: Examples of H3K9me3 immuno-staining (green) with DNA counterstaining (red) on
595 nuclei from mouse oocytes in NSN (non-surrounded nucleolus) versus SN (surrounded
596 nucleolus) oocytes and in 1-cell stage embryos: either fertilized (zygotes), cloned (obtained by
597 nuclear transfer - NT) or parthenotes. Clear compaction of chromatin and accumulation of
598 H3K9me3 can be observed in SN oocytes. After fertilization, asymmetric distribution can then
599 be observed between the maternal and paternal pronuclei (mPN and pPN respectively) with
600 H3K9me3 accumulation around the nucleolus precursor; whereas cloned embryos present no
601 asymmetry with much more aggregates of H3K9me3, especially at the nuclear periphery. Bar=
602 10 μ m

603

604 **Figure 1**



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607 **Figure 2**

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