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Functional Antagonism between OTX2 and NANOG Specifies a Spectrum of Heterogeneous Identities in Embryonic Stem Cells

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SUMMARY

Embryonic stem cells (ESCs) cultured in leukemia inhibitory factor (LIF) plus fetal bovine serum (FBS) exhibit heterogeneity in the expression of naive and primed transcription factors. This heterogeneity reflects the dynamic condition of ESCs and their versatility to promptly respond to signaling effectors promoting naive or primed pluripotency. Here, we report that ESCs lacking *Nanog* or overexpressing *Otx2* exhibit an early primed identity in LIF + FBS and fail to convert into 2i-induced naive state. Conversely, *Otx2*-null ESCs possess naive identity features in LIF + FBS similar to *Nanog*-overexpressing ESCs and convert poorly into FGF-induced early primed state. When both *Nanog* and *Otx2* are inactivated, ESCs cultured in LIF + FBS exhibit primed identity and weakened ability to convert into naive state. These data suggest that, through mutual antagonism, NANOG and OTX2 specify the heterogeneous identity of ESCs cultured in LIF + FBS and individually predispose them for optimal response to naive or primed inducing factors.

INTRODUCTION

Pluripotency is the capability of a single cell to generate all embryonic and adult cell types. *In vivo* this ability is exhibited by the epiblast, and *in vitro* by pluripotent stem cells (Nichols and Smith, 2009; Rossant and Tam, 2009; Gardner and Beddington, 1988). Mouse ESCs may be derived from both the inner cell mass and early preimplantation epiblast; they can be indefinitely propagated in culture by ensuring provision of leukemia inhibitory factor (LIF) plus fetal bovine serum (FBS) and may efficiently integrate into host blastocysts and contribute to all body tissues (Nichols and Smith, 2009; Silva and Smith, 2008; Martin, 1981; Evans and Kaufman, 1981). However, their state depends strictly on a regulatory network controlled by core pluripotency transcription factors OCT4, SOX2, KLF2/4, NANOG, and ESRRB as well as LIF, WNT, and BMP4 signaling pathways (Kalkan and Smith, 2014; Festuccia et al., 2012; Martello et al., 2012; ten Berge et al., 2011; Silva et al., 2009; Ying et al., 2008). ESCs cultured in LIF + FBS are characterized by cell heterogeneity in both expression of specific transcription factors and sensitivity to signaling molecules, which together define a state ensuring self-renewal and opportunity to convert into naive or primed pluripotency. This cell heterogeneity is exemplified by the fluctuating expression of *Nanog* and by the detection of naive and primed markers in specific ESC sub-type compartments (Smith, 2017; Acampora et al., 2013, 2016; Torres-Padilla and Chambers, 2014; Cahan

and Daley, 2013; Martinez Arias et al., 2013; Muñoz Descalzo et al., 2012; Nichols and Smith, 2011; Kalmar et al., 2009; Hayashi et al., 2008; Chambers et al., 2007). A similar heterogeneity exists in the preimplantation mouse embryo at E4.5–E4.7 when the epiblast gradually loses naive identity and begins to induce early primed pluripotency (Acampora et al., 2016). Recently, the state of the early primed epiblast has been discussed as representing a new phase of pluripotency, named formative, which is interposed between naive and primed pluripotency (Smith, 2017). Formative pluripotency is hypothesized to represent an essential staging post required to enable naive cells to successfully remodel transcriptional, epigenetic, signaling, and metabolic networks in preparation for transit into a mature primed state responsive to differentiation cues (Smith, 2017). ESCs cultured in LIF + FBS may be committed to naive or primed pluripotency if adequately stimulated. For example, ESCs cultured in LIF may convert into a naive state of pluripotency if provided with the two inhibitor molecules (2i), which respectively inhibit FGF signaling and activate WNT signaling (Marks et al., 2012; Nichols et al., 2009; Ying et al., 2008); alternatively ESCs may also convert to a primed state of pluripotency if LIF is replaced with FGF and Activin A (Kunath, 2011; Lanner and Rossant, 2010; Brons et al., 2007; Tesar et al., 2007). Signaling-pathway-mediated modification of the pluripotent state is associated with a response in the expression of specific genes, which ultimately determine the state of pluripotency. This implies that the precise dosage and



relationship between pluripotency factors should determine optimal functioning of the entire circuitry (Smith, 2017; Torres-Padilla and Chambers, 2014; Karwacki-Neisius et al., 2013; Muñoz Descalzo et al., 2012; Takahashi and Yamanaka, 2006; Niwa et al., 2000). For example, *Nanog* overexpression is sufficient to drive LIF-independent self-renewal, and the gene dosage of *Nanog* determines the efficiency with which ESCs can self-renew (Chambers et al., 2007). *Nanog* null ESCs, although predisposed to differentiate, retain an undifferentiated phenotype and ability to self-renew (Theunissen et al., 2011; Silva et al., 2009; Chambers et al., 2003, 2007; Mitsui et al., 2003). On the other hand, OTX2 opposes LIF-independent self-renewal, is required for transition into early primed pluripotency, and stabilizes the EpiSC state (Buecker et al., 2014; Yang et al., 2014; Acampora et al., 2013). OTX2 may also bind to the *Nanog* promoter and positively sustain *Nanog* expression to optimize *in vivo* the specification of the epiblast and *in vitro* the integrity of ESC sub-type compartments (Acampora et al., 2016). We hypothesize that integration of the OTX2 and NANOG antagonistic networks specifies the heterogeneous identity of ESCs cultured in LIF + FBS. To gain insights into this mechanism, we studied whether different dosages of OTX2 and/or NANOG may intrinsically modify the pluripotent state of ESCs cultured in LIF + FBS without the contribution of signaling effectors. Our data suggest that OTX2 and NANOG cooperate to deliver the heterogeneous and flexible identity of ESCs cultured in LIF + FBS from their antagonistic regulatory networks.

RESULTS

OTX2 and NANOG Antagonism Controls the Specification of ESC Sub-types Cultured in LIF + FBS

We recently reported that the combined analysis of NANOG, OCT6, and OTX2 protein expression identified five OCT4⁺ compartments in ESCs cultured in LIF + FBS. In addition to a large naive-like (NANOG⁺-OTX2⁻-OCT6⁻) and a small primed-like (NANOG⁻-OTX2⁺-OCT6⁺) compartment, three potentially transitional compartments were identified that we refer to as pre-naive-like (NANOG⁺-OTX2⁺-OCT6⁻), pre-primed-like (NANOG⁻-OTX2⁻-OCT6⁺), and unassigned (NANOG⁻-OTX2⁺-OCT6⁻) (Acampora et al., 2016). To test our hypothesis that the antagonism between the OTX2 and NANOG regulatory networks coordinates the specification of ESC compartments, we analyzed the expression of these transcription factors in a new allelic series of ESC mutant lines, which have different gene dosages of *Otx2* and/or *Nanog*, including *Nanog* null and *Otx2* null ESC lines.

We first targeted *CreER* into the *Rosa26* locus of wild-type (WT) ESCs (*R26^{CreER/+}*) and used this ESC line to generate all

mutant ESCs (Figure S1). To monitor expression from the *Nanog* locus in *Nanog* null ESCs (*Nanog^{Ch/-}*; *R26^{CreER/+}* referred to as *NanogKO*) *mCherry* (*Ch*) was inserted into one of the *Nanog* alleles. To generate the *NanogKO* mutant, *Nanog^{flox/Ch}*; *R26^{CreER/+}* (referred to as *NanogCKO*) ESCs (Figures S1A–S1C) were plated at clonal density and treated with 4-OH-tamoxifen (Tx) 3 hr later. Only very few *NanogKO* clones could be cultured (Figure S2A) for more than 12 passages (P12). Although these clones generated abundant OCT4⁻ cells with primitive endoderm (PE)-like identity, they also exhibited numerous small and flat OCT4⁺ colonies (Figures S2B and S2C) (Mitsui et al., 2003). These OCT4⁺ cells accounted for 13% of total cells (Figures S2E and S3A; Table S1). We next analyzed the identity of OCT4⁺ cells by monitoring the expression of CH, KLF4, OTX2, and OCT6. Cell-counting experiments showed that compared with WT, in *NanogKO* ESCs the number of OCT4⁺ cells expressing CH or KLF4 was severely decreased, while that of OTX2 and OCT6 increased (Figures S2D, S2F, and S3A; Table S1). OCT4⁺ ESCs were then analyzed for co-expression of OCT6, CH, and OTX2. Compared with WT, in *NanogKO* ESCs the naive-like and the pre-naive-like compartments were heavily reduced, and the primed-like compartment markedly expanded (Figures S2G and S3B; Table S1). Noteworthy, the sum of all compartments accounted for the expected percentage of OCT4⁺ ESCs (Figure S2H). This indicates that the cell-counting analysis is not excluding OCT4⁺ ESC sub-types negative for CH, OTX2, and OCT6 expression. These data revealed that loss of *Nanog* causes a marked expansion of the primed-like compartment at the expense of the naive-like and pre-naive-like compartments (Figure S2I). Based on this, we reasoned that, despite the large expansion of the primed-like sub-type, a fraction of *NanogKO* ESCs retained naive-like and pre-naive-like identity, suggesting that CH⁺ ESCs may include a fraction of cells protected from conversion into primed-like identity. To test this possibility, we sorted CH⁺ ESCs (Figure 1A) from *NanogKO* ESCs and studied whether these *NanogKO* sorted (*NanogKOS*) ESCs generated colonies with a different identity. Alkaline phosphatase (ALP) assay showed that none of the *NanogKOS* ESC colonies was uniformly ALP⁺, but most of them retained an evident mixed staining (Figure 1B). Immunohistochemistry assays and cell counting showed that compared with *NanogKO* ESCs, *NanogKOS* ESCs generated much less PE-like OCT4⁻ cells (Figures 1C, 1D, S2B, S2C, and S4A) and showed a remarkable increase in the percentage of total cells expressing OCT4 (Figures 1F and S2E; Table S1). Cell counting showed that in *NanogKOS* ESCs, the number of CH⁺ and KLF4⁺ cells was decreased compared with WT ESCs but increased compared with *NanogKO* ESCs; in a complementary trend, the number of OCT6⁺ and OTX2⁺ cells in *NanogKOS* ESCs was increased

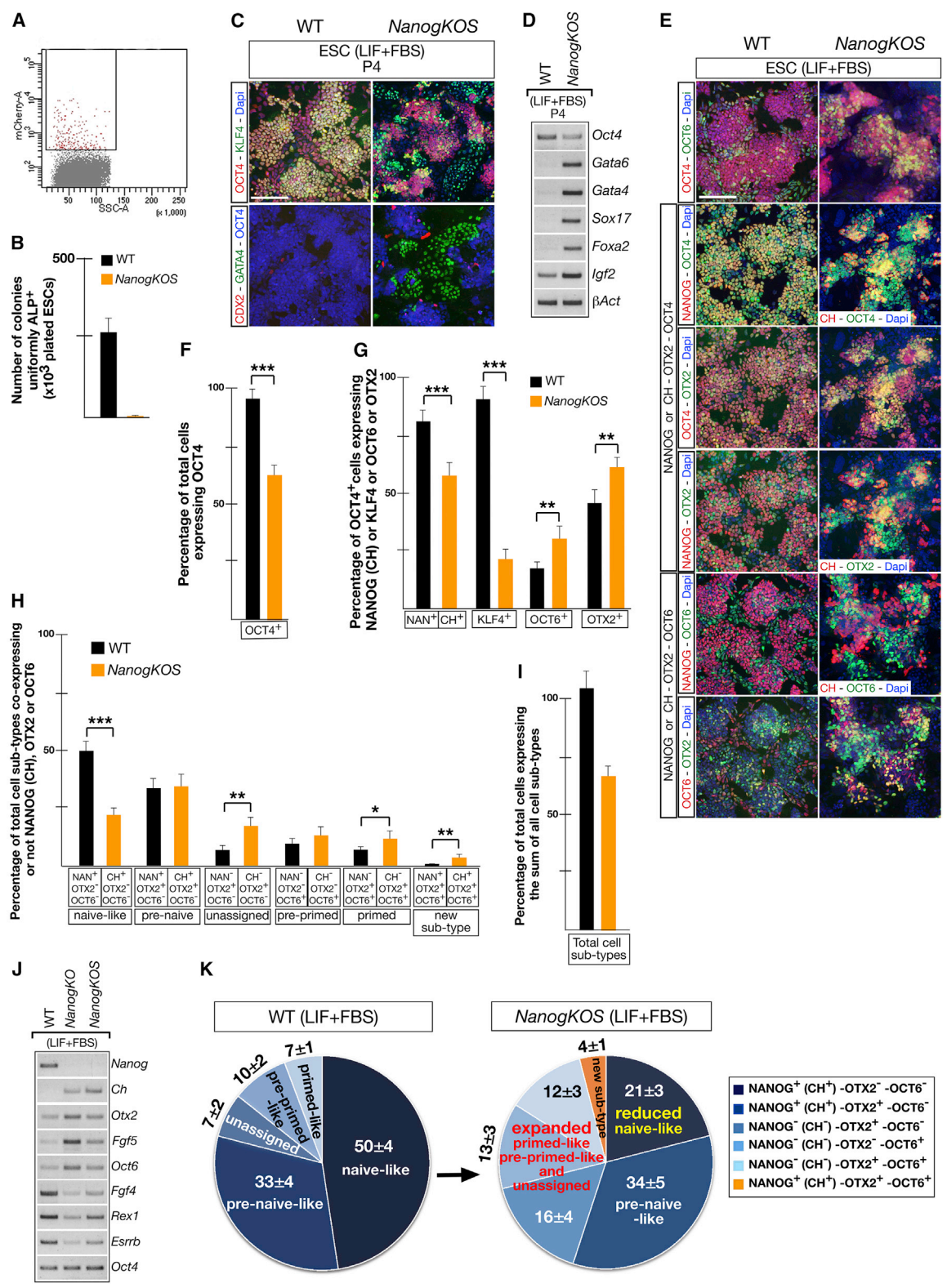


Figure 1. *NanogKOS* ESCs Exhibit Partial Recovery of Naive-like and Pre-naive-like ESC Sub-types

(A) Flow cytometry plot of *NanogKO* ESCs showing CH⁺ cells versus side scatter (SSC).
 (B) Number of WT and *NanogKOS* colonies uniformly ALP⁺. Data are the means ± SD from four independent experiments.

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compared with WT ESCs but diminished compared with *NanogKO* ESCs (Figures 1G, S2E, and S3A; Table S1). Analysis of *NanogKOS* sub-types showed that, compared with WT, the naive-like compartment was severely reduced, the pre-naive-like compartment was unaffected, and the unassigned, pre-primed-like, and primed-like compartments were expanded (Figures 1H and 1K). Compared with *NanogKO*, the size of the naive-like, pre-naive-like and primed-like compartments of *NanogKOS* ESCs was partially restored (Figures 1K and S2I). Coherently, compared with *NanogKO*, in *NanogKOS* ESCs the expression of *Fgf4*, *Rex1*, and *Esrrb* was moderately increased and that of *Oct6* and *Fgf5* decreased (Figures 1J and S4B) (Chambers et al., 2007). These data suggest that the inactivation of *Nanog* has a remarkable impact on the identity of OCT4⁺ ESCs, which become shifted toward the primed-like identity (Figure S2I). Our analysis also reveals that the CH⁺ ESC fraction is protected from complete conversion into primed-like identity (Figure 1K).

Since we initially hypothesized that NANOG and OTX2 act through reciprocal antagonism to control the state of ESCs cultured in LIF + FBS, we next studied mutant ESCs lacking *Nanog* in an *Otx2* null background (*Nanog*^{Ch/-}; *Otx2*^{Gfp/-}; *R26*^{CreER/+} referred to as *DKO*) in comparison with *Nanog*^{Ch/+}; *Otx2*^{Gfp/+}; *R26*^{CreER/+} (referred to as *DHet*) and *Otx2*^{Gfp/-}; *R26*^{CreER/+} (referred to as *Otx2KO*) ESCs (Figure S1). First, *Nanog*^{fllox/Ch}; *Otx2*^{Gfp/-}; *Rosa26*^{CreER/+} (referred to as *NanogCKO*; *Otx2KO*) (Figure S1D) ESCs were plated at clonal density and with Tx to select *DKO* ESCs. Importantly, about 80% of the *DKO* ESC clones could be extensively and stably passaged (Figure 2A). Analysis of ten of these clones showed flat colonies and no sign of differentiation into PE-like cells (data not shown). This indicates that loss of *Nanog* in an *Otx2* null background prevents both loss of efficient self-renewal and differentiation into PE-like cells. We next monitored the ability of mutant ESC lines to generate colonies uniformly stained with ALP.

Compared with WT and *DHet*, *DKO* ESCs similar to *NanogKOS* ESCs produced very few uniformly ALP⁺ colonies, whereas *Otx2KO* ESCs showed a higher number of fully stained ALP⁺ colonies (Figure 2B) (Acampora et al., 2013). Analysis of trypsinized living cells showed that compared with *DHet*, the ratio between CH⁺ and GFP⁺ cells is reversed in *DKO* ESCs (Figure 2C). Of note, the CH⁺-GFP⁻ ESC subset was absent in *DKO* mutant (Figure 2C). Cell counting of immunohistochemistry experiments showed that OCT4 was ubiquitously expressed in all mutant ESCs (Figures 2D and 3A). The number of NANOG⁺ or CH⁺ and KLF4⁺ cells was similar in WT and *DHet* and reduced in *DKO* ESCs, and the number of OTX2⁺ or GFP⁺ and OCT6⁺ ESCs was severely diminished in *Otx2KO*, moderately reduced in *DHet*, and massively increased in *DKO* ESCs (Figures 2D and 3B; Table S2). We next analyzed how these abnormalities in cell number affected the size of ESC sub-type compartments. Compared with WT, *DHet* ESCs exhibited a moderate expansion of the naive-like compartment and a corresponding contraction of the pre-naive-like sub-type; *Otx2KO* ESCs lacked the pre-primed-like sub-type and displayed a significant reduction of the primed-like and pre-naive-like compartments, which was counterbalanced by the marked expansion of the naive-like sub-type; and *DKO* ESCs lacked the naive-like sub-type and heavily expanded the size of the primed-like compartment (Figures 3C, 3D, and S3C; Table S2). Noteworthy, *DKO* ESCs, like *Nanog* null ESCs (Figures 1K and S2J), showed a small new sub-type co-expressing CH, GFP, and OCT6 (Figures 3C and 3D). These data were further corroborated by the expression of additional markers and signaling effectors. Compared with WT, naive markers such as *Klf4*, *Esrrb*, *Rex1*, the phosphorylated form of the LIF signaling effector STAT3 (p-STAT3) and the phosphorylated form of the β CATENIN (p- β CAT) were almost unaffected in *DHet*, downregulated in *DKO*, and several of them such as p-STAT3, *Esrrb*, *Klf4*, and

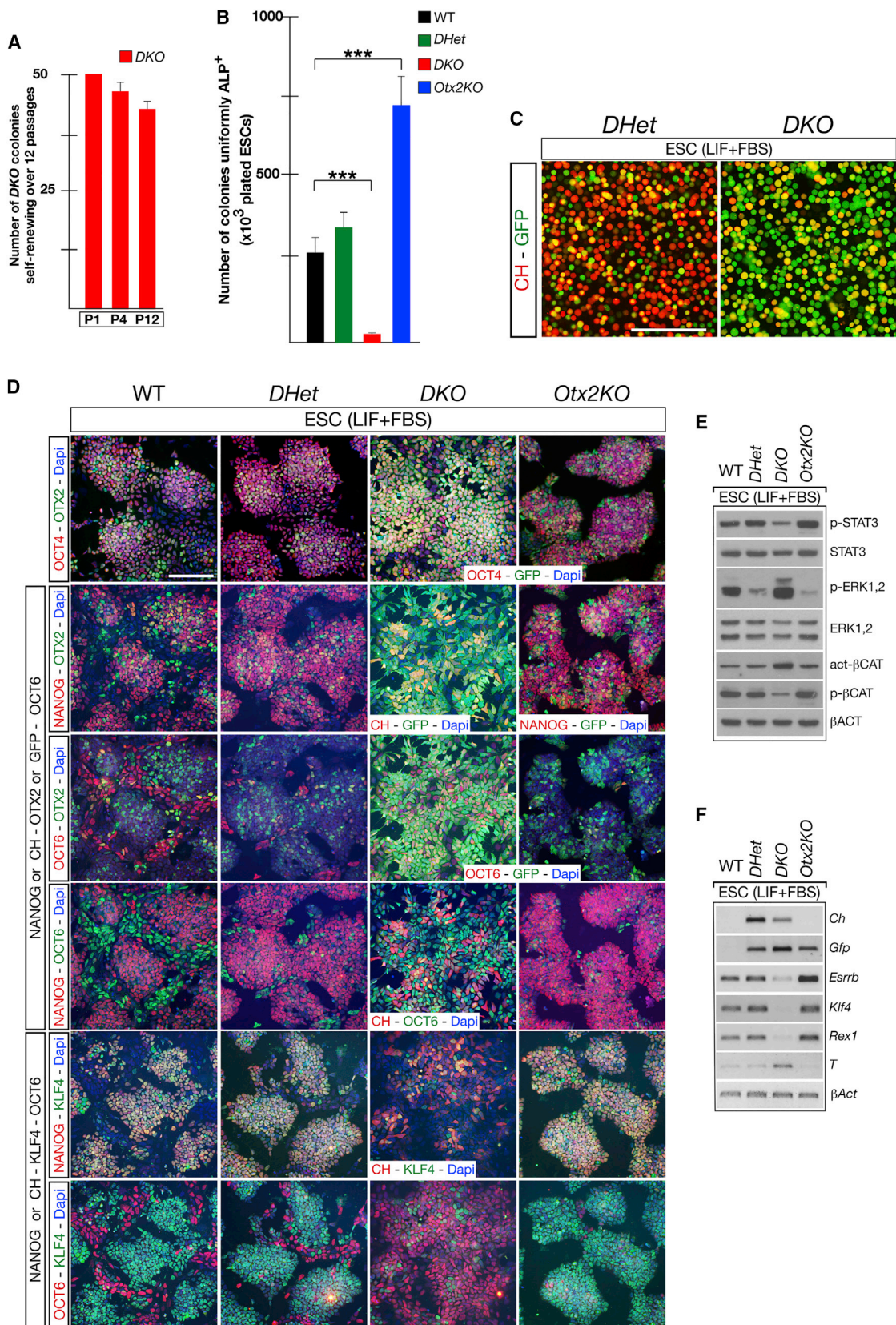
(C) Representative images of WT and P4 *NanogKOS* ESCs immunostained with OCT4 and KLF4 and with CDX2, GATA4, and OCT4. Scale bar, 100 μ m. (D) RT-PCR assays with *Oct4* and PE (*Gata6*, *Gata4*, *Sox17*, *Foxa2*, and *Igf2*) markers on WT and *NanogKOS* RNAs normalized by β -Actin (n = 3 independent experiments).

(E) Representative images of WT and *NanogKOS* ESCs immunostained with OCT4 and OCT6, with NANOG or CH, OTX2, and OCT4 and with NANOG or CH, OTX2, and OCT6; ESCs are also stained with DAPI. Scale bar, 100 μ m.

(F–I) Cell-counting analysis showing in WT and *NanogKOS* ESCs the percentage of total cells expressing OCT4 (F), the percentage of OCT4⁺ cells expressing NANOG or CH, KLF4, OCT6, and OTX2 (G), the percentage of total cell sub-types showing naive-like, pre-naive-like, unassigned, pre-primed-like, and primed-like identity (H), and the percentage of total cells expressing the sum of all cell sub-types (I). Data are reported as means \pm SD from four independent experiments. ***p \ll 0.001; **p < 0.001; *p value is between 0.005 and 0.001. (J) RT-PCR assays showing the expression of *Nanog*, *Ch*, *Otx2*, *Fgf5*, *Oct6*, *Fgf4*, *Rex1*, and *Esrrb* in WT, *NanogKO*, and *NanogKOS* RNAs normalized by *Oct4* (n = 3 independent experiments).

(K) Schematic representation of ESC sub-type compartments showing that *NanogKOS* ESCs exhibit a reduction of the naive-like compartment and moderate expansion of the unassigned, pre-primed-like, and primed-like compartments. A color code of the different compartments is also shown.

See also Figures S1–S4 and Table S1.



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Rex1, were upregulated in *Otx2KO* ESCs (Figures 2E, 2F, and S4C). Primed markers such as *T* and p-ERK1,2 were upregulated in *DKO*, while in *Otx2KO* and *DHet* ESCs, p-ERK1,2 was downregulated (Figures 2E, 2F, and S4C). These data indicate that (1) in contrast to *Nanog* null ESCs, loss of *Nanog* in an *Otx2* null background generates a compensatory effect resulting in a restored LIF-dependent self-renewal and suppression of PE-like differentiation, and (2) in contrast to *Otx2KO*, the identity of *DKO* ESCs exhibits a marked expansion of the primed-like compartment. Overall, these data suggest that NANOG and OTX2 cooperate through their antagonistic transcriptional networks to specify and maintain the identity of ESCs cultured in LIF + FBS.

Identity Assignment of *DKO* and *Otx2KO* ESCs by Transcriptome Analysis

To evaluate the identity of *DKO* and *Otx2KO* cells, we used global transcriptome analyses. We compared *DKO* and *Otx2KO* ESCs cultured in LIF + FBS with WT ESCs cultured in LIF + 2i, LIF + FBS, or in FGF2 + Activin A for 44 hr to generate EpiLCs (44 hr) (Figures 4 and S5; Tables S3–S5). We first performed pairwise analysis of *DKO* ESCs (LIF + FBS) and WT EpiLCs (44 hr) both compared with WT ESCs (LIF + FBS) or WT ESCs (LIF + 2i) to identify *DKO* ESCs transcripts in common with or differentially expressed in WT EpiLCs (Figures S5A–S5D; Tables S3–S5). These data were integrated to generate comparative Venn diagrams showing that *DKO* exhibited 999 (632 upregulated and 367 downregulated) transcripts in common with those specifically expressed in WT EpiLCs (44 hr) (box [j] highlighted by red circle in Figures 4A and 4B; Tables S3–S5). In contrast, *DKO* shared only 360 and 208 differentially expressed genes with those specifically expressed in WT ESCs (LIF + FBS) and WT ESCs (LIF + 2i), respectively (box [j] highlighted by red circle in Figures 4C, 4D, S5M, and S5N; Tables S3–S5). RT-PCR assays validated a number of *DKO* ESC (LIF + FBS) transcripts selectively up- or downregulated in WT EpiLCs (44 hr) or WT ESCs (LIF + FBS) (Figures 4I, 4J, S5I, and S5J). These data suggest that *DKO* ESCs exhibit a transcriptome profile with a marked signature in common with WT EpiLCs

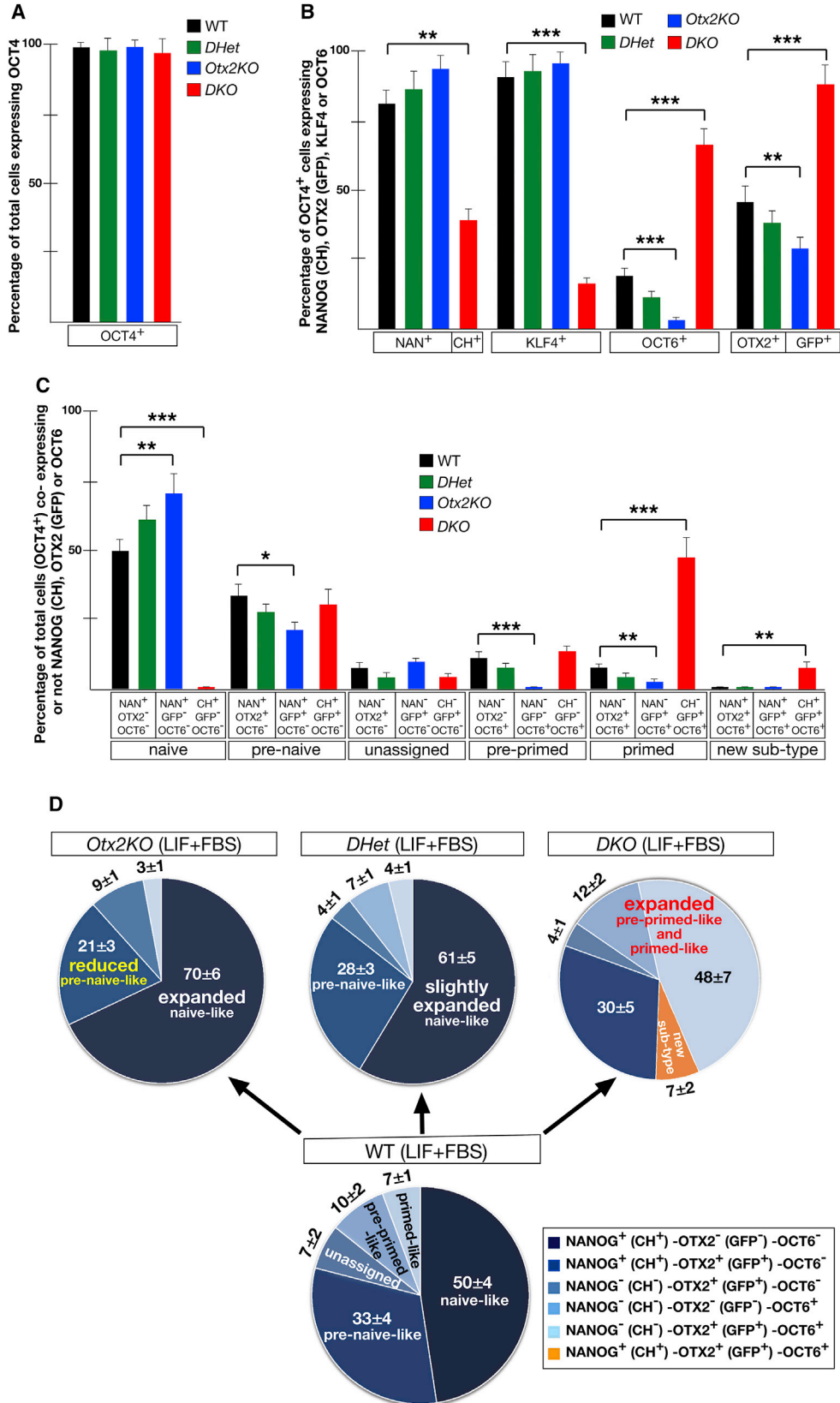
(44 hr). A similar analysis was performed for the transcriptome of *Otx2KO* ESCs (LIF + FBS). Pairwise analysis of *Otx2KO* ESCs (LIF + FBS) and WT ESCs (LIF + 2i) both compared with WT EpiLCs (44 hr) (Figures S5E and S5F), and pairwise analysis of *Otx2KO* ESCs (LIF + FBS) and WT ESCs (LIF + FBS) both compared with WT ESCs (LIF + 2i) (Figures S5G and S5H) were performed to identify the *Otx2KO* ESCs (LIF + FBS) transcriptome fraction in common with or differentially expressed in WT ESCs (LIF + 2i) or WT ESCs (LIF + FBS). Comparative Venn diagrams showed that, although the highest overlap was with WT ESCs (LIF + FBS), specific transcripts (box [j] highlighted by red circle in Figures 4E and 4F), *Otx2KO* ESCs (LIF + FBS) also showed 215 gene transcripts in common with those specifically downregulated in WT ESCs (LIF + 2i) (box [j] highlighted by red circle in Figure 4H). *Otx2KO* ESCs (LIF + FBS) shared very few differentially expressed transcripts ($n = 55$) with WT EpiLCs (44 hr) (box [j] highlighted by red circle in Figures S5O and S5P). This suggests that OTX2 is required to activate genes repressed by LIF + 2i in WT ESCs. These data were also validated by RT-PCR assays (Figures 4K, 4L, S5K, and S5L). Then we compared the transcriptome of *DKO* ESCs (LIF + FBS) with that of *Otx2KO* ESCs (LIF + FBS). We identified 4,209 differentially expressed transcripts, which included 264 of the 500 genes reported as those more sensitive to NANOG (Festuccia et al., 2012) (Figure S5Q; Table S3, sheet 25). Based on this, we compared these 500 NANOG-sensitive genes (250 up- and 250 downregulated) with the OTX2-sensitive genes (230 up- and 378 downregulated) previously identified (Table S3, sheets 19 and 20). Pairwise analysis showed an overlap of 82 genes (Figure S5R), which included 18 genes downregulated in *Nanog* null and upregulated in *Otx2KO* ESC lines, 58 genes with the opposite expression profile and 6 genes downregulated in both *Nanog* and *Otx2* null ESCs (Table S3, highlighted in sheet 26). Noteworthy, 17 of the 18 genes were in common with those selectively upregulated in WT ESCs (LIF + FBS) and included *Klf4* and *Tbx3*, which are required to maintain pluripotency (Niwa et al., 2009); on the other hand, 23 out of the 58 genes were shared with those selectively upregulated in WT EpiLCs (44 hr). This suggests that the

Figure 2. NANOG and OTX2 Antagonism Controls the Identity and Size of ESC Sub-type Compartments and Influences Self-Renewal (A and B) Number of *DKO* colonies with efficient self-renewal (A) and number of ESC colonies uniformly stained with ALP in WT and mutant ESCs (B). Data are reported as means \pm SD of four independent experiments. *** $p < 0.001$.

(C) Representative images showing that *DHet* and *DKO* trypsinized ESCs exhibit mirror distribution of GFP⁺ and CH⁺ cells. Scale bar, 100 μ m. (D) Representative immunohistochemistry assays performed in WT, *DHet*, *DKO*, and *Otx2KO* ESCs with OTX2 or GFP and OCT4, with NANOG or CH, OTX2 or GFP and OCT6, and with NANOG or CH, KLF4 and OCT6; ESCs are also stained with DAPI. Scale bar, 100 μ m.

(E and F) Western blots (E) and RT-PCR assays (F) performed on WT, *DHet*, *Otx2KO*, and *DKO* ESCs to assess the expression level of p-STAT3, STAT3, p-ERK1,2, ERK1,2, activated- β CAT and p- β CAT (E) as well as Ch, Gfp, Esrrb, Klf4, Rex1, and T (F). β -Actin is used as quantitative internal standard ($n = 3$ independent experiments).

See also Figures S1 and S4.



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antagonism between OTX2 and NANOG may depend, at least in part, on opposing transcriptional regulation exerted directly or indirectly by these two transcription factors on a limited number of common target genes.

DKO ESCs Colonize Preimplantation Embryos with Reduced Efficiency

We next examined whether *DKO* ESCs may colonize host blastocysts and be maintained in postimplantation embryos and/or adult mice. The contribution from mutant ESCs was assessed by immunostaining with the ER antibody recognizing *R26*-driven constitutive expression of CreER (Figure S1), PCR assays, and coat-color inspection. *DHet* and *DKO* ESCs cultured in LIF + FBS or in LIF + 2i up to P5 were injected into E2.5 host embryos, which were allowed to develop *in vitro* for 24–30 hr before implantation. We found that, in contrast with *DHet*, *DKO* ESCs did not generate evident chimerism in E12.5 embryos and mice (Figure S6A). This may be contributed by intrinsic loss of their preimplantation colonizing ability and/or loss of pluripotency. To rule out the last possibility, we analyzed *DKO* teratomas and found that they generated derivatives of all germ layers (Figure S6B). Then, we studied the colonization efficiency of *DKO* ESCs in preimplantation embryos. Mutant ESCs were injected at E2.5 in WT embryos cultured up to day 4.75 and tracked using CreER immunofluorescence. Cell counting of ER⁺ ESCs versus DAPI⁺ total cells showed that, compared with *DHet* and *Otx2KO*, *DKO* ESCs were less efficiently integrated into host embryos (Figure 5A; Table S6). Then we studied the identity of both injected mutant ESCs and cells of the host embryos. We first determined the percentage of ER⁺ cells expressing NANOG or CH and OTX2 or GFP. We found that compared with *DHet*, the percentage of ER⁺ cells expressing NANOG or CH was increased in *Otx2KO* and diminished in *DKO* ESCs, while the percentage of ER⁺ cells expressing GFP was significantly increased in *Otx2KO* and *DKO* ESCs (Figures 5B and 5E; Table S6). When extended to KLF4, GATA4, OCT6, and CDX2, this analysis showed that compared with *DHet* ESCs, the percentage of ER⁺ cells expressing KLF4 was remarkably diminished in blastocysts injected with *DKO* ESCs; ER⁺ cells co-expressing OCT6 or

GATA4 or CDX2 were not detected (Figures 5B and 5E; Table S6). This analysis suggests that the host environment slightly modifies the expression of NANOG or CH, OTX2 or GFP and KLF4 described *in vitro* for mutant ESCs, while it efficiently prevents/suppresses OCT6. These data were further confirmed by determining the percentage of total cells co-expressing ER with NANOG or OTX2 or KLF4 (Figure 5C; Table S6). Then, we studied whether the percentage of total cells expressing NANOG, OTX2, or KLF4 in host cells (ER⁻) was affected. For these markers, this percentage was slightly increased only in blastocysts injected with *DKO* ESCs (Figures 5D and 5E; Table S6), suggesting that it may be a consequence of the reduced colonizing ability of *DKO* ESCs.

Otx2 Overexpression Is Sufficient to Induce Early Primed-like Identity in ESCs Cultured in LIF without the Contribution of FGF

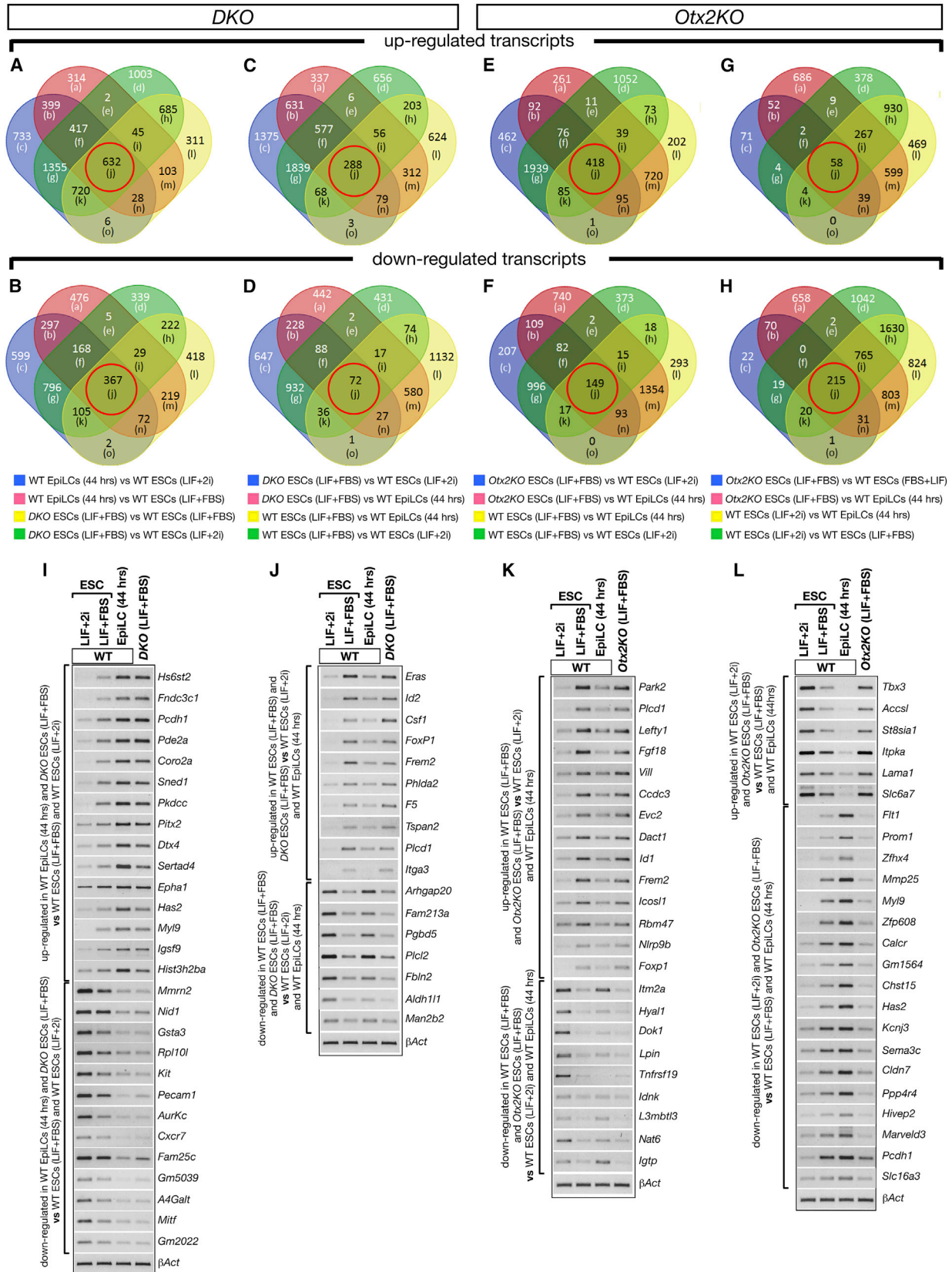
Transition of ESCs into early primed pluripotency is characterized by rapid and wide activation of OTX2 and OCT6 (Acampora et al., 2016). Therefore, we investigated whether ubiquitous activation of OTX2 in ESCs cultured in LIF + FBS was sufficient to impart an early primed-like identity similar to that observed when FGF2 was administered to WT ESCs. To this aim, we generated a further mutant ESC line (*pPyCAGOtx2-ER*) showing constitutive expression of a fusion protein between OTX2 and ERT2 (OTX2-ER) (Figure S7A). Western blot analysis of the Tx-untreated *pPyCAGOtx2-ER* selected clone (red arrow in Figure S7B) showed that the amount of the OTX2-ER fusion protein was about four times that of the endogenous OTX2. Assuming that the endogenous OTX2 was expressed in about 40% of ESCs and that the OTX2-ER protein is present in all ESCs, we concluded that the OTX2-ER level/cell should be about two times that of the endogenous OTX2. Shortly after Tx administration, OTX2-ER was translocated to the nucleus (Figure S7C). Compared with WT, in *pPyCAGOtx2-ER* ESCs, the distribution and/or expression level of a number of markers including NANOG, OCT4, KLF4, OCT6, *Esrrb*, *Rex1*, *Fgf5*, *T*, *Foxa2*, and *Sox1* was apparently not influenced by OTX2-ER when Tx was absent (Figures 6A, 6B, and S4D). However, when Tx was added

Figure 3. Cell-Counting Analysis of Mutant ESC Sub-type Compartments

(A–C) Cell-counting data showing for WT, *DHet*, *DKO*, and *Otx2KO* ESCs the percentage of total cells expressing OCT4 (A), the percentage of OCT4⁺ cells expressing NANOG or CH, KLF4, OCT6 and OTX2 or GFP (B), and the percentage of total cells (OCT4⁺), corresponding to the naive-like, pre-naive-like, unassigned, pre-primed-like, primed-like compartments (C); note that *DKO* ESCs show a new small sub-type (CH⁺-GFP⁺-OCT6⁺) (C). Data are reported as the means ± SD from four independent experiments. ***p << 0.001; **p < 0.001; *p value is between 0.005 and 0.001.

(D) Schematic representation of ESC compartments showing that *Otx2KO* ESCs exhibit a relevant expansion of the naive-like compartment counterbalanced by a significant contraction of the pre-naive-like and loss of the pre-primed-like compartments; and *DKO* ESCs show loss of the naive-like compartment and expansion of primed-like and pre-primed-like compartments.

See also Figure S3 and Table S2.



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to LIF + FBS-containing medium, nuclear translocation of OTX2-ER induced a profound change in the identity of ESCs, which exhibited marked similarity with WT ESCs primed with FGF2 (Figure 6A). Indeed, Tx-treated *pPyCAGOttx2-ER* ESCs (LIF + FBS) and WT ESCs exposed to FGF2 in KSR retained an ubiquitous distribution of OCT4 and exhibited a relevant contraction of NANOG⁺ and KLF4⁺ cells counterbalanced by the expansion of OCT6⁺ cells (Figures 6A, 6C, 6D, and S7D; Table S2). Analysis of ESC sub-types showed that in Tx-treated *pPyCAGOttx2-ER* ESCs, the naive-like sub-type was lost, the pre-naive-like sub-type was reduced, and the primed-like sub-type was expanded (Figures 6E and 6F; Table S2). Importantly, FGF2-mediated priming of WT ESCs induced a similar organization of sub-type compartments (Figures 6E and 6F; Table S2). Moreover, in Tx-treated *pPyCAGOttx2-ER* ESCs, additional naive factors such as *Esrrb* and *Rex1* were downregulated, whereas primed markers such as *Fgf5*, *T*, and *Foxa2* were upregulated (Figures 6B and S4D). Noteworthy, when Tx administration was interrupted at P1, the expression of NANOG, OTX2, OCT4, KLF4, and OCT6 reverted and, at P3, was very similar to that of WT ESCs (Figure S7E). These findings were confirmed in a second independent clone (black arrow in Figure S7B) (data not shown). These data indicate that in LIF + FBS medium, OTX2 is sufficient, without the contribution of FGF2, to induce an expansion of the primed-like ESC compartment as observed in FGF2-primed WT ESCs and *NanogKO* ESCs in LIF + FBS medium. Then, we analyzed the response to FGF2 of *NanogKOS*, *DKO*, and *Otx2KO* ESCs. Indeed, on the basis of their identity in LIF + FBS, the *NanogKOS* and *DKO* ESCs should be primed more efficiently than *Otx2KO* ESCs. Consistently, when compared with WT, *NanogKOS* and *DKO* cells exhibited higher induction of OCT6 and lower expression of CH, while *Otx2KO* cells induced OCT6 in fewer cells and retained a higher number of NANOG⁺ cells (Figure S7F). OTX2 or GFP were detected in most of the *NanogKOS* and *Otx2KO* cells and virtually in all *DKO* cells (Figure S7F). We next analyzed the identity of Tx-treated *pPyCAGOttx2-ER* ESCs in chimeric blastocysts. We found that the percentage of total cells expressing ER was not

affected by Tx treatment (Figure S7G). Marker analysis showed that, compared with Tx-untreated blastocysts, in Tx-treated blastocysts, KLF4 and NANOG were detected in a reduced percentage of ER⁺ cells, OCT6 was activated in a small but significant percentage of ER⁺ cells, and GATA4 and CDX2 were detected only in ER⁻ cells (Figures S7H and S7K; Table S6). These data were confirmed by determining the percentage of total cells expressing NANOG, KLF4, and OCT6 in ER⁺ donor cells (Figures S7I and S7K; Table S6). The complementary analysis showed that the percentage of total cells expressing NANOG or KLF4 in ER⁻ host cells was not significantly affected (Figures S7J and S7K; Table S6). These findings suggest that the wide activation of OCT6 observed in cell-culture experiments is partially prevented by the host environment, which instead is unable to recover loss of naive markers.

Loss of *Otx2* and/or *Nanog* Affects the Responding Competence to 2i-mediated Conversion into Naive State

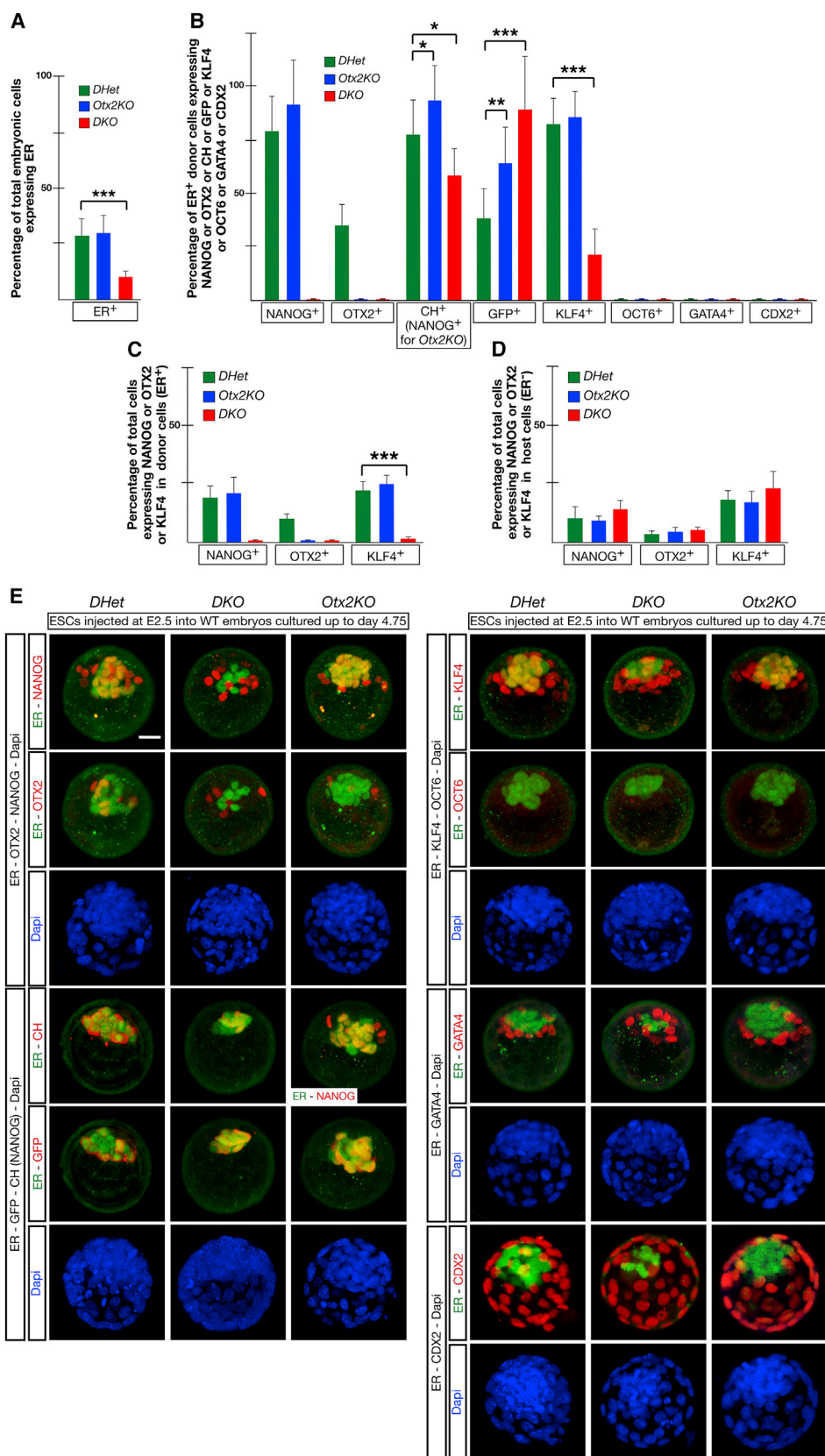
Finally, we tested whether the transition to the naive state was affected in mutant ESCs. In WT ESCs cultured in LIF + 2i, this transition was characterized by increased expression of naive markers such as OCT4, NANOG, KLF4, *Esrrb*, *Fgf4*, and *Rex1* and loss of primed markers such as OTX2 and OCT6 (Figures 7A, 7B, and S4E). At P1, WT, *DHet* (data not shown), and *Otx2KO* ESCs exhibited a stable naive-like state while *DKO* ESCs showed small OCT4⁺ colonies expressing KLF4, CH and, unexpectedly, GFP (Figures 7A, 7B, and S4E). Moreover, most of the OCT4⁻ *DKO* cells expressed PE (GATA4, KLF4, *Foxa2*, *Sox17*) or trophectoderm (TE) (CDX2, *Hand1*, *Gata3*) markers (Figures 7A, 7B, and S4E), which was reminiscent of *NanogKO* and *NanogKOS* ESCs in LIF + FBS. Nevertheless, under sustained culture in LIF + 2i, *DKO* ESCs gradually expanded the naive-like cell fraction and at P5 were similar to WT ESCs (Figures 7A–7C). We suggest that the phenotype of *DKO* ESCs may reflect the fact that, although NANOG is ablated, the simultaneous loss of OTX2 restrains but does not suppress naive conversion. To test this possibility, we first cultured *pPyCAGOttx2-ER* (-Tx) ESCs in LIF + 2i for two passages to allow conversion into the naive-like state and then administered Tx for a third

Figure 4. Analysis of *DKO* and *Otx2KO* Transcriptomes

(A–H) Venn diagrams showing the different categories of gene transcripts (boxes from (a) to (o) in each Venn diagram) up- or downregulated in *DKO* (A–D) and *Otx2KO* (E–H) ESCs in LIF + FBS when compared with WT ESCs (LIF + FBS) or WT ESCs (LIF + 2i) or WT EpiLCs (44 hr); this analysis allows us to identify *DKO* (LIF + FBS) gene transcripts shared with those up- or downregulated specifically in WT EpiLCs (44 hr) or WT ESCs (LIF + FBS) (red circle in box [j] in A–D); the same analysis was performed to identify *Otx2KO* ESCs (LIF + FBS) transcripts shared with those specifically expressed in WT ESCs (LIF + FBS) and WT ESCs (LIF + 2i) (red circle in box [j] in E–H).

(I–L) RT-PCR assays of selected genes up- or downregulated specifically in *DKO* (LIF + FBS) and WT EpiLCs (44 hr) (I), or in *DKO* (LIF + FBS) and WT ESCs (LIF + FBS) (J), or in *Otx2KO* (LIF + FBS) and WT ESCs (LIF + FBS) (K), or in *Otx2KO* (LIF + FBS) and WT ESCs (LIF + 2i) (L). The selected genes belong to the box (j) highlighted with red circles in the Venn diagrams (n = 3 independent experiments).

See also Figure S5 and Tables S3–S5.



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passage. Compared with untreated *pPyCAGOt2-ER* ESCs, Tx treatment generated hypomorphic OCT4⁺ colonies expressing NANOG and KLF4 and numerous OCT4⁻ cells showing neural specific markers SOX1 and TUJ1 (Figure 7D). RT-PCR assays showed that Tx administration downregulated *Esrrb*, *Rex1*, and *Fgf4* expression and activated strong *Sox1* expression (Figures 7E and S4F). Then, we analyzed *NanogKOS* ESCs cultured in LIF + 2i. We found that at P1, *NanogKOS* ESCs exhibited small OCT4⁺-CH⁺ colonies expressing KLF4 only in a few OCT4⁺ cells (Figure 7F). Noteworthy, as observed for GFP in *DKO* ESCs, OTX2 was partially suppressed also in *NanogKOS* ESCs. Furthermore, *NanogKOS* ESCs generated a relevant number of PE-like OCT4⁻ cells (Figures 7E, 7G, and S4G). At P5, in contrast with *DKO*, *NanogKOS* ESCs did not recover a naive-like phenotype (Figures 7E, 7G, and S4G). Together these and previous findings highlight the relevance of the antagonism between NANOG and OTX2 for the control of the heterogeneous identity of ESCs in LIF + FBS. Without integration of this mutual antagonism, the plasticity of ESCs is lost, and the capacity for bidirectional conversion into naive or primed pluripotency is forced in a more unidirectional manner either forward (primed) or reverse (naive), which is established by the dominance of the OTX2 regulatory network over that controlled by NANOG or vice versa. When both networks are abolished, ESCs retained a degree of plasticity sufficient to accomplish bidirectional conversion.

DISCUSSION

Heterogeneity in gene expression is a typical feature of ESCs cultured in LIF + FBS. Understanding the molecular basis controlling this heterogeneous condition is important to decipher the plasticity of ESCs (Smith, 2017; Torres-Padilla and Chambers, 2014; Martinez Arias et al., 2013; Nichols and Smith, 2009, 2011; Chambers et al., 2007). Although different mechanisms may operate together to determine the final state of ESCs, the control exerted by specific transcription factors may have a prominent role. In this context, allelic control of NANOG dosage has been proposed to have a causative role for the heteroge-

neity of ESCs cultured in LIF + FBS (Torres-Padilla and Chambers, 2014; Miyanari and Torres-Padilla, 2012). On the other hand, OTX2 exhibits heterogeneous expression in ESCs cultured in LIF + FBS and is required to promote the transition into the early primed state (Acampora et al., 2013, 2016; Buecker et al., 2014; Yang et al., 2014). These previous findings led us to investigate the possibility that OTX2 and NANOG may be antagonistic determinants that together specify the heterogeneity of ESCs cultured in LIF + FBS. We therefore studied how these two factors influence the identity and size of ESC sub-type compartments to assess whether they privilege or affect specific states of pluripotency co-existing in ESCs self-renewing in LIF + FBS. To this aim, we analyzed an allelic series of mutant ESCs carrying different dosages of OTX2 and/or NANOG. We have shown that, without the addition of exogenous factors that promote naive or primed pluripotency, loss of *Otx2* causes expansion of the naive-like compartment, which is consistent with previous data on *Nanog* overexpression (Chambers et al., 2003). In contrast, loss of *Nanog* analyzed in *NanogKO* ESCs or *Otx2* overexpression generate a strikingly similar phenotype characterized by remarkable contraction of the naive-like compartment and expansion of the primed-like sub-type. When both genes are deleted, ESCs acquire a pluripotent state exhibiting identity features in common with early primed-like cells. These data indicate that antagonism between NANOG and OTX2 is an essential requirement to specify and maintain ESC heterogeneity, which we visualize through the analysis of OCT4⁺ cells co-expressing different combinations of NANOG, OTX2, and OCT6 (Acampora et al., 2016). We envisage that between naive-like and primed-like compartments, there are at least three transitional and dynamic compartments—the pre-naive-like, the pre-primed-like, and the unassigned—ready to be converted and stabilized into naive or primed pluripotent states. Therefore, in addition to previous studies indicating that NANOG and OTX2 are important coordinators of naive and primed pluripotency, respectively (Acampora et al., 2013, 2016; Buecker et al., 2014; Yang et al., 2014; Silva et al., 2009; Chambers et al., 2003, 2007; Mitsui et al., 2003), our new data suggest that the combined action of these two transcription factors

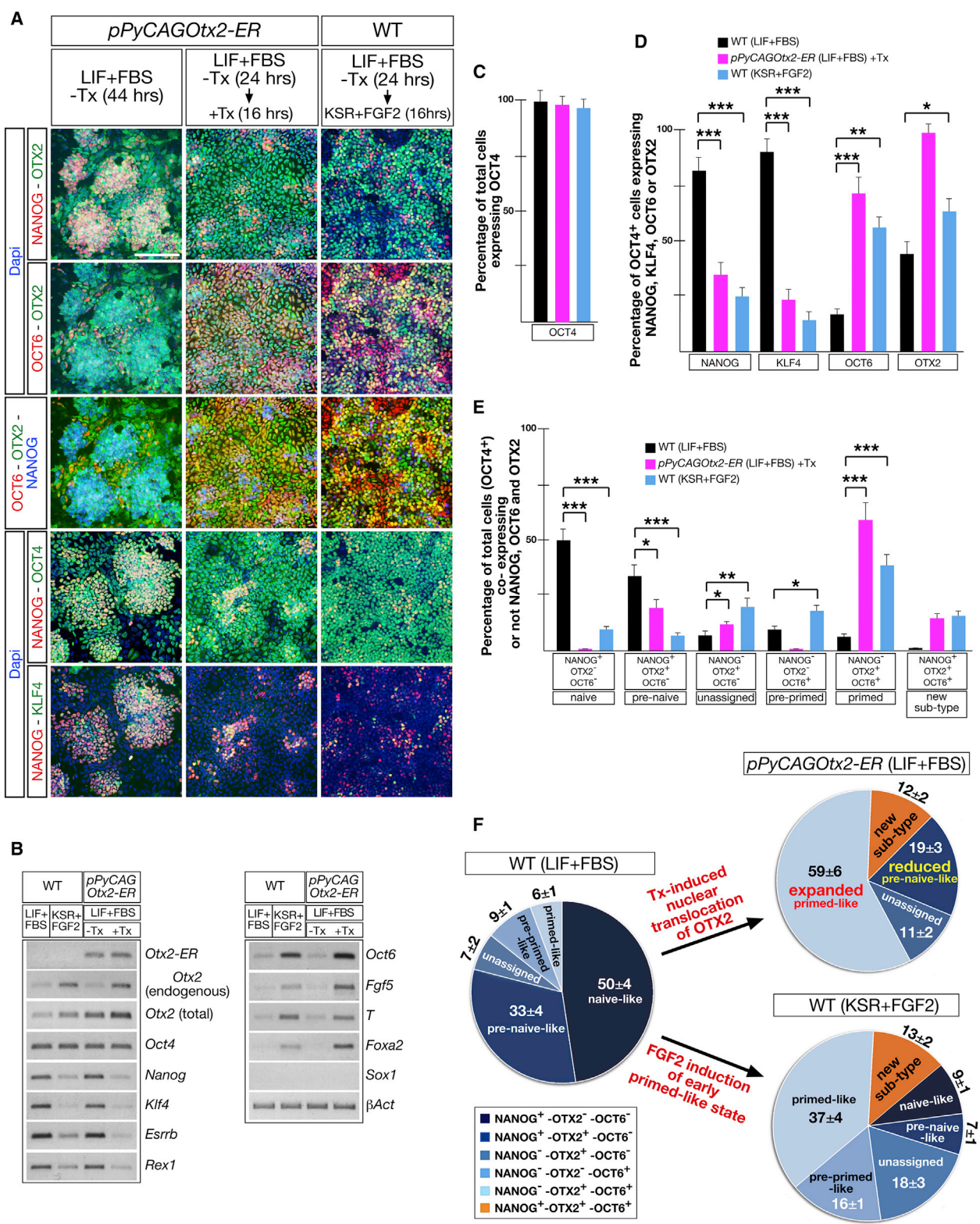
Figure 5. Abnormalities of Mutant ESCs in Chimeric Blastocysts

(A) Graphic representation showing that, compared with *DHet* and *Otx2KO*, *DKO* ESCs exhibit reduced efficiency of integration into host embryos. Data are the means \pm SD from $n = 10$ blastocysts for each ESC line; *** $p \ll 0.005$.

(B–D) Cell counting showing the percentage of ER⁺ donor cells expressing NANOG, OTX2, CH, GFP, KLF4, OCT6, GATA4, and CDX2 (B) or the percentage of total cells (DAPI⁺) expressing NANOG, OTX2, or KLF4 in the donor (ER⁺) (C) or in the host cells (ER⁻) (D). Data are the means \pm SD from $n = 7$ blastocysts for each combination of antibodies reported in (E) and for each ESC line. * $p < 0.05$; ** $p < 0.01$; *** $p < 0.005$.

(E) Representative confocal images of embryos injected at E2.5 with *DHet*, *DKO*, and *Otx2KO* ESCs and immunostained at E4.75 with ER plus NANOG and OTX2, or CH and GFP, or KLF4 and OCT6, or GATA4, or CDX2. Note that the externalmost plane of acquisition is removed to allow a better view of the inner cell mass. DAPI staining is also shown. Scale bar, 30 μ m.

See also Figure S6 and Table S6.





is instrumental in determining the heterogeneous identity of ESCs cultured in LIF + FBS. Importantly, a similar heterogeneity can be identified *in vivo* between E4.5 and E4.7 when the naive pluripotency circuit is being shut down but before the early primed state becomes dominant. Most of the ESC sub-types detected in LIF + FBS cultures are present in the E4.5–4.7 epiblast (Acampora et al., 2016). We suggest that at this stage, the transitional heterogeneity of the epiblast is captured in cultured ESCs by LIF + FBS and is maintained by antagonism between OTX2 and NANOG. These observations are relevant to the recently elaborated concept of formative pluripotency, a state hypothesized to interpose between naive and primed pluripotency and suggested to occur abruptly (Smith, 2017). However, the finding that the epiblast of E4.5–4.7 embryos and ESCs in LIF + FBS exhibit a spectrum of distinct and precise combinations of naive and primed markers leads us to suggest that this transition may not necessarily occur abruptly. In this context, an imbalance in favor of the OTX2-dependent regulatory network due to downregulation of *Nanog* expression and upregulation of *Otx2* might trigger conversion of pluripotency from a metastable and heterogeneous condition toward a formative state.

The co-existence of different pluripotent sub-types is an intermediate gate preceding the unidirectional transition of the epiblast into the early primed state *in vivo*, and allowing bidirectional conversion into naive or early primed state *in vitro* (Smith, 2017; Torres-Padilla and Chambers, 2014; Martínez Arias et al., 2013; Nichols and Smith, 2011; Silva et al., 2009; Ying et al., 2008). The capability of NANOG to impose a unimodal naive-like identity is counterbalanced by OTX2, which, in turn, promotes primed-like identity. We propose that integration of these opposing regulatory circuits underlies the mechanism that specifies heterogeneity in ESCs. This identity can be established only in LIF-containing medium where exogenous signal constraints (FGF or 2i) are minimized and ESCs are allowed to manifest their flexible nature. Signaling pathway activation perturbs this state and determines the dominance of one of the two regulatory networks. This process requires the contemporary and co-ordinated response by both NANOG and OTX2. Indeed, 2i-mediated induction of naive state could not be efficiently accom-

plished without *Nanog* activation and *Otx2* repression; similarly, FGF-mediated induction of primed state is incomplete without *Otx2* activation and *Nanog* downregulation. Specification and maintenance of this mechanism require that the dosage and expression pattern of OTX2 and NANOG should be established within precise parameters, which ultimately determine the degree of integration between their antagonistic transcriptional networks. According to a hierarchically high role of OTX2 and NANOG in interpreting and responding to primed and naive inducers, we report that transitions between naive and primed compartments can be coherently mimicked in LIF + FBS only by genetic manipulations of OTX2 and/or NANOG dosage.

EXPERIMENTAL PROCEDURES

Generation of ESC Lines

All mutant ESC lines (Figure S1) were generated in an E14Tg2a-derived ESC line, whose *Rosa26* locus was previously targeted with a *CreER-pac* gene cassette (Acampora et al., 2013). The sequential steps for the generation of mutant ESCs are described in the Supplemental Experimental Procedures.

Cell-Culture Experiments

ESC culture procedures are described in the Supplemental Experimental Procedures.

RT-PCR and Western Blotting

RT-PCRs were performed in non-saturating conditions using the primers and cycles listed in Table S7. Western blots were probed with antibodies and at the dilutions listed in Table S7.

FACS Analysis

*Nanog*KO ESCs were trypsinized, centrifuged, and resuspended in ESC medium containing 2% FBS and 2 mM EDTA. The CH⁺ fraction was sorted using a BD FACSAria III (Becton Dickinson) into a 6-well plate. The data were analyzed using the BD FACSDiva software.

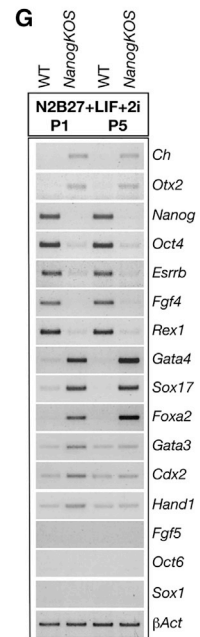
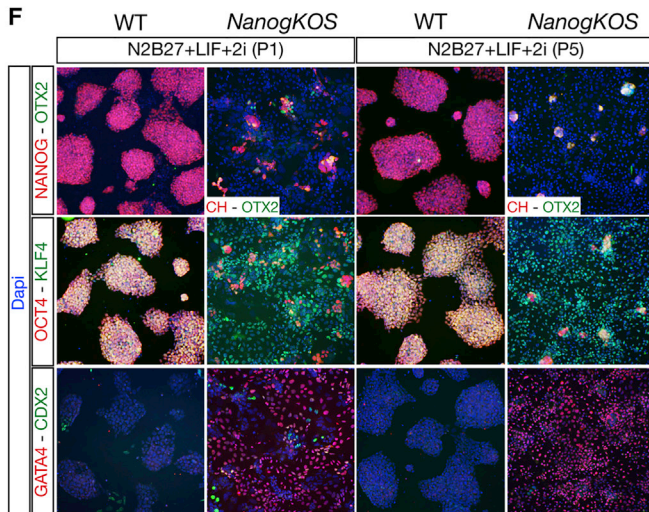
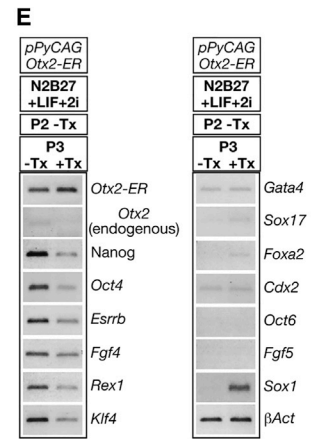
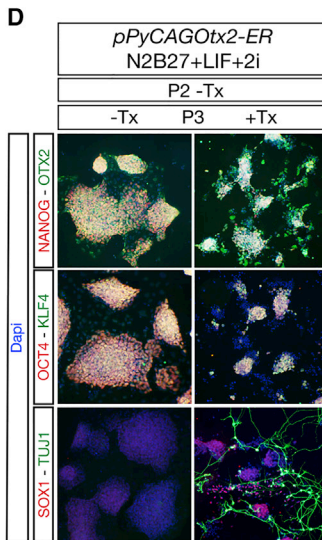
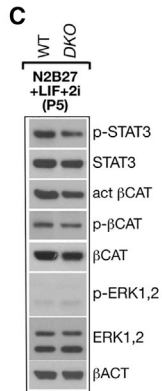
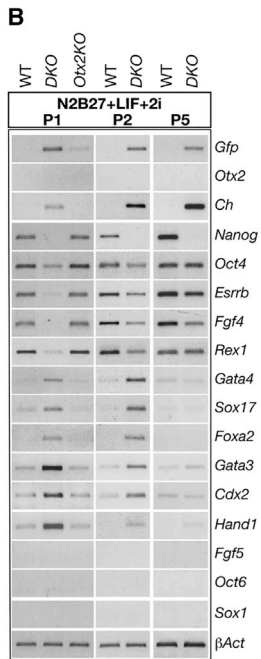
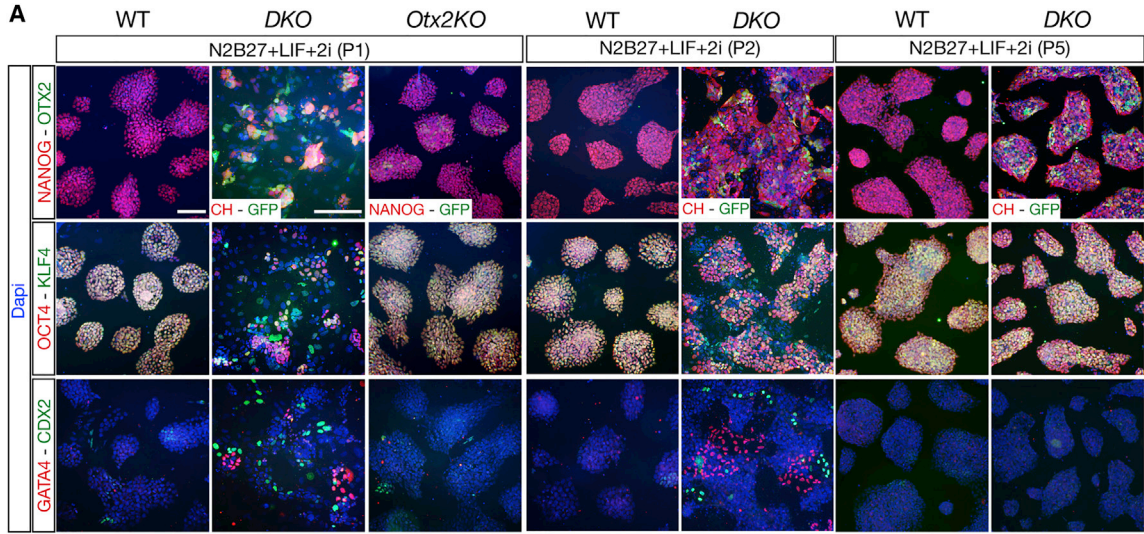
Immunohistochemistry

Immunohistochemistry assays on ESCs and embryos and ALP assays were performed as described (Acampora et al., 2013). Antibodies and dilutions used in this study are listed in Table S7.

(B) Representative RT-PCR assays show that in WT and pPyCAG0tx2-ER ESCs, FGF2 treatment and Tx-induced nuclear translocation of OTX2-ER downregulate *Nanog*, *Klf4*, *Esrrb*, and *Rex1* and upregulate endogenous *Otx2*, *Fgf5*, *T*, and *Foxa2* ($n = 3$ independent experiments). (C–E) Cell counting showing for WT ESCs (LIF + FBS), Tx-treated pPyCAG0tx2-ER in LIF + FBS, and WT ESCs in KSR plus FGF2 the percentage of total cells expressing OCT4 (C), the percentage of OCT4⁺ cells expressing NANOG or KLF4 or OCT6 or OTX2 (D), and the percentage of total cells (OCT4⁺) expressing the different sub-types identified by the combinatorial analysis of NANOG, OCT6, and OTX2 (E). Data are presented as means \pm SD from four independent experiments. *** $p \ll 0.001$; ** $p < 0.001$; * p value is between 0.005 and 0.001.

(F) Cell-counting data show that nuclear translocation of OTX2-ER is sufficient to induce a re-organization of sub-type compartments similar to that induced by FGF2 in WT ESCs.

See also Figures S4 and S7 and Table S2.



(legend on next page)



RNA-Seq Experiments, Analysis of RNA-Seq Data, and RNA-Seq Validation Assays

RNA-seq experiments were performed on DKO ESCs (LIF + FBS), Otx2KO ESCs (LIF + FBS), WT ESCs (LIF + FBS), WT ESCs (LIF + 2i), and WT EpiLCs (44 hr). Indexed sequencing libraries were generated and analyzed as described in the [Supplemental Experimental Procedures](#).

Chimerism Experiments

Chimerism experiments are described in the [Supplemental Experimental Procedures](#). Animals were handled in accordance with the authorization 1196/2015-PR released by the Italian Ministry of Health.

Cell-Counting Experiments

Cell counting of ESC sub-types was manually performed on immunohistochemistry images printed in A4 format ([Acampora et al., 2013, 2016](#)). SD was calculated from the analysis of four independent experiments. p values were determined using the one-tailed Student's t test. Cell counting of chimeric blastocysts was performed on n = 10 blastocysts for colonization efficiency experiments and on n = 7 blastocysts for each combination of antibodies and for each injected ESC line ([Table S6](#)). Images were printed in A4 format for manual cell counting. Data are reported as means ± SD; p values were determined using the one-tailed Student's t test. For details see [Supplemental Experimental Procedures](#).

ACCESSION NUMBERS

RNA-seq data are available in European Nucleotide Archive database (<https://www.ebi.ac.uk/ena/submit/sra/#home>) under accession number ENA: PRJEB19241/ERP021225.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, seven figures, and seven tables and can be found

with this article online at <https://doi.org/10.1016/j.stemcr.2017.09.019>.

AUTHOR CONTRIBUTIONS

D.A. performed and supervised most of the *in vitro* and *in vivo* experiments. L.G.D.G. generated targeting molecules and contributed to the generation and analysis of mutant ESCs. A.G. and V.N. performed and analyzed RNA-seq experiments. D.O. generated the pPyCAGOtx2-ER ESC line. J.Z. generated the pPyCAGOtx2-ER plasmid. A.L. contributed to the generation of mutant ESCs. I.C. contributed to the preparation of the manuscript. A.S. conceived the experiments and wrote the manuscript.

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Figure 7. Conversion of Mutant ESC Lines into Naive-like and Primed-like Pluripotent States

- (A) Representative immunohistochemistry assays with OTX2 or GFP and NANOG or CH, OCT4 and KLF4, and with GATA4 and CDX2 performed at passage (P) 1, P2, and P5 in WT, Otx2KO, and DKO ESCs cultured in LIF + 2i show that DKO ESC colonies acquire a domed morphology, ubiquitous expression of OCT4, KLF4, and CH and stop generating OCT4⁺ cells. Scale bars, 100 μm.
- (B) RT-PCR assays show that in DKO ESCs, the expression of naive, PE, and TE markers progressively recovers a profile similar to that of WT ESCs (n = 3 independent experiments).
- (C) Western blots show that at P5 also the level of p-STAT3, p-ERK1,2, act-βCAT, and p-βCAT is similar in WT and DKO ESCs (n = 3 independent experiments).
- (D) Representative immunohistochemistry assays with OTX2 and NANOG, OCT4 and KLF4, and SOX1 and TUJ1 on pPyCAGOtx2-ER ESCs cultured in LIF + 2i up to P3 without Tx or cultured up to P2 without Tx and for an additional passage (P3) with Tx.
- (E) RT-PCR assays of Tx-treated pPyCAGOtx2-ER ESCs show reduced expression of Esrrb, Fgf4, and Rex1 and Sox1 upregulation; expression of PE, TE, and primed markers appear unaffected (n = 3 independent experiments).
- (F) Representative immunohistochemistry assays with OTX2 and NANOG or CH, OCT4, and KLF4 and with GATA4 and CDX2 show that NanogKOS ESCs cultured in LIF + 2i exhibit small and rare OCT4⁺ colonies, abundant differentiation of PE-like cells expressing KLF4 and GATA4, and do not recover a naive-like phenotype at P5.
- (G) RT-PCR assays performed on WT and NanogKOS ESCs cultured in LIF + 2i show that naive markers are downregulated, PE markers are stably upregulated, TE markers are transiently upregulated, and primed markers are unaffected (n = 3 independent experiments). See also [Figure S4](#).



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