Live analysis and function of Myc-mediated cell competition in mouse pluripotent stem cells

Doctoral thesis

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Director: Dr. Miguel Torres Sánchez
À la plus folle. JTA.
La duda es uno de los nombres de la inteligencia

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In the early mouse embryo and in embryonic stem cell (ESC) cultures, the transcription factor Myc exhibits a cell-to-cell heterogeneous pattern. Cells expressing low levels of Myc are eliminated from the population by cell competition. Myc has been reported to promote cell reprogramming to pluripotency and to regulate cell anabolism and proliferation in ESCs; however, the biological role of Myc-dependent endogenous cell competition and the dynamics and regulation of Myc during this process remain unknown.

Here we develop a new image analysis tool that allows us to track the temporal evolution of endogenous Myc levels, perform neighborhood analysis in ESC cultures and generate 3D+t computerized data. We show that despite Myc degradation and resynthesis during mitosis, Myc levels are mostly heritable in ESC lineages. Cell competition results from random interactions between cells with high discrepancies in Myc levels. Myc-low cells (“losers”) temporally integrate contacts with Myc-high cells (“winners”), which leads to a progressive decrease in their own Myc levels until dying. Interestingly, endogenous Myc levels correlate with the pluripotency status; differentiation-primed cells express low Myc levels and are outcompeted by Myc-high naive ESCs. Indeed, cell competition inhibition results in an accumulation of primed cells. These observations in ESCs correlate with findings in the mouse epiblast. Moreover, we show that Myc levels directly determine the competitive ability of ESCs irrespective of the pluripotency status.

Our results identify Myc as a mediator between differentiation status and competitive ability of pluripotent cells. Myc-driven endogenous cell competition thus acts as a mechanism to protect pluripotent stem cell pools from differentiation.
Tanto en el embrión temprano de ratón como en cultivos de células madre embrionarias (CMEs), el factor de transcripción Myc muestra un patrón de expresión heterogéneo. Las células que expresan menos niveles de Myc son eliminadas de la población mediante competición celular. Se ha demostrado que Myc promueve la programación celular hacia un estado de pluripotencia y regula el anabolismo y la proliferación en CMEs; sin embargo, la función biológica de la competición celular endógena dependiente de Myc, así como la regulación y dinámica de la expresión este gen durante este proceso aún se desconocen.

En esta tesis desarrollamos una novedosa herramienta de análisis de imagen, a partir de la cual realizamos análisis de células individuales y sus vecindarios y generamos un sistema de datos 3D+t computarizados. De esta manera demostramos que, a pesar de que Myc se degrada y se vuelve a sintetizar rápidamente durante la mitosis, sus niveles de expresión son en su mayor parte heredables. Además, la competición celular se desencadena por interacciones aleatorias entre células con grandes diferencias en sus niveles de Myc; de forma que las células que poseen bajos niveles de Myc (“perdedoras”) integran en el tiempo contactos con células con mayor expresión (“ganadoras”) lo que lleva a una reducción progresiva en sus propios niveles hasta que mueren. Cabe resaltar que los niveles endógenos de Myc correlacionan con el estado de pluripotencia; las células en un estado más avanzado de diferenciación expresan menos Myc siendo eliminadas por células más indiferenciadas y con altos niveles de expresión. De hecho, la inhibición de la competición celular produce una acumulación de células en proceso de diferenciación. Estas observaciones correlacionan con resultados obtenidos en estudios realizados en epiblasto de ratón. Además, demostramos que la eficiencia competitiva viene directamente determinada por los niveles de Myc independientemente del estado de pluripotencia.

Estos resultados señalan a Myc como un intermediario entre el estado de diferenciación y la capacidad competitiva de las células pluripotentes. Por lo tanto, la competición celular endógena promovida por Myc funciona como un mecanismo de protección de poblaciones de células madre pluripotentes frente a la diferenciación.
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Abbreviations

2i: 2 inhibitors
5mC: DNA methylation
BMP: Bone Morphogenetic Protein
Bst: Belly spot and tail
Cdk4: Cyclin-Dependent Kinase4
CHIRON: CHIR99021 (inhibitor)
Dgl: Discs Large
Dpp: Decapentaplegic
ECFP: Enhanced Cyan Fluorescent Protein
EDAC: Epithelial Defence Against Cancer
EMT: Epithelial-Mesenchymal Transformation
EPC: ectoplacental cone
EpiLCs: Epiblast-like Cells
EpiSCs: Epiblast Stem Cells
ERK: Extracellular Regulated Kinase
ESC: Embryonic Stem Cell
ExE: Extraembryonic Ectoderm
EYFP: Enhanced Yellow Fluorescent Protein
FGF: Fibroblast Growth Factor
GSC: Germline/Spermatogonial Stem Cell
GSK: Glycogen Synthase Kinase 3
ICM: Inner Cell Mass
iMOS: Inducible Mosaic
iPSC: induced Pluripotent Stem Cell
JAK: Janus Kinase
JNK: c-Jun N-terminal Kinase
Klf4: Kruppel Like Factor 4
KO: Knockout
Lgl: Lethal Giant Larvae
LIF: Leukemia Inhibitory Factor
MAPK: Mitogen-Activated Protein Kinase
MDCK: Madin-Darby Canine Kidney
MEF: Mouse Embryonic Fibroblast
MHN: Myc-Highest Neighbour
ML-NDC: Myc-Low Non-Dying Cell
Myc-H: Myc-High
Myc-L: Myc Low
Myc-M: Myc-Medium
Myc: Myelocytomatosis oncogene
Mycel: v-myc avian myelocytomatosis viral oncogene lung carcinoma derived
Mycn: v-myc avian myelocytomatosis viral related oncogene, neuroblastoma derived
NDC: Non-Dying Cell
PD03: PD0325901 (inhibitor)
PDC: Prospective-Dying Cell
PE: Primitive Endoderm
PGC: Primordial Germ Cell
PI3K: Phosphatidylinositol 3-Kinase
RpL24: Ribosomal Protein L24
Sox2: SRY (sex determining region Y)-box 2
SPARC: Secreted Protein Acidic Cysteine-Rich
SSL: Spline Smooth Line
STAT: Signal Transducer and Activator of Transcription
SWH: Salvador/Warts/Hippo
TE: Trophectoderm
TET: Ten-eleven translocation
TFCP2l1: Transcription Factor CP2 like1
TGC: Trophoblast Giant Cell
TGFb: Transforming Growth Factor beta
TS: TE stem
VprBP: Vpr (HIV-1) binding protein
Vps25: Vacuolar Protein-Sorting-Associated protein 25
WT: Wild Type
INTRODUCTION
Cell competition consists in the fitness comparison between viable neighbouring cells resulting in the elimination of the less fit cells (Figure 1). This phenomenon was firstly described in *Drosophila* and currently has been characterized in many biological scenarios and is envisioned as an important tissue homeostatic mechanism in metazoans.

In mammals, pluripotent cells of the early mammalian embryo show a natural cell-to-cell heterogeneity in Myc levels, which leads to the competitive elimination of Myc-low cells from the embryonic lineage. The mechanism and the biological relevance of this phenomenon, however, remain unknown. In this thesis, we aim to elucidate the mechanisms and biological basis of endogenous cell competition regulated by Myc in the context of Embryonic Stem Cells (ESCs) and the early mammalian embryo.

1. Cell competition

1.1. The discovery of Cell Competition: early studies in flies

Forty years ago, Ginés Morata and Pedro Ripoll reported the phenomenon while studying the proliferation of *Drosophila* cells mutant for *Minute* genes, in which ribosomal proteins are affected. *Minute* mutations affect protein synthesis ability and produce cellular lethality in homozygosis, while heterozygous flies are viable and nearly normal but show slower development due to a reduced proliferative rate. When a mosaic of heterozygous (*M/+*) and wild-type (WT) cells were induced in wing imaginal discs, *M/+* cells, which otherwise would be viable, were eliminated due to confrontation with WT cells (Morata and Ripoll, 1975) (Figure 1). Simpson confirmed the previous observations and observed that elimination of *M/+* clones in a WT background did not take place when larvae were starved. Since starvation induces generally reduced growth rates, these experiments strongly suggested that a limitation of growth/proliferation of *M/+* determines the competitive elimination of *M/+* cells and that the mechanism underlying *M/+* cell depletion was related to the metabolic control of cell growth (Simpson, 1979). The requirement of a differential growth rate for the elimination of *M/+* cells was later confirmed by the confrontation
of Minute mutations with variable severity (Simpson and Morata, 1981). In these models, competition did not affect the final size of the wings or compartments, suggesting the compensatory proliferation of WT cells. Moreover, cell competition was shown by these studies to result from local interactions between slow- and faster-growing cells (Simpson and Morata, 1981). However, cell competition was ineffective across compartment borders and WT clone expansion was restricted to well-defined and reproducible frontiers, which suggests that wing disc are divided in non miscible cell populations (compartment boundary) (Simpson and Morata, 1981).

It was not until many years later that cell competition became again a subject of interest when the phenomenon was reported to require short-range interactions and to be executed by apoptosis of loser cells in Drosophila imaginal discs (Abrams, 2002; Milan et al., 2002; Moreno et al., 2002).
1.2. Supercompetition and the transcription factor Myc

Supercompetition is as a type of cell competition in which cells with experimentally increased fitness eliminate neighbouring WT cells, thereby overcolonising tissues (Abrams, 2002) (Figure 1). The first gene to be linked to this definition was the proto-oncogene dMyc. dMyc is the fly homologue of the mammalian family of Myc transcription factors and regulates many downstream targets involved in cell growth and cell anabolism (Johnston et al., 1999; de la Cova and Johnston, 2006). In 2004 two pioneer papers presented the role of this proto-oncogene in cell competition by inducing dMyc overexpressing clones in the wing disc (de la Cova et al., 2004; Moreno and Basler, 2004). These clones were able to expand at the expense of WT cells until they filled the compartment (Figure 1). Apoptosis via JNK (Moreno and Basler, 2004) and hid activation in the loser cells (de la Cova et al., 2004) was required in this process. In this context, dMyc-overexpressing cells acted as “supercompetitors” and the WT cells behaved as “losers” (Figure 1). The discovery of supercompetition induced by Myc led to propose that early steps of tumour formation could be driven by this phenomenon (Rhiner and Moreno, 2009), which generated a new wave of interest in cell competition.

On the contrary, downregulation of dMyc in cells led to their elimination, in similarity to observations with M/+ clones (Johnston et al., 1999; de la Cova et al., 2004; Moreno and Basler, 2004). This suggested that dMyc was a general driver of cell competition by controlling cell’s anabolic machinery. Interestingly, cells with two extra dMyc copies, which behave as supercompetitors in front of WT cells, instead behaved as “losers” when confronted with cells with four extra copies of dMyc (Moreno and Basler, 2004) supporting the idea that competition is based on intercellular comparison of relative Myc levels and not on cell-autonomous sensing of absolute Myc level.

In addition, these studies ruled out a strict relationship between cell growth or proliferative activity and competitive ability, since overexpression of factors inducing cell growth (PI3K Dp110) or cell cycle regulators (CyclinD and Cdk4) generated large clones but were unable to eliminate surrounding WT cells, therefore Myc and Minute
and no other proliferation/growth regulators affect specific competitive pathways (de la Cova et al., 2004; Moreno and Basler, 2004).

1.3. Pathways involved in cell competition in *Drosophila*

After the discovery of *Minute* and Myc-driven competition, many studies have tried to shed light on this field by finding out new pathways and molecules involved in this process. These factors can be classified into two groups depending of the kind of cell competition they induce (Figure 2).
The first group includes factors that affect metabolic pathways and are likely to induce cell competition by producing contrasting anabolic ability in neighbouring cells (Figure 2). This group includes the classical factors inducing cell competition, like *Minute* mutations, the BMP/Dpp pathway and Myc. This group also includes other pathways involved in growth/survival signal reception or transduction to the cell’s anabolic machinery, such as the Rab5 and Vps25 endocytic pathways (Moreno and Basler, 2004; Herz *et al.*, 2006), the Ras/MAPK pathway (Prober and Edgar, 2000) and the JAK/STAT pathway (Rodrigues *et al.*, 2012). In addition, cell competition can also be induced by mutations in the Salvador/Warts/Hippo pathway, which is a central regulator of growth, cell death and tissue homeostasis (Tyler *et al.*, 2007; Neto-Silva *et al.*, 2010; Ziosi *et al.*, 2010; Chen *et al.*, 2012). The nature of competitiveness regulation by these pathways is likely similar, since hierarchical regulatory interactions have been established between the various pathways involved in metabolic cell competition. Myc acts downstream of various cell competition-inducing pathways like the BMP/Dpp (Doumpas *et al.*, 2013), Ras/MAPK (Prober

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**Figure 2. Pathways involved in metabolic and structural competitive fitness.**

Schematic of the main pathways involved in structural and metabolic cell competition. Cell selection is initiated by mutations or pathways that generate differential fitness activity between neighbouring cells triggering cell competition. On one hand, metabolic fitness is determined by the anabolic activity of the cell, specifically by protein synthesis machinery. This machinery is activated by Myc, on which cellular growth and proliferative pathways converge. Even though the JAK/STAT pathway does not depend on Myc signalling, it also affects cell metabolic fitness. On the other hand, structural fitness is determined by the apico-basal polarity machinery. When polarity pathways are mutated the structural fitness is decreased and the SWH pathway is repressed leading to an increase in growth/proliferation. Since competitive behaviours between isolated mutant and surrounding WT cells counteract SWH pathway repression by JNK, polarity-deficient cells are thus outcompeted. However, when polarity-deficient cells elude competition, SWH pathway is effective and trigger overgrowth of mutant cells. The Notch pathway and the endocytic pathway (Rab5, Vps25) may interact with both metabolic and structural fitness pathways, but their specific involvement has not been reported yet. Ellipses in red represent factors whose gain of function increases fitness and blue ellipses represent those factors whose gain of function reduce fitness. Filled arrows indicate activation, whereas bars mean inhibition or blockade of target activity. Dashed lines show a deduced effect and a solid line indicates a known downstream target. (Based on (Claveria and Torres, 2016)).
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and Edgar, 2000; Moreno and Basler, 2004) and SWH (Neto-Silva et al., 2010; Ziosi et al., 2010). Moreover, BMP/Dpp signal transduction is enhanced by a positive feedback that involves Myc and enhanced protein synthesis is required downstream of Myc to induce cell competition (Moreno and Basler, 2004). Although the STAT pathway does not show any apparent epistatic relationship during cell competition with the Myc, SWH, or Minute pathways (Rodrigues et al., 2012), it seems to affect the same cellular parameters, because Myc-low cells are rescued by STAT hyperactivation and STAT-deficient cells are rescued when confronted with Minute mutant cells (Rodrigues et al., 2012).

The second group of cell competition inducers includes factors affecting polarity and epithelial integrity and therefore define a different class of competitive interactions known as structural cell competition (Figure 2). Extrusion from the epithelium is the typical way of eliminating cells in this type of competition and cell death is not always involved in this process. Structural cell competition is induced by alterations affecting components of the canonical pathway regulating apico-basal polarity, such as Lgl (Menendez et al., 2010; Tamori et al., 2010), Scribble (Brumby and Richardson, 2003), Dlg (Woods and Bryant, 1991), Mahjong/VprBP (Tamori et al., 2010) and Crumbs (Hafezi et al., 2012). In addition, the role of Rab5 and Vps25 in cell competition may also be related to the structural organization of the cell, since the endocytic pathway also affects apico-basal polarity (Lu and Bilder, 2005). Yet, although there is also cross talk between structural and metabolic cell competition, these pathways have very different natures. In fact, in various cases, the mutations that induce the loser phenotype in structural competition, however promote metabolic changes typical of winner cells in metabolic competition, indicating that structural alterations in epithelial tissues overrule metabolic activity in competitive interactions. An additional difference is that both winner and loser cells in metabolic competition usually respect normal tissue architecture, while winners or losers in structural cell competition can produce tumorous growth.

Finally, there are some other pathways that induce cell competition but whose mechanisms and relationships to the canonical pathways have not been completely resolved, such as the Wnt pathway (Vincent et al., 2011; Rodrigues et al., 2012) or patterning mismatches (Adachi-Yamada et al., 1999; Milan et al., 2002).
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1.4. Cell competition in mammals

The first study suggesting that cell competition could also take place in other metazoans than insects was the analysis of a mouse mutation equivalent to the fly Minute mutants. The Bst mice mutants are defective in ribosomal protein L24. While Bst homozygous mutant mice are not viable, heterozygous mice showed a kinked tail, decreased pigmentation and retinal defect but are viable and normal-sized (Oliver et al., 2004). As in the Drosophila system, cells heterozygous for RpL24 were disproportionately disadvantaged with respect to WT cells in mouse chimeras (Oliver et al., 2004). However, whether this observation was due to cell autonomous features of the Bst mutants or indeed reflects competitive interactions has not been addressed.

A second study pointed out to the occurrence of cell competition in liver regeneration. In aged rat livers, host hepatocytes were outcompeted by transplanted fetal hepatocytes, apparently by local interactions that promoted resident hepatocyte death, which facilitated long-term repopulation post-transplantation (Oertel et al., 2006).

Following these pioneering studies, cell competition based on stress and mediated by p53 was described (Figure 2). Stress signals induce p53-mediated cell death in Drosophila and either cell death or cell cycle arrest and repair in mammals. In bone marrow transplants, cells sublethally irradiated behaved as losers in a p53-dependent manner when confronted with non-irradiated cells (Bondar and Medzhito, 2010). In addition, when a mosaic p53 mutant bone marrow is irradiated only p53-deficient clones were recovered (Marusyk et al., 2010). In these two studies, a senescence-like phenotype, rather than apoptotic elimination, was induced in the outcompeted cells (Bondar and Medzhito, 2010; Marusyk et al., 2010). In this context, cell competition would be the result of cells comparing their p53 activity levels, resulting in the outcompetition of the p53-high population. Given the role of p53 in transducing stress situations, this mechanism would ensure the elimination of stressed cells whenever unstressed cells are available in the surroundings.

Epithelial monolayers in culture derived from the Madin-Darby canine kidney (MDCK) have also emerged as an important in vitro system for studying cell
competition in mammalian cells. Studies with structural cell competition pathways have shown that mutations in *Lgl*, its partner *Mahjong* (Tamori et al., 2010) or *Scribble* (Norman et al., 2012) induce cell competition in the MDCK culture system.

**Figure 3. Schematic representation of the iMOS system construction.**

**A.** A recombinase-based system for genetic mosaic induction was obtained by using two different *loxP* sequence variants that recombine with homotypic but not heterotypic sequences. The three cassettes flanked by the *loxP* sites end in a triple polyadenylation signal blocking the expression of the following transcriptional units. Upon Cre expression, either the transcriptional unit T0 is excised, leading to the expression of T1 or both T0 and T1 are eliminated leading to the expression T2. This system allows to randomly generate a mosaic with two labelled cell populations. The construct is targeted to the ubiquitous Rosa26 locus, whose expression gives rise to a policistronic mRNA expressing a buffer sequence or a protein of interest and a fluorescent reporter protein, either the yellow fluorescence protein (EYFP) or the cyan fluorescent protein (ECFP). The recombination frequency of EYFP cells is 75% and that of the ECFP cells 25%. This difference is due to the different distances between the *loxP* sites. **B.** Schematic representation of the iMOS lines used in this thesis. In iMOS\(^{WT}\), both labelled population are WT. iMOS\(^{T1-Myc}\) generates an EYFP Myc-overexpressing cell population and an ECFP WT population. In iMOS\(^{T2-p35}\), EYFP cells are WT while ECFP cells express the baculoviral apoptosis inhibitor p35. (Based on (Claveria et al., 2013)).
This model also showed the so-called Epithelial Defence Against Cancer (EDAC) which consists in the competitive elimination of cell carrying oncogenic mutations, like Rasv12 or Src, by apical extrusion from the epithelial sheet (Hogan et al., 2009; Kajita et al., 2010). In this model maintenance of epithelial homeostasis only occurs when WT cells win, whereas when the defective cells escape from cell competition, they give rise to tumour formation (Hogan et al., 2009; Kajita et al., 2010).

Additional cases of cell competition have been recently described when cells lacking the Notch pathway are confronted with WT cells in the adult mouse esophageal epithelium (Alcolea et al., 2014) (Figure 2). WT cells were progressively eliminated from the tissue by their Notch-deficient neighbour clones, which underwent overgrowth (Alcolea et al., 2014).

Furthermore, recent studies have shown that Myc-dependent cell competition also takes place in the early mouse embryo. Myc-induced supercompetition was characterised in mouse embryos by using a system of inducible mosaics based on Cre recombination, which generated two different populations (Claveria et al., 2013) (Figure 3). One cell population overexpressed Myc under the expression of the ubiquitous Rosa26 promoter and the other population remained WT and was selectively eliminated from the epiblast (Claveria et al., 2013). This demonstrated that supercompetition could also be induced in the embryo and that it provoked the phenotypically silent replacement of WT cells. In addition, it was shown that apoptosis of WT loser cells was needed for this replacement and that competition was contact-dependent, was apoptotic rate was increased in WT cells in direct contact with Myc overexpressing cells (Claveria et al., 2013). In this study cell competition was also explored in vitro in mouse ESCs, where Myc induced cell competition could also be shown (Claveria et al., 2013). In a similar study, it was reported that Bmpr1a (BMP receptor 1a) knock out ESCs decreased their Myc levels when compared with their WT counterparts and were subsequently outcompeted (Sancho et al., 2013). Interestingly, it was also reported that Myc activation led to induced cell competition in several tissues and organs of the developing and adult mouse (Claveria et al., 2013; Villa del Campo et al., 2014).
All these assays performed in mammalian models demonstrated that cell competition is widespread in metazoans and not restricted to just *Drosophila* wing discs or Minute mutations. In addition, they identify the Myc pathway as a conserved central regulator of cell competition.

### 1.5. Endogenous cell competition

All of the mentioned findings so far rely on experimental introduction of stress or mutations and did not address the potential roles of spontaneous cell competition in normal development or tissue homeostasis. Endogenous cell competition however has been detected in the mouse epiblast, where cells express naturally heterogeneous levels of Myc (Claveria *et al.*, 2013; Sancho *et al.*, 2013) (Figure 4). During the different phases of epiblast development, abundant cell death is observed, showing a peak at 6 days of development, just before the onset of differentiation and gastrulation (Poelmann, 1980; Heyer *et al.*, 2000; Plusa *et al.*, 2008). This maximal incidence of death is partially due to sensitivity to DNA damage and mediated by a p53-dependent quality check (Heyer *et al.*, 2000) and in part regulated by Myc levels (Claveria *et al.*, 2013). Epiblast cells with lower Myc levels preferentially undergo apoptosis due to their confrontation with neighbours with higher Myc levels. This elimination thus results in the recruitment of the pool of cells with higher Myc levels to the developing embryo (Figure 4). When cell death is inhibited by p35 overexpression, rescued cells express relatively lower Myc levels than the rest, indicating that in normal conditions the Myc-low cell population is eliminated (Claveria *et al.*, 2013). Spontaneous cell competition between cells expressing contrasting Myc levels was also detected in mESC cultures (Claveria *et al.*, 2013). However, the biological basis, dynamics and mechanisms of Myc-mediated cell competition remain unknown.

An additional example of endogenous competition has recently been reported for T-cell progenitors in the mouse thymus (Martins *et al.*, 2014). Bone marrow derived colonizing cells outcompeted resident progenitor cells upon competition for the survival factor Interleukin7. Cell competition was needed to continuously renovate the pool of thymic T cell precursors with fresh cells from the bone marrow.
(Martins et al., 2014). Since abolishment of this replacement led to cancerous formation of thymic cells, cell competition would be playing a tumour suppressor role in this phenomenon.

![Diagram of cell competition](image)

**Figure 4. Endogenous cell competition.**

During normal mouse development, Myc levels in the epiblast are intrinsically expressed in a heterogeneous pattern. This produces contacts between cells with high and low-Myc levels, which are fitter and less fit cells, respectively. Myc discrepancies between neighbouring cells leads to outcompetition of the less fit population and a selection of the epiblast cell pool. This process refines the epiblast cell population, giving rise to a final embryo composed by the Myc-high fitter cells from the initial pool. (Based on (Claveria et al., 2013)).

Finally, cell competition has been shown to counter-select naturally viable, but suboptimal cell clones in mammalian fibroblasts (Penzo-Mendez et al., 2015).

**1.6. Mechanisms of cell competition**

The mechanisms by which cells are able to compare their fitness have been intensively investigated. Although several hypotheses have been proposed trying to clarify how competitive interactions occur, complete understanding of the
phenomena observed is far from complete. The different mechanisms and interactions proposed to take place between competing cells are explained here.

1.6.1. Ligand capture model

This was the first model discussed (Morata and Ripoll, 1975), and received support from studies indicating that cells compete for growth/survival factor signals (Moreno et al., 2002). This model was supported by the observation that loser cells presented lower level of Dpp signalling (a secreted protein member of the TGFβ family and the fly orthologue of mammalian BMPs) and an excess of Brinker activated by the JNK pathway and triggering apoptosis (Moreno et al., 2002). However, in several studies the Dpp distribution does not correlate with the intensity of cell competition (Burke and Basler, 1996; de la Cova et al., 2004; Martin et al., 2009). Moreover, in various scenarios where Dpp growth factors are present in excess still competitive behaviours take place (Senoo-Matsuda and Johnston, 2007; Hogan et al., 2009; Tamori et al., 2010; Norman et al., 2012; Claveria et al., 2013; Sancho et al., 2013; Mamada et al., 2015). All these observations suggest that Dpp signalling might result from the execution of competitive program more than being its driving force.

Even though this model gave gratifying explanations, the trophic theory did not address many aspects related to the phenomena, such as the related compensatory proliferation or the mechanism of fitness comparison (Adachi-Yamada and O’Connor, 2002; Gibson and Perrimon, 2005; Shen and Dahmann, 2005).

1.6.2. Models based on fitness reporting

This type of models propose that cells directly report their fitness status to neighbouring cells resulting in the dismissal of cells with comparative lower fitness. The Flower pathway has been the first model proposed for this kind of interaction (Rhiner et al., 2010). Flower is a transmembrane protein ubiquitously expressed in Drosophila. This protein has three isoforms differing in their C-terminal regions that are exposed to the extracellular space. Less fit cells specifically express one of the isoforms (Flower-lose isoforms), which promotes context-dependent apoptosis only.
when surrounding cells express the ubiquitous form (Rhiner et al., 2010). The Flower code has been observed to function in a variety of scenarios, such as cell competition driven by Minute, Myc, Dpp and Scribble, however it is not detected when apoptosis is induced by direct cell death inducers (Rhiner et al., 2010). Since loss of Flower neither affects growth nor homeostasis of tissues, Flower has been proposed to function as a label of loser cells, being considered a dedicated effector of cell competition.

Finally, Crumbs has also been shown to report fitness between prospective winner and loser cells (Hafezi et al., 2012) (Figure 2). As mentioned previously, Crumbs is a transmembrane protein component of the apico-basal polarity system. This protein differentially expressed in neighbouring cells determining the winner and loser fate in a context-dependent manner and mutations that alter fitness also alter Crumbs levels (Hafezi et al., 2012).

However, the molecular mechanisms involved in the ability of the Flower and Crumbs systems to mediate quantitative fitness comparison between cells is still unresolved.

On the other hand, other authors have proposed the release of diffusible signals as a mechanism to report fitness. This concept emerged from experiments where Drosophila S2 WT cells were out-competed when co-cultured with Myc-overexpressing cells separated by a permeable membrane (Senoo-Matsuda and Johnston, 2007). Additional experiments showed that conditioned medium from cultures undergoing Myc-induced competition, but not from cultures expressing Myc in a homogenous manner, was able to induce WT cell death (Senoo-Matsuda and Johnston, 2007). These studies suggested a model in which diffusible factors produced during competition induce loser cell elimination. In this model, prospective loser cells are not specifically targeted, and therefore winner cells should activate a mechanism that protects them or sensitizes loser cells to the killer signal. The combination of the killing signal with winner-specific resistance to it would constitute the fitness comparison mechanism. These experiments however may only apply to certain models of cell competition, since similar experiments in Drosophila (Portela et al., 2010) and mammalian models showed inconsistent results. In mouse ESCs and
mouse fibroblasts, conditioned medium experiments did not induce cell competition and apoptosis was only increased in cells adjacent to Myc-overexpressing cells (Claveria et al., 2013; Penzo-Mendez et al., 2015). In contrast, in BMP-induced mESC cell competition models, diffusible signals were reported to be responsible for competitive interactions (Sancho et al., 2013). Remarkably, in vivo studies show that very short range or direct contact was needed for the elimination of less fit cells (Simpson and Morata, 1981; Moreno et al., 2002; Moreno and Basler, 2004; Li and Baker, 2007; Claveria et al., 2013) and that cell competition does not cross compartment boundaries (Simpson and Morata, 1981), therefore even in the scenarios in which diffusible factors are involved, they would mostly act locally. These results suggest that the molecular mechanism that determine competitive interactions may depend on the model being studied and further analysis needs to be done to elucidate this question and identify the putative signals involved.

Possible examples of competition based on diffusible molecules have been recently reported. An innate immunity-related recognition mechanism has been characterised in cell competition in Drosophila imaginal discs (Meyer et al., 2014). In this study, it was shown that activation mediated by the diffusible signal Späztle of Toll-related receptor is required for low-Myc and elimination of Minute mutant cells (Meyer et al., 2014). Another diffusible killer activity has also been identified during neuronal competition for target innervation in mammals (Deppmann et al., 2008).

1.6.3. Mechanical models

Several studies have proposed mechanical stress as an independent inducer of cell competition in epithelia, where cells compete for limited available space (Shraiman, 2005; Eisenhoffer et al., 2012; Marinari et al., 2012; Paoli et al., 2013). These models may be applied to structural cell competition, where alterations in the apico-basal polarity can alter attachment forces among cells and between cells and the basal membrane. Mechanical stress seems not to be involved in metabolic cell competition (Senoo-Matsuda and Johnston, 2007; Claveria et al., 2013; Sancho et al., 2013; Mamada et al., 2015; Penzo-Mendez et al., 2015). In addition, entosis has been shown to be the predominant mechanism of cell competition between Ras-Rac-transformed cells and WT human cells (Sun et al., 2014b). Entosis is a type of cell
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death initiated when living cells are engulfed by other cells of the same kind (Overholtzer et al., 2007). The ability of a cell to engulf can determine the winner and loser phenotype in some models and is determined by mechanical differences between cells. High deformability controlled by RhoA and actomyosin was reported to be sufficient to explain the winner status, whereby this higher deformability enables cell to engulf their neighbours (Sun et al., 2014b). In addition, E-cadherin overexpression transformed cells induce entosis-mediated cell competition (Sun et al., 2014a). Nevertheless, in some cases the inner cell escapes and does not die, suggesting that cell engulfment events do not always lead to cell death and that the strive for survival of the engulfed cell might represent a fitness status test (Overholtzer et al., 2007).

1.7. Loser cell clearance and replacement

Several mechanisms have been proposed for the clearance of cell corpses after the activation of the apoptotic pathway in those cells. Extrusion from the epithelial layer and phagocytosis are the two main mechanisms for eliminating loser cells. In vertebrate models apical extrusion is predominant (Gu and Rosenblatt, 2012; Lolo et al., 2013) while basal extrusion is predominant in Drosophila epithelia (Grzeschik et al., 2010; Tamori et al., 2010; Lolo et al., 2013), where dead cells are engulfed and remove by haemocytes (Lolo et al., 2012). The sphingosine-1-phosphate and Rho pathways have been shown to be involved in extrusion of dead cells (Gu et al., 2011) and release of tyrosyl-tRNA synthetase by loser cells stimulates hemocyte recruitment and phagocytic activity (Casas-Tinto et al., 2015).

In the Minute model, some studies report that engulfment seems to be the main system to eliminate Minute mutant cells (Li and Baker, 2007). In this context, mutations in essential components for the phagocytic process prevent competition suggesting that engulfment may be upstream of the initiation of the death machinery (Li and Baker, 2007), however these findings have not been confirmed in further experiments (Lolo et al., 2012). Studies have also reported the elimination of polarity mutant cells by engulfment (Ohsawa et al., 2011), contrasting with others in which this process seems not to be required for elimination of the Minute mutant cells, polarity defective cells or WT loser cells in the dMyc supercompetition model (Lolo et
Engulfing by neighbouring cells is also observed in mouse ES and epiblast cells (Claveria \textit{et al.}, 2013). Notably, in these scenarios complete engulfment of intact cells is observed, suggesting that entosis may be involved in the competition process, although further studies would be needed to confirm this (Claveria and Torres, 2016).

After dead cell elimination, expansion of the winner cells allows the replacement of the loser population. Proliferation signals produced by loser cells stimulate this compensatory growth, and these signals may be stress-induced or apoptosis-induced (Claveria and Torres, 2016). Interestingly, a proliferative state is not required for cell competition. In a model of postmitotic competition in the \textit{Drosophila} ovarian epithelium, winner cells undergo hypertrophy instead of overproliferation to fill the space left by the loser population (Tamori and Deng, 2013).

\textbf{1.8. Physiological roles of cell competition}

Cell competition is a mechanism senses and optimizes cell fitness in a wide variety of contexts. An important remark is that while cellular fitness is an intrinsic property cells, the outcome of cell competition is determined by fitness comparison with neighbours. The role of cell competition as a fitness-quality control suggests that this mechanism is important in maintaining healthy tissues and its failure might be involved in a wide range of diseases. However, most cell competition models reported are experimentally induced, which limits the identification of its physiological roles. Investigations of natural cell competition need to be explored further, however this is hampered by the pleiotropy of the majority of the pathways involved in this phenomenon (Claveria and Torres, 2016).

\textbf{1.8.1. Tissue quality control}

The first study describing cell competition, already suggested that it could function as a potential quality control mechanism through eliminating less fit cells (Morata and Ripoll, 1975). Later experiments also suggested a role in regulating organ size during development (de la Cova \textit{et al.}, 2004; Moreno and Basler, 2004).
Myc-driven endogenous cell competition in the pregastrulation embryo (Claveria et al., 2013) suggests the implication of competitive behaviours in a natural quality control checking in the epiblast. Interestingly, due to the pluripotency of epiblast cells, these can be eliminated and replaced without compromising embryo development (Claveria and Torres, 2016).

Recently, Azot has been characterized as a mediator of apoptosis induction specifically in several types of cell competition, but not in other stress-response pathways, in Drosophila (Merino et al., 2015). The function of Azot in cell death induction is therefore strongly linked to cell competition-induced cell death. Loss of Azot leads to accumulation of defective cells and premature aging (Merino et al., 2015), indicating an important role of tissue quality control during Drosophila development and adult tissue homeostasis.

An interesting example of natural cell competition controlling tissue quality is the analysis of lineage displacement in the Drosophila female germline, in which cell competition has been proposed to maintain the boundaries between the GSC niche and the differentiating compartment (Jin et al., 2008; Rhiner et al., 2009). Therefore, the proposed function of cell competition is GSC stemness maintenance, although its relevance in maintaining GSC quality during normal oogenesis has not been explored (Jin et al., 2008; Rhiner et al., 2009).

1.8.2. Regeneration

Several studies suggest the implication of cell competition in regeneration. When fetal hepatocytes are transplanted into a damaged adult liver, transplanted cells replace resident cells by progressively eliminating them (Oertel et al., 2006). This colonization of the tissue is not only due to their increased proliferative activity, but also to the induction of cell death of the adult hepatocytes localized close to transplanted cells (Oertel et al., 2006). In addition, experiments in mouse adult heart show that Myc-overexpressing cardiomyocytes can replace WT cells without modifying the function and anatomy of the heart, suggesting a potential role of cell competition also in the maintenance of the myocardium (Villa del Campo et al.,
More recently, in a model of *Drosophila* neural regeneration, it has been shown that the comparison of neuronal fitness leads to the elimination of non-scarred injured tissue, helping the damage resolution (Moreno et al., 2015).

### 1.8.3. Cancer

Finally, the relevance of cell competition for cancer has been extensively hypothesized and explored. Interestingly, cell competition has been characterized both as a tumour-suppressing and as tumour-promoting mechanism.

On one hand, although the expression levels that drive metabolic cell competition do not disrupt normal development or tissue homeostasis, the pathways involved in promoting the winner phenotype promote oncogenesis in humans and tumour formation in experimental models (Claveria and Torres, 2016). Supercompetition has been linked to field cancerization models in which cells carrying pre-oncogenic lesions silently expand in tissues at the expense of surrounding normal cells (Rhiner and Moreno, 2009). Examples of this are the expansion of *Kras* mutation in the intestine (Snippert et al., 2014) and *Notch* mutations in the oesophagus (Alcolea et al., 2014), both driving experimental field cancerization. Other studies connect flower deficiency (Petrova et al., 2012) and SPARC overexpression (Petrova et al., 2011; Yamada et al., 2015) with tumour formation. More recently, cell competition has also been implicated in tumour expansion and invasiveness once the tumour is established (Eichenlaub et al., 2016; Suijkerbuijk et al., 2016).

On the other hand, the pathways involved in structural cell competition are also linked with oncogenesis and carcinoma formation in a variety of tissues. In this case, the tumour-promoting phenotype is mostly associated with the loser fate, indicating a tumour suppressing role for cell competition (Kajita and Fujita, 2015; Claveria and Torres, 2016). Other experiments show a tumour-suppressing role when mutations affect individual cells or small number of cells but a tumour-promoting role of cell competition in larger growing tumours (Ballesteros-Arias et al.).
2. Early development of the mouse embryo, ESCs and pluripotency

2.1. From fertilization to gastrulation

Development of the zygote or fertilized egg proceeds through symmetric cleavage divisions until the morula is formed. Cells at the early morula stage are totipotent blastomeres, which are able to give rise to all embryonic and extra-embryonic lineages (Gardner, 1998). Removal of one or more cells at this stage (6 to 8-cell embryo) does not affect normal development (Nichols and Smith, 2009). Moreover, the fate of the individual cells can be manipulated by changing their position within the embryo (Hillman et al., 1972) (Figure 5).

After a fixed number of cell cycles, the individual blastomeres compact and at 32-cell stage those blastomeres on the periphery differentiate and form an epithelium called trophectoderm (TE) (Selwood and Johnson, 2006) while a group of cells known as the inner cell mass (ICM) remain internal and pluripotent. Pluripotency is defined as the ability to generate all embryonic cell lineages but not the complete set of extraembryonic tissues. The appearance of the blastocoel, a cavity below the TE produced by fluid pumping, generates the blastocyst (Nichols and Smith, 2009). The ICM then segregates in epiblast, constituted by pluripotent cells precursors of the embryo proper, and the hypoblast or primitive endoderm (PE), which mostly contributes to extraembryonic or non-definitive endoderm. At E3.5, before implantation in the uterus, the three lineages of the blastocyst are well established and this process is autonomous with respect to external cues from fertilization to implantation (Figure 5).

Initial models of epiblast-primitive endoderm segregation considered the early ICM as a homogeneous population in which cells were allocated to the two lineages according to their position with respect to the blastocoel (Chazaud et al., 2006). However, later studies showed that the ICM is already heterogeneous at E3.5. The random “salt and pepper” expression of two transcription factors in a mutually exclusive manner, Nanog and Gata6, determines the two lineages within a single cell population and then these lineages segregate by sorting out into the primitive
endoderm, facing the blastocoel, and the epiblast, positioned between the trophectoderm and the primitive endoderm (Chazaud et al., 2006) (Figure 5).

**Figure 5. Overview of some stages of early mouse embryogenesis.**

Frontal view of the pre- and post-implantation embryo at different stages. The zygote undergoes symmetric cleavages giving rise to the morula, which is then compacted. At E3 blastomeres start to segregate and form the trophectoderm, which is the first event of differentiation. At E3.5, the second event of differentiation takes places and some cells from the inner cell mass form the primitive endoderm. Later, the blastocyst hatches from the zona pellucida and implants in the uterine wall. The primitive endoderm migrates to become the parietal and visceral endoderm, which covers the epiblast. At E6, the epiblast is a monostratified epithelium. Gastrulation begins at E6.5 when some cells of the posterior pole undergo an epithelial-mesenchymal transformation to generate mesoderm and form the primitive streak.
Later, the blastocyst hatches from the zona pellucida and implants in the uterine wall. As a natural selection checkpoint, only embryos able to hatch continue developing, invading the maternal tissue and implanting (Bedzhov et al., 2014). After implantation, the trophectoderm becomes the trophoblast and contributes to extraembryonic membranes, while the epiblast generates the embryo plus some extra-embryonic tissues. In the first two days post-implantation, the mural trophectoderm (cells that surrounds the blastocyst cavity that are not in contact with the ICM) gives rise to polyploid Trophoblast Giant Cells (TGCs), which represent the first terminally differentiated cell type (Bedzhov et al., 2014). Contrary to this, the polar TE, which surrounds the outer surface of the ICM, is the source of multipotent progenitors, the TE stem (TS) cells. These cells proliferate and become the ectoplacental cone (EPC) that builds the proximal half of the egg cylinder and later contributes to the placenta, and the extraembryonic ectoderm (ExE). The primitive endoderm migrates to become the parietal endoderm, which covers the inner surface of mural TE, and the visceral endoderm, which covers the egg cylinder and epiblast. By six days after fertilization, the epiblast is a cup-shaped monostratified epithelium of ~1000 cells (Figure 5).

At 6.5 days after fertilization the first morphological sign of embryonic patterning is evident when gastrulation begins (Beddington and Robertson, 1999). Some cells of the future posterior pole of the epiblast, which is apparently a homogeneous epithelium, undergo epithelial-mesenchymal transformation (EMT) to generate mesoderm and form the primitive streak (Beddington and Robertson, 1999). Cells move through the streak and spread forward to form mesoderm. The visceral endoderm is replaced by the definitive endoderm that derives from the epiblast (Figure 5). During the next hours, the streak elongates from the rim to the distal tip of the cup and the node forms. This specialized structure is considered homologous to the organizer of other vertebrate embryos and gives rise to the axial gastrulating cells that comprise the axial mesoderm (future prechordal plate and notochord) and the definitive gut endoderm (Beddington and Robertson, 1999).
2.2. Embryonic Pluripotency: defining the Naive and Primed pluripotent states in mammalian development

The mammalian egg is totipotent as it can give rise to the entire embryo. However, this does not mean that the zygote can directly differentiate into the all cell types. In fact, the egg and the blastomeres derived from it produce directly only two types of cells: trophectoderm cells and cells from the inner cell mass (Nichols and Smith, 2009).

As mentioned before, pluripotency is the capacity of cells to produce all three embryonic germ layers, but not trophectoderm-derived extra-embryonic tissues (Hanna et al., 2010). Pluripotency is not a single status, but evolves in a highly dynamic manner through different stages of pre- and post-implantation development. To develop, the ICM must be able to generate other cell types in a flexible manner (Gardner and Beddington, 1988; Nichols and Smith, 2009). The early ICM is the founder pluripotent population. Once it segregates into PE and epiblast at the late blastocyst stage, the epiblast retains the pluripotency, while the PE loses it and commits with the endodermal fate. Following segregation of PE, the epiblast of the late blastocyst (embryonic day 4.0) acquires a pluripotency state known as “naive” or “ground state” (Hackett and Surani, 2014). The naive state pluripotent epiblast is functionally and molecularly different from blastomeres and early inner cell mass (Gardner, 1998; Kurimoto et al., 2006; Kaji et al., 2007; Nichols and Smith, 2009). The naive state is characterized by homogenous expression of key pluripotency transcription factors and by the elimination of gametic epigenetic silencing, which results in a generalized hypomethylated “open” chromatin state, including both X-chromosomes in female embryos. The “naive state” is therefore considered as the unrestricted pluripotent state in which cells show an unbiased developmental potential.

Departure from the naive status starts in the epiblast shortly after implantation; however, pluripotency is maintained for several days before differentiation starts during gastrulation (E6.5). The transit between naive-state epiblast cells takes place in a continuum manner in which cells transit through a “formative” phase into an “primed” pluripotency state before they actually commit.
with a particular fate and differentiate (Nichols and Smith, 2009; Hackett and Surani, 2014; Kalkan et al., 2017).

In addition, both states are characterized by expressing the core pluripotency network. Initially this regulatory network was almost exclusively referred to a small set of transcription factors, Oct4, Sox2 and Nanog (Orkin, 2005). These three factors converge and form the primary, yet central, network that governs the robust pluripotent state. On one hand, Oct4 is essential for early mouse development and maintenance of the ES cell pluripotency (Nichols et al., 1998). On the other hand, Sox2 is generally considered a transcriptional partner of Oct4 (Avilion et al., 2003). And finally, the presence of Nanog in tissue culture relieves the requirement for LIF by promoting ES cell self-renewal (Chambers et al., 2003; Mitsui et al., 2003). In spite of the relevance of these key transcription factors and their cooperative autoregulation to maintain ESCs properties, some studies also point to the involvement of additional factors that either provide feed-forward regulations to stabilize the pluripotency network, or exert other cellular functions required for ESC identity maintenance (Ivanova et al., 2006; Wang et al., 2006; Zhou et al., 2007).

### 2.3. Naive and Primed stem cell pluripotent states in culture

The different types of pluripotency observed in vivo can be captured in apparently equivalent states in cultured cells. The transient nature of naive pluripotency is captured in vitro in ESCs derived from the ICM or through experimental reprogramming (Evans and Kaufman, 1981; Martin, 1981; Yamanaka and Blau, 2010; Hackett and Surani, 2014) (Figure 6). ESCs represent a surrogate in vitro model of the naive epiblast status that, under permissive culture conditions, keeps naive pluripotent capacity while undergoing self-renewal. On the other hand, both in vitro models of epiblast stem cells (EpiSCs) derived from late epiblast (Brons et al., 2007; Tesar et al., 2007) and epiblast-like cells (EpiLCs), which are derived from ESCs showing a cellular state highly similar to pregastrulating epiblasts but distinct from EpiSCs (Hayashi et al., 2011), capture the primed pluripotent state (Figure 6), which provides another system to study pluripotency and cell fate decisions.
The definition of both states implies that the naive ESCs correspond to a more immature state of pluripotency whereas the primed state is prone to differentiate, which is reflected, for example, in the state of X-chromosome inactivation. In naive female cells, both chromosomes are active; on the contrary, primed EpiSCs have already undergone X-chromosome inactivation (Hanna et al., 2010).

Furthermore, the two pluripotent states exhibit distinct molecular signatures and in vitro growth properties (Figure 7). In addition, the two states depend on signalling pathways that often antagonize each other (Hanna et al., 2010). Although the core pluripotency network is expressed in both naive and differentiating-primed states (Wang et al., 2006), they differ in the signalling pathway they rely on (Figure 7). The BMP/SMAD signalling pathway (Ying et al., 2003) and LIF/STAT3 (Smith et al., 1988) maintain naive pluripotency of ESCs, whereas FGF/MAPK promotes

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**Figure 6. Scheme of different types of mouse pluripotent stem cells and their sources.**

Several pluripotent cell types can be derived by explanting cells at various stages of early mouse embryonic development. ESCs are derived from cells harvested at the blastocyst stage. Pluripotent cells derived from post-implantation embryo give rise to epiblast stem cells (EpiSCs). Embryonic germ cells and spermatogonial germ cells (GSC) are derived from primordial germ cells (PGCs) from embryos and spermatogonial progenitors from adults respectively. Alternatively, somatic cells can be directly reprogrammed in vitro by exogenous expression of transcription factors generating induced pluripotent stem cells (iPSCs). In addition, epiblast-like cells (EpiLCs) can be generated from either ESCs or iPSCs in a progressive process.
transition to the primed state and, together with Activin/Nodal promote definitive differentiation (Brons et al., 2007; Tesar et al., 2007) (Figure 7).

Based on this knowledge, different culture conditions can promote the different states of pluripotency found in the embryo. Fetal bovine serum, together with mouse embryonic fibroblasts (MEFs), was originally used to derive ESCs (Martin and Evans, 1975). Serum contains BMP4, which is crucial for activating Inhibitor of Differentiation (Id) genes via SMAD pathway (Figure 7). Serum can be thus substituted by overexpression of Id genes or exogenous addition of BMP4 (Hackett and Surani, 2014). However, serum contains as well FGF and activin-like activities that promote exit from the ground state and differentiation and MEFs were important in limiting this differentiation promotion. LIF, which activates the JAK-STAT3 signalling pathway (Figure 7), was later reported to be essential for enabling the proliferation of ESCs without MEFs (Smith et al., 1988; Williams et al., 1988).

Maintenance of naive pluripotency can be achieved by serum/LIF culture conditions. However, in these conventional conditions self-renewal is maintained by abrogation or counterbalance of differentiation stimuli downstream of their effects. Therefore, in serum/LIF ESCs exist in a battleground of competing signals and exhibit heterogeneous expression of pluripotency markers that gives rise to metastability. In fact, only a fraction of these cells correlate with the pre-implantation epiblast, as shown by transcriptional and epigenetic profiling (Ying et al., 2008; Marks et al., 2012; Boroviak et al., 2014).

It is thus preferable to insulate, rather than counteract, differentiation stimuli in order to stabilize upstream naive state. Two inhibitor (2i) molecules have been found sufficient to protect ESCs from differentiation stimuli and select against differentiating cells. PD03 (PD0325901) inhibits FGF/ERK signalling pathway by blocking ERK1/2 phosphorylation (Ying et al., 2008), which appears to drive transition from naive to primed state (Kunath et al., 2007) (Figure 7). In the presence of PD03 LIF is still required to support ESC naive and self-renewing state. GSK3 inhibition enhances ESC self-renewal (Sato et al., 2004; Ogawa et al., 2006), and the inclusion of the GSK3 inhibitor, CHIRON (CHIR99021) (Figure 7) in addition to PD03 sustains ESC naive state propagation, even in the absence of LIF or
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serum/BMP4 (Ying et al., 2008). Collectively PD03 and CHIRON inhibitors are known as the “2i” culture system. In 2i/LIF cultures most of ESCs exhibit molecular and functional properties corresponding to an optimized state of naive pluripotency that is closely comparable to the developmental ground state in vivo. Therefore, ESCs in 2i cultures retain the essential features of early epiblast pluripotent cells (Boroviak et al., 2014), being considered the best current approach as a model for naive epiblast cells.
Additionally, investigation about pluripotency has been enhanced after the discovery of cell reprogramming by forced expression of a set of transcription factors, which also provided an enormous potential for regenerative medicine. Previous studies show that somatic cells can be reprogrammed by transferring their nuclear contents into oocytes (Wilmut et al., 1997) or by fusion with ESCs (Tada et al., 2001; Cowan et al., 2005) indicating that unfertilized eggs and ES cells contain factors that can confer totipotency or pluripotency to somatic cells. However, a later works demonstrated the flexibility of cellular identity by the full reprogramming of differentiated somatic cells to pluripotent stem cells through forced expression of a set of transcription factors. These new generated pluripotent cells are called induced pluripotent stem cells (iPSCs) (Takahashi and Yamanaka, 2006). The initial genes identified as responsible for reprograming adult cells areOct3/4, Sox2, Klf4 and Myc(Takahashi and Yamanaka, 2006).
3. The proto-oncogene Myc family in development

3.1. Functions of the transcription factors of the Myc family in mammalian development

During the last decades, the role of the proto-oncogene Myc gene family (comprising 3 transcription factors: Myc, Mycn and Mycl) in normal and cancer cells has been extensively studied. The fact that Myc genes are deregulated in most tumours has attracted attention across the cancer research field. Nevertheless, in this thesis we are interested in non-oncogenic roles of this family in normal development, pluripotency and cell competition. The three main members of the proto-oncogenic Myc family are involved in several cellular processes, such as the regulation of cellular proliferation, differentiation and apoptosis (Meyer and Penn, 2008). The role of Myc proteins in development has been intensively investigated. Even though Mycl is widely expressed in the mouse embryo, it appears not to be necessary for embryonic development, as Mycl knockout mice develop normally (Hatton et al., 1996). Contrary to this, Myc deficient embryos die before E10.5 due to hematopoietic and placental defects (Trumpp et al., 2001; Dubois et al., 2008) and embryos lacking Mycn are die at E11.5 exhibiting neuroectodermal and heart defects (Charron et al., 1992), indicating that endogenous Myc and Mycn are critical for completing embryogenesis. However, they have also been shown to be separately dispensable for murine embryonic stem cell function. The lack of phenotype of the single KOs suggests highly redundant functions of Myc and Mycn (Malynn et al., 2000). Moreover, recently it has been shown that Myc and Mycn cooperate in maintaining the ESC proliferative program (Scognamiglio et al., 2016).

3.2. Myc factors and pluripotency

Myc has been reported to regulate cell cycle regulation, cell growth and metabolism in ESC, by regulating a different subset of genes than those targeted by the core pluripotency network transcription factors (Chen et al., 2008; Hu et al., 2009; Kim et al., 2010; Claveria et al., 2013; Gu et al., 2016; Scognamiglio et al., 2016) and has multiple roles in several stem and progenitor cell types (Laurenti et al.,
2009). Myc also controls the balance between self-renewal and differentiation in adult hematopoietic stem cells (Wilson et al., 2004).

Additionally, in ESCs cultured in serum plus LIF, Myc proteins have been suggested to sustain pluripotency through repression of the primitive endoderm master regulator GATA6 and contribute to the cell cycle control by regulating the mir-17-92 miRNA cluster (Smith et al., 2010).

Furthermore, Myc has been described to promote cell reprogramming of somatic cells into a naive pluripotent status in cooperation with transcription factors of the core pluripotency network, such as Oct4, Sox2 and Klf4 (Cartwright et al., 2005; Takahashi and Yamanaka, 2006). In addition, primed EpiSCs can also be reverted to the naive state by overexpressing Klf4 or Myc in EpiSCs under LIF-exposure conditions (Guo et al., 2009; Hanna et al., 2009). More recently, ESC self-renewal capacity has been reported to be maintained thanks to a Myc-dependent self-propagating epigenetic memory, which is established by a Myc-driven positive feedback loop (Fagnocchi et al., 2016).

Even though several studies have tried to shed light on the relationship between Myc and pluripotency and some roles have been reported both in vivo and in vitro, the precise role of Myc with respect to pluripotency states in embryos and cultured cells remain elusive.
In the field of Cell Competition, many studies have tried to shed light on how fitness comparison is produced between neighbouring cells and results in the elimination of the less fit cells. Natural cell-to-cell heterogeneity in Myc levels has been reported to trigger cell competition both in early mouse development and ESCs. In this thesis, we aimed to perform further studies to understand the dynamics of Myc-related endogenous cell competition and explore the biological relevance of this phenomenon during normal mouse development and ESC maintenance.

The specific aims of this thesis were:

Generate a new time-lapse microscopy and image analysis tool to track Myc levels during cell competition in ESCs.

Determine the dynamics of Myc heterogeneity and how it is regulated in pluripotent cells.

Determine how the cell-to-cell Myc heterogeneous levels lead to cell competition and the dynamics of Myc expression during the process.

Determine the biological basis of Myc-driven endogenous cell competition in vitro and in vivo.
1. **Mouse lines and mESC derived lines**

The \textit{iMOS}^{WT}, \textit{iMOS}^{T1-Myc} and \textit{iMOS}^{T2-p35} mouse lines have been previously described (Claveria \textit{et al.}, 2013). After recombination both labelled populations are WT in the \textit{iMOS}^{WT} cell line. In the \textit{iMOS}^{T1-Myc} the EYFP population overexpresses Myc and ECFP cells are WT. In the \textit{iMOS}^{T2-p35} the EYFP population is WT and the ECFP population expresses the baculoviral apoptosis inhibitor p35.

\textit{Sox2Cre} mice (Hayashi \textit{et al.}, 2002) were used to activate the \textit{iMOS}^{T2-p35} line when the experiment was performed in embryos and to generate the Myc KO from the Myc^{flx/flx} line.

The GFP-Myc reporter line has been previously described (Huang \textit{et al.}, 2008). Homozygous mice are viable and show no apparent phenotypic alterations.

WT mice used are of the CD1 strain.

Mouse embryonic stem cells were derived from E3.5 embryos from the following mouse lines: \textit{iMOS}^{WT}, \textit{iMOS}^{T1-Myc}, \textit{iMOS}^{T2-p35} and GFP-Myc. The protocol for establishing ESC lines is explained below.

2. **E5-6 embryo harvest**

Mice embryos were extracted at different developmental stages. Vaginal plugs were checked every morning. Midday of the day when vaginal plug is detected is considered gestational day 0.5 (E0.5). To obtain the embryos, females were sacrificed by CO$_2$ inhalation and abdominal cavity was opened to expose the uterus, which was transferred to sterile PBS. Then working in cold PBS under the scope, muscular uterine wall was carefully ripped and then the decidual layer was removed. Exposed embryos were fixed in paraformaldehyde (PFA) (Merck) 2% in PBS overnight at 4\textdegree{}C. After fixation embryos were washed in PBS several times.
3. Mouse embryonic fibroblasts culture

Mouse embryonic fibroblasts (MEFs) were only used as a feeder layer for the stem cells. Fibroblasts have been previously extracted from E10.5 embryos by the Pluripotent Cell Technology Unit staff. Approximately 5 million of cells from one vial were plated on 100-mm 0.1% gelatine-coated plastic plates (Falcon). After 2 days cells were passed to 2 150-mm 0.1% gelatine-coated plastic plates (Falcon) and cultured again for another 2 days. Fibroblasts were then inactivated with mitoMycin-C (Sigma), trypsinized (Trypsin-EDTA 10x, Gibco) and frozen. MEF culture medium contained the following reagents:

- High glucose DMEM, GlutaMAX™ Pyruvate (LifeTech)
- 15% Fetal Bovine Serum (FBS)
- 1% Penicillin/Streptomycin (10,000U/ml; 100x)
- 0.2% 2-beta-mercaptoethanol (50mM)

MEFs freezing medium contained:

- 60% MEF culture medium
- 20% FBS
- 20% DMSO (Sigma)

4. Mouse ESC derivation and establishment

Mouse ESCs were derived from E3.5 embryos. Female sacrifice and uterus extraction was done as described. Embryos extraction was performed by flushing the blastocysts out of the uterus under a dissection scope using a 1-ml syringe with 23-G needle. Blastocysts were collected and transferred individually to wells of a 24-well plate (Falcon) containing fresh mitoMycin-C-inactivated MEF feeder layer. Culture was maintained for 5-8 days in 2i (GSK3β and Mek 1/2 inhibitors) medium. Medium was not changed until the blastocysts attached, then it was changed every day. Those hatching blastocysts whose inner cell mass expanded were properly trypsinized and seeded on a well of 12-well plate (Falcon) and then went through several passages in order to expand the clone.
2i derivation medium contained:

- High glucose DMEM, GlutaMAX™ Pyruvate (LifeTech)
- 15% KO-Serum Replacement (Invitrogen)
- Leukemia Inhibitor Factor (LIF)
- 1% nonessential amino acids (100x)
- 1% Penicillin/Streptomycin (10,000U/ml; 100x)
- 0.3 μM GSK3β inhibitor CHIR99021 (Stemolecule™)
- 0.1 μM Mek 1/2 inhibitor PD0325901 (Stemolecule™)

Cultures were checked for Mycoplasma by the Cell Culture Unit staff and for normal karyotype (protocol explained below) at their establishment.

For time-lapse studies, the ESC lines were stably transfected with a PGK-ttdTomato construct that drives expression of a membrane-attached Tomato fluorescent protein.

### 4.1. Karyotyping

Approximately 1.5x10⁶ cells in a p-60mm plate (Falcon) were treated with 0.5μg/ml colcemid (KARYO MAX COLCEMID, Gibco) for 1 hour at 37°C. Cells were then trypsinized and resuspended in 6 ml of a hypotonic solution (75mM KCl) for 7 minutes at room temperature. 8 drops of a fixative solution (3:1 methanol/glacial acetic acid) were added and then centrifuged for 5 minutes at 900rpm at room temperature. Supernatant was removed and 6ml of the fixative solution were added again, letting 20 minutes in ice. This last step was repeated twice. Finally, a small volume of the solution was dropped on a slice, dried and mounted with DAPI. Karyotype of a cell line was considered suitable when more than 80% of the cells exhibited normal karyotype.
5. Mouse ESC routine culture

ESCs were routinely maintained on mitoMycin-C-inactivated MEF feeder layers. Cells were passed every 2 days and frozen 800,000-900,000 cells per vial. ESC routine culture medium (SR-ES medium) contained the following reagents:

- High glucose DMEM, GlutaMAX™ Pyruvate (LifeTech)
- 15% KO-Serum Replacement (Invitrogen)
- Leukemia Inhibitor Factor (LIF)
- 1% nonessential amino acids (100x)
- 0.1% 2-beta-mercaptoethanol (50mM)

mESCs freezing medium contained:

- 60% ESC culture medium
- 20% FBS
- 20% DMSO (Sigma)

6. iMOS induction for functional assays in ESCs

For the iMOS$^{T1-Myc}$ and iMOS$^{T2-p35}$ induction, cells were plated without feeder on fibronectin-coated 3.5-mm glass bottom dishes (MatTek) and cultured in 2xLIF SR-ES medium. After 24h iMOS$^{T1-Myc}$ cells were treated with 10 µM 4-hydroxytamoxifen (4-OHTAM) (Sigma-Aldrich) and iMOS$^{T2-p35}$ cells with 20 µM for 24 h and cultured for additional 24 h.

7. Immunofluorescence

7.1 Whole mount embryo immunofluorescence

For E5-E6.5 embryos, same procedure was followed for all the stainings. Embryos were permeabilized in 0.5% Triton X-100 (Calbiochem) for 30 minutes. Excess of triton was washed in PBS during 15 minutes. Blocking was performed in
10% goat serum (Gibco-BRL Life-Technologies) 0.1% Triton X-100 for one hour and then incubation with the primary antibodies was performed overnight at 4°C. Embryos were washed several times and incubated with secondary antibodies and DAPI (1:2000) overnight at 4°C. Following day and after several washes, embryos were ready to be imaged and store in PBS or Vectashiel mounting medium (Vector Laboratories, USA). Primary and the secondary antibodies were diluted in the blocking solution.

7.2. Mouse ESC immunofluorescence

For mouse ESCs the same protocol was followed for all the stainings, excepting for pSTAT3 immunofluorescence. Fixed samples were permeabilized in 0.25% Triton X-100 (Calbiochem) from 10 to 30 minutes depending on the antibody, washed in PBS and blocked with 10% goat serum 0.1% Triton X-100 for one hour. Incubation with the primary antibodies was performed at room temperature for one hour. After several washes, secondary antibodies and DAPI (1:2000) were incubated during 1 hour at room temperature. Cells were then washed and mounted in Vectashiel mounting medium.

For pSTAT3 staining, cells were permeabilized in cold 100% methanol for 10 minutes at -20°C and the blocking solution contains 5% goat serum in 0.25% Triton X-100. Primary and the secondary antibodies were diluted in the blocking buffer.

7.3. Primary antibodies

- anti-Myc polyclonal antibody (06-340, Millipore) 1:300
- anti-Myc T58 (ab85380, abcam)
- phospho-p44/42 MAPK (Erk1/2) (Thr202/Tyr204) (#4370, Cell Signaling) 1:200
- anti-phospho-histone H3 (Ser 10) ‘mitosis marker’ (06-570, Millipore) 1:300
- Phospho-Histone H3 (Ser10) (6G3) (#9706, Cell Signaling) 1:300
- cleaved-caspase3 (Asp175) Antibody (#9661 Cell Signaling) 1:100
7.4. Secondary antibodies

All of them were used 1:500.

- 488 goat anti-rabbit (Life Tech)
- 488 goat anti-mouse (Life Tech)
- Cy3 goat anti-rabbit (Jackson)
- 568 goat anti-mouse (Life Tech)
- 568 goat anti-chicken (Life Tech)
- 633 goat anti-rabbit (Life Tech)
- 633 goat anti-mouse (Life Tech)
- 647 donkey anti-chicken (Jackson)

8. Confocal microscopy

Immunofluorescence of fixed cells was imaged with either a Leica TCS SP5 inverted conventional confocal microscope fitted with a HCX PL APO CS 40x 1.25 oil objective, an SP8 inverted conventional confocal microscope fitted with HC PL Apo CS2 40x/1.3 oil objective, or a Zeiss LSM 780 upright microscope fitted with W Plan-Apochromat 40x/1,0 DIC M27 objective. Whole mouse embryos were imaged with a Zeiss LSM 780 upright microscope fitted with a W Plan-Apochromat 20x/1,0 DIC M27 75mm WD 1.8mm objective in confocal mode at 405, 488, 568 or 663 nm. Z-stacks were acquired in order to image the whole embryo. Acquisitions were commonly 1024x1024 pixels, A.U. set to 1.5.
9. Time-lapse video microscopy of ESC cultures

200,000 GFP-Myc ESCs were seeded on 3.5-mm 0.1% gelatine-coated plastic plates (Falcon) without feeder and cultured in 2xLIF SR-ES medium. 24 h later cells were transferred to the environmental chamber of a Zeiss LSM 780 upright microscope. In every imaging experiment, the incubation chamber was previously stabilized to 37°C and 5% CO2 by mixing and humidification. To avoid liquid evaporation, embryo-grade mineral oil was added on top of the medium after objective immersion. Samples were imaged with a W Plan-Apochromat 40x/1.0 DIC M27 objective every 7 minutes for 24 h. Z-stacks with sections spaced at 2 µm are acquired at each timepoint. In addition, a multi-position map was generated, and every ESC colony was manually assigned an x-y-z location. We used confocal lasers with two excitation channels (488 nm excitation for GFP-Myc visualization and 561 nm excitation for the tdTomato membrane marker), and all images were 512x512 pixels (0.415x0.415 µm resolution), taken at speed 7 and with 2 scans/Z-slice. A trade-off between spatial resolution, temporal resolution and total acquisition time was carefully selected to minimize photobleaching while enabling tracking of cell division, apoptosis and cell-cell interactions. For apoptosis detection, we added either Hoechst33258 (B-2883, Sigma) (1:2000) or caspase3-reporter (CaspGLOW™ Red Caspase Staining kit, MBL) (1:1000) to the medium immediately before imaging.

For iMOS<sup>T1-Myc</sup> induction analysis, we started time-lapse confocal analysis 10 h after the induction. The timings of new EYFP<sup>+</sup> cells and apoptotic events were scored manually.

In the GFP-Myc and iMOS<sup>T1-Myc</sup> coculture, both populations were seeded in a proportion 1:1. The timings of new EYFP<sup>+</sup> cells and apoptotic events were scored manually.

10. Clonal analysis assay

150,000 ESCs were plated without feeder on fibronectin-coated 3.5-mm glass bottom dishes (MatTek) and cultured in 2xLIF SR-ES medium. To generate clones we used the iMOS<sup>WT</sup> cell line (Claveria et al., 2013) and induced EYFP clones at a very
low recombination rate. For this, cells were treated 48h after plating for 1 h with 5 μM 4-hydroxytamoxifen (4-OHTAM) (Sigma-Aldrich) and cultured for an additional 60h. Only 15% of colonies were recombined, so the expected frequency of polyclonal groups was around 2% (0.15±X100). Cells were fixed and analyzed by confocal immunofluorescence for Myc, and subject to confocal analysis. We then calculated the variation of Myc levels within clonal groups identified by EYFP signal and for a set of equivalent randomly chosen cell groups. We repeated the process at least 100 times per colony to have a sample representative of the whole population and considered the median standard deviation of the 100 subsamples.

11. Flow Citometry

For cell cycle analysis, cells were trypsinized, incubated in culture medium with 5 μg/ml Hoechst33342 (B2261-Sigma) for 45 minutes at 37°C, washed and resuspended in 2% FBS PBS and analysed by flow cytometry in a BD FACSCanto™ II (laser wavelengths 405, 488, 561, 640). Propidium iodide (P4864-Sigma) was used to identify dead cells.

For the DNA demethylation experiments, 5-Azacytidine was previously dissolved at 20.5 mM in water:acid acetic (1:1) and cells were treated with 0.25 μM 5-Azacytidine (A2385, Sigma). After treatment cells were washed, trypsinized, resuspended in 2% FBS PBS and stained with Hoechst33258 (B-2883, Sigma) for dead cell exclusion. Cell suspensions were analysed in a BD LSRFortessa™ Special Order Research Product (laser wavelengths 405, 488).

For the LIF withdrawal assay, cells were trypsinized, resuspended in 2% FBS PBS and stained with Hoechst33258 (B-2883, Sigma) for dead cell exclusion. Cell suspensions were analysed in a BD LSRFortessa™ Special Order Research Product (laser wavelengths 405, 488, 561).

For the analysis, FACSDiva and FlowJo softwares were used.
12. RNA isolation from cells

In order to perform RNA differential expression analysis, RNA from cells was isolated. For assess the difference in gene expression depending on Myc levels, total RNA was purified from GFP-Myc ES cells sorted in BD FACSARia™ II flow cytometer according to GFP-Myc (three biological replicas). RNA was extracted using RNeasy Mini Kit (Qiagen). Isolated RNA was then handed to Genomic Unit in CNIC to proceed with sequencing.

13. RNASeq. Data analysis

13.1. RNAseq library production

In order to assess the difference in gene expression among the three populations depending on Myc levels, total RNA was purified from ES cells sorted in a cytometer according to GFP-Myc (three biological replicas). A cDNA library was prepared according to Illumina recommendations and sequenced in a GAI1 analyzer using the paired-end 50bp protocol. Sequencing reads were pre-processed by means of a pipeline that used FastQC to assess read quality, and Cutadapt v1.3 (Martin, 2011) to trim sequencing reads, eliminating Illumina adaptor remains and discarding reads shorter than 30 bp. Reads were mapped against the mouse transcriptome (GRCm38 assembly, Ensembl release 76) and quantified using RSEM v1.2.3 (Li and Dewey, 2011).Raw data have been deposited in the GEO database with reference number GSE89196 .Raw expression counts were then processed with an analysis pipeline that used the Bioconductor package EdgeR (Robinson et al., 2010) for normalization and differential expression testing. Only genes expressed at a minimal level of 1 count per million, in at least 3 samples, were considered for differential expression analysis.

13.2. RNAseq meta-analysis

Gene set enrichment analysis was performed using the online tools in the Broad Institute GSEA web site (http://software.broadinstitute.org/gsea/). Genes with significant variation between Myc-H and Myc-L cells (adj pval <0.001) were
screened against the curated “chemical and genetic perturbations” database. 8 of the top-ten hits corresponding to experiments in embryonic stem cells were gene sets identifying targets of pluripotency factors (p-values between $<3.78^{-25}$ and $<1.73^{-62}$).

For Principal Components Analysis, we compared our RNA-seq dataset to Gene Expression Omnibus RNA-Seq datasets GSE67259 and GSE56138. For GSE67259 dataset, expression levels in RPMs were compared with the TPMs calculated from FPKMs of dataset GSE56138 and to the TPMs of our dataset. For each gene symbol, we considered the TPMs of all samples from all datasets. 16,998 genes were expressed in all experiments. Combat (Robinson et al., 2010) without covariate was then used on the log2-TPMs matrix to remove batch differences among the datasets. ggfortify and gplots packages were used for visualization of the Principal Components Analysis and main gene set contributing to the Principal Component 1.

14. Image analysis

14.1. Nuclear and cytoplasmic signal detection

Confocal images were analyzed with ImageJ (http://imagej.nih.gov/ij). To quantify nuclear signal, nuclei were detected by DAPI staining and segmented by using threshold tool to create a mask. Manual correction was applied to ensure the segmented objects belonged to individual cells. Masks were applied to the corresponding channel and a measure tool gave the mean intensity for every single object. For cytoplasm signal quantification masks were obtained by subtracting DAPI mask to the whole cell mask, which was created thanks to Tomato membrane marker. Signal intensity measuring was performed as in the nuclei.

14.2. Video editing

Time-lapse videos were edited using Imaris x64 software. Imaris allowed to generate a video file from lsm format files (Zen lite files from Zeiss microscopy). Videos were re-edited with Adobe After Effects software.
14.3. 4D tracking workflow

Image processing and analysis methods were developed using Definiens Developer version XD2.4 (Definiens AG, Germany) and Matlab (R2015a, Bioinformatics, Image Processing, Optimization, Statistics and Machine Learning toolboxes). A manual-editing tool was implemented, allowing the user to visualize and correct errors. Quantitative cellular parameters in the pipeline are automatically extracted from the segmented and tracked nuclei, generating tables ready for data mining. The tables contain information on cell and nuclear position and shape, nuclear GFP intensity, cell neighbourhood, nuclear size, mitosis and apoptosis. The complete set of primary data generated is in Table 1.

During time-lapse analysis, photobleaching of GFP-Myc fluorescence produced a progressive signal reduction throughout the first third of the tracking period (Figure 8). After the initial photobleaching, the GFP-Myc signal was stable during the remainder of the tracking period (Figure 8). To avoid interference from photobleaching during the analysis, we used established methods (Gopinath et al., 2008) to normalize the values, using the endpoint mean and standard deviation as reference values. All quantitative analyses were made using the normalized data. This procedure allowed to perform meaningful in silico synchronizations.

![Figure 8. Correction of GFP-Myc signal after photo-bleaching.](image)

Graph displays individual data (dots) and locally weighted scatterplot smooth lines showing raw and corrected GFP-Myc levels of all recorded data synchronized to the start of time-lapse analysis.
15. Statistical analysis

Comparisons and graphs were made with either GraphPad Prism 5.0a or NCSS 11 Statistical Software (2016). Statistical analysis have been included in Figure legends.
1. Myc is the main driver of spontaneous cell competition in ESCs

Myc has been reported to be involved in natural cell competition in the epiblast and ESCs (Claveria et al., 2013). Nevertheless, taking into account the similar regulation and cooperation in development and ESCs between Myc and Mycn (Malynn et al., 2000; Scognamiglio et al., 2016), we also wanted to explore the implication of Mycn in this phenomenon.

It has been shown that Mycn can largely replace most Myc functions necessary for normal murine growth and development (Malynn et al., 2000). We thus explored whether Mycn could be compensating for Myc in ES cells. To address this, we first analysed the correlation between Myc and Mycn endogenous expression levels (Figure 9 A). Interestingly, in similarity to Myc, Mycn expression also showed a heterogeneous pattern in ESC cultures (Figure 9 A). However, quantification of both Myc and Mycn did not show any negative or positive significant correlation between both signals (Figure 9 A, B). This suggests that Mycn does not depend on Myc activity or undergoes compensatory regulation depending on Myc levels in ESCs.

Engulfing of loser cells by neighbouring winner cells has been observed in mESC and epiblast cells (Claveria et al., 2013). It has been reported that in these winner-loser pairs Myc expression levels are much higher in the winner than in the loser cells (Claveria et al., 2013). To directly assess the implication of Mycn in cell competition, we then analysed Mycn expression in pairs of engulfing/engulfed cells (Figure 9 C). Quantification of Mycn levels of engulfing and engulfed cells showed no differences between them in Mycn expression levels (Figure 9 C, D).

Given that not always cell competition involves engulfing events, we also examined caspase3-positive cells for Mycn levels (Figure 10 A). In WT cultures, Myc levels of apoptotic cells were reduced by half in comparison with Myc levels of non-apoptotic cells (Figure 10 B), as already shown in previous works (Claveria et al., 2013). On the contrary, Mycn levels did not show any difference between the apoptotic and non-apoptotic population (Figure 10 C). These data suggest that Mycn is not involved in ESC competitive behaviour.
**Figure 9. Mycn expression pattern and its implication in cell competition.**

A. Co-localization of Myc and Mycn in ESCs. B. Quantification of the correlation between Myc and Mycn immunofluorescence in single cells of ESC cultures. C. DAPI staining of an ESC colony; The magnified frame shows an engulfing event and Mycn immunofluorescence. D. Mycn level paired ratios in engulfing and engulfed ES cells in WT culture. Dotplot in D shows mean and standard error of the mean (SEM) and one sample t-test was used for statistical analysis; ns, adjusted p-val>0.05.
These observations suggest that, even though endogenous Mycn is not homogeneously expressed in ESCs, the differences in Mycn levels between cells are not determinant in spontaneous cell competition in ESCs. Myc is thus the only factor of its family in determining the competitive ability of ESCs.

Figure 10. Study of Mycn role in apoptotic-mediated cell competition.

A. Confocal detection of caspase-3, Myc and Mycn immunofluorescence in an ESC colony. B. Myc levels in non-apoptotic (general population) and apoptotic (caspase-3 positive) cells. C. Mycn levels in non-apoptotic (general population) and apoptotic (caspase-3 positive) cells. Boxplots in B and C show median, 25 and 75 percentiles (boxes), 10 and 90 percentiles (whiskers). Mann Whitney test was used for statistics. ns, adjusted p-val>0.05; p<0.001***.
2. Live imaging analysis of Myc levels in mESCs

2.1. The GFP-Myc knock-in line faithfully reports endogenous Myc expression

Although the implication of Myc in cell metabolism and proliferation in ESCs (Hu et al., 2009; Kim et al., 2010; Claveria et al., 2013; Gu et al., 2016; Scognamiglio et al., 2016) and cell reprogramming (Cartwright et al., 2005; Takahashi and Yamanaka, 2006) has been studied, the biological basis and dynamics of Myc-mediated cell competition remains elusive. To address this question, we used live reporting and tracking of endogenous Myc levels in mouse pluripotent cells. As a tool for this we derived a homozygous ESC line from previously described knock-in mice expressing a GFP-Myc fusion protein from the Myc locus (Huang et al., 2008) (see Material and Methods).

In the first place, as Myc protein (Cartwright et al., 2005) and mRNA (Dani et al., 1984; Hann and Eisenman, 1984) have very short half-lives, we wanted to determine the reliability of this GFP-Myc knock-in line in reporting endogenous Myc levels. To this end, we studied the mouse embryo epiblast and found that GFP-Myc faithfully reported Myc levels in this tissue, despite the cell-to-cell highly heterogeneous Myc pattern (Claveria et al., 2013) (Figure 11 A, B). In addition, in ESCs we also found a strong and fast variation of Myc during mitosis (Figure 11 C). Both Myc, detected by immunofluorescence, and GFP-Myc levels showed a drastic decreased in the transition from prophase to metaphase and a sharped increase during late telophase, being almost undetectable during metaphase, anaphase and cytokinesis (Figure 11 C, D). Moreover, we observed that this decrease during mitosis matches an increase in T58-phosphorylation (Figure 11 D), which targets Myc for degradation (Welcker et al., 2004). During interphase, T58-phosphorylated Myc was not detected, which indicates that Myc is specifically degraded during mitosis. These results demonstrated that the GFP-Myc knock-in line is a reliable reporter for endogenous Myc expression, even when the protein is rapidly degraded and re-synthesized as occurs during mitotic phases.
RESULTS

A

B

C

D

early prophase
prophase
pro-meta-phase
metaphase
anaphase
telophase
late telophase

DAPI
5μm

pH3

anti-Myc

GFP-Myc

Myc-T58
2.1. Setting up 24h 3D+t time-lapse and computer workflow for analysis

In order to characterize Myc dynamics during endogenous cell competition we performed time-lapse experiments in which homozygous Myc-GFP cells were scanned in 3D during 24 hours. These 24h 3D+t time-lapses were then used to track all cells. As cells moved very actively in ESC colonies (video 1), 3D scans were acquired every 7 minutes in order to allow an accurate tracking (Figure 12 A). Nuclear GFP-Myc signal and tdTomato cell membrane marker were acquired with green (488nm) and red (555nm) channels respectively (Figure 12 B) in a Zeiss LSM 780 upright microscope (see more details in Materials and Methods).

Next, in collaboration with the Cellomics Unit from CNIC, we developed a computer workflow for automatically segmentation and tracking the 3D+t acquisitions (Video 2) (Fernández de Manuel et al, 2017). This software also allowed to inspect and correct manually any incorrect outcome from the automatic segmentation and tracking (Figure 12 C) (see more details in Materials and Methods). With this approach, we were able to track cells across cell division, generating lineage trees in which the information on the progenitor cell identity was incorporated to daughter cell identities (Figure 12 D, E; videos 2 and 3). In this way, we obtained not only lineage trees for every cell family in the analysed colonies, but also multiple quantitative parameters, such as nuclear volume, cell trajectories, shapes and their fluctuations, GFP-Myc levels and information on neighbour cells (Figure 12 E; Table 1).
Figure 12. Multiparametric 4-dimensional analysis of embryonic stem cells, including lineage analysis and tracking of endogenous Myc levels.

A. Live confocal microscopy set-up. B. Example of 3D raw data acquisition from live GFP-Myc, membrane-Tomato ESCs (top) and whole volumes 3D rendering from these images (bottom). C. 3D segmentation of the nuclei of the ESC colony in A, with GFP-Myc levels displayed as green brightness intensity. D. Example of the tracking of segmented nuclei during mitosis. E. Cell lineage tree deduced from the tracking data, displaying the temporal pattern of GFP-Myc levels.
3. Myc is mostly regulated by cell-intrinsic heritable features

3.1. Study of Myc stability along the tracking

As it has been shown above and described in previous works (Claveria et al., 2013), Myc is heterogeneously expressed both in ESCs and in the embryo. Therefore, we wanted to test what was the nature of this heterogeneity. We previously observed no detection of T58-phosphorylated Myc in ES cultures (excepting mitotic cells) (Figure 11 D), discarding any implication of Myc degradation in driving Myc heterogeneity.

We then determined the stability of Myc levels in the tracked cells. Classification of the whole cell population into three groups according to mean GFP-Myc level (top 30%, Myc-high (H); lower 30%, Myc-low (L); the rest, Myc-medium (M)) showed that Myc-H cells have 2.5X higher GFP-Myc intensity than Myc-L cells (Figure 13 A). Next, we performed in silico synchronisation of the cell cycle in order to determine the fluctuations of Myc during cell cycle. The resulting expression profile captured the drastic reduction of GFP-Myc levels during mitosis and the rapid recovery during telophase (Figure 13 B; Figure 11 C, D). In addition, mitosis was preceded by a moderate increase in Myc levels, and typical fluctuations were detected afterwards (Figure 13 B). However, the severe fluctuations during mitosis did not extend enough time to explain the cell-to-cell heterogeneity observed in GFP-Myc levels (Figure 13 A).

Next, we examined the stability of Myc-levels within the three groups previously established. During 24h tracking these groups revealed little variation (Figure 13 C). Interestingly, even after in silico synchronization, Myc levels were maintained through cell division, despite rapid Myc degradation and resynthesis (Figure 13 D). The fact that Myc levels are maintained through mitosis suggests that they are to some extent imprinted in ESC lineages.
RESULTS

Figure 13. Myc is to some extent imprinted in ESC lineages.

A. Distribution of GFP-Myc levels after *in silico* classification of ESCs into groups according to live-recorded average GFP-Myc levels before cell’s first division. B. Display of individual (dots) and spline smooth line (SSL) of GFP-Myc levels of all dividing cells *in silico* synchronized to cell division. C. Tracking of GFP-Myc levels (SSLs) in Myc-H, Myc-M and Myc-L ESCs during the observation period. D. Similar data to C, synchronized to cell division time. Boxplots show median, 25 and 75 percentiles (boxes), 10 and 90 percentiles (whiskers). Mann Whitney test was used for statistical analysis, p<0.001***.

3.2. Retrospective clonal analysis of Myc

Since from the time-lapse experiments we only have information of 24-hour periods, we decided to perform retrospective clonal analysis in ESCs. We used the iMOS\textsuperscript{WT} cell line (Claveria et al., 2013) to generate EYFP+ clonal cell marks by inducing recombination at very low rate (see Materials and Methods for further
Cells were cultured for additional 60 hours after 4-hydroxytamoxifen removal. We then determined the variability of Myc levels between clonal groups of cells and randomly chosen equivalent groups. We found that Myc levels showed much lower variation in clonally related cells than in randomly chosen cells, with an estimated heritability of 0.8 per cell division, indicating that Myc levels display high heritability (Figure 14 A, B). In contrast with the predominant regulation of Myc in somatic cells, in which extracellular signals such as nutrients, mitogens or growth factors are described to be the main regulators (Wierstra and Alves, 2008), the clonal analysis, together with the previous observations in time-lapse and the cell-to-cell heterogeneity in Myc levels, indicates that Myc is mostly regulated by some intrinsic heritable mechanism in ESCs. Thus, sub-families of Myc expression levels seem to coexist in ESC cultures and largely transmit their Myc levels to their progeny.

**Figure 14. Retrospective clonal analysis of Myc.**

A. Example of a clonal mark induced by iMOS<sup>WT</sup> activation in an ESC colony together with immunofluorescence detection of Myc levels. B. Variation of Myc levels in clonal groups and equivalent groups of randomly chosen cells. Boxplots show median, 25 and 75 percentiles (boxes), 10 and 90 percentiles (whiskers). Mann Whitney test was used for statistical analysis, p<0.001***.

This form of heterogeneity is thus different from that reported for other transcription factors, such as Nanog, which has been shown to be regulated by stochastic transitions from low to high states of Nanog expression (Chambers et al.,...
RESULTS

2007; Kalmar et al., 2009). Moreover, quantification of both Myc and Nanog expression in ES cells did not show any significant correlation between both transcription factors (Figure 15 A, B).

![Figure 15. Myc and Nanog expression do not correlate.](image)

A. Confocal detection of Myc and Nanog immunofluorescence in an ESC colony showing colonization of both signals. B. Quantification of the correlation between GFP and Myc immunofluorescence in single cells of ESC colonies.

4. Temporal integration in losers of discrepant Myc levels with neighbours leads to cell competition

4.1. Identification and characterization of apoptotic cells

To study how MYC heterogeneity leads to cell competition we performed neighbour analysis by manual inspection using the computer workflow. First, we tracked apoptosis events during the 24-hours time-lapse analysis, identifying 24 apoptotic cells, that we called prospective-dying cells (PDC) (videos 4-6). Three of these apoptotic events were discarded from further analysis as they were considered Myc-independent death events. First, we analysed the average Myc-GFP level of
PDCs during the observation time, finding that, in agreement with previous works of Myc-driven competition (Claveria et al., 2013; Sancho et al., 2013), it was lower than that of the non-dying cells (NDCs) (Figure 16 A) and that it approximately matched the level found in MYC-L cells (Figure 13 A). Moreover, PDCs showed persistently lower GFP-Myc levels than NDCs, as revealed by the temporal profile of GFP-Myc levels synchronized to the time of PDC death (Figure 16 B). These results indicate that pre-existing low Myc levels correlate with the loser fate. Interestingly, this synchronization also revealed that PDCs show a tendency to reduce their Myc-GFP levels starting approximately 500 minutes before death (Figure 16 B). For comparison, we characterized a group of cells that showed similar Myc-GFP levels to those found in PDCs (Figure 16 C) but did not die during the tracking; this group was named Myc-L non-dying cells (ML-NDCs). These cells also showed continuously lower Myc-GFP levels than the general population during the observation time but did not undergo any GFP-Myc level reduction (Figure 16 D).
4.2. Analysis of PDC neighbourhood

We then performed neighbourhood analysis of the GFP-Myc expression profiles. NDCs showed a GFP-Myc profile similar to that of their neighbours (Figure 17 A). In contrast, in PDC neighbours GFP-Myc levels are initially lower than those of the general population but rise from 800 minutes before loser death up to matching the levels in NDCs and maintain a plateau until the death event (Figure 17 B, E). The increase in PDC neighbours’ GFP-Myc levels therefore precedes the decline in PDC GFP-Myc levels (Figure 17 B). The difference in GFP-Myc levels between PDCs and their neighbours was thus exacerbated before the death event (Figure 17 D). Conversely, ML-NDCs neighbours did not show any increase in their GFP-Myc level profile and they stably maintained lower levels than NDCs (Figure 17 C). The contrast in GFP-Myc levels between ML-NDCs and their neighbours is therefore always below that preceding PDCs death (Figure 17 D). Furthermore, by synchronizing ML-NDCs neighbours to the timepoints in which their GFP-Myc levels increase to match those of the plateau preceding PCD death, we observed that ML-NDC neighbours did not experience any increase in their GFP-Myc levels extending longer than 500 minutes (Figure 17 F). These results suggest that Myc-low cells survive in ESC colonies as long as the discrepancy in Myc expression with their neighbours is moderate. However,
sustained high discrepancies for periods above 600–800 minutes lead to their elimination.

**Figure 17. Analysis of Prospective Dying Cells neighbourhood.**

A. Temporal evolution of GFP-Myc levels in NDCs and their neighbour (means). B. Temporal evolution of GFP-Myc levels in PDCs and PDC-neighbour means synchronized to PDC death. Significant increases between consecutive 100-minute sections are shown (arrowheads). C. Temporal evolution of GFP-Myc levels in Myc-Low non-dying cells (ML-NDCs), ML-NDC neighbour means and NDCs. D. Temporal evolution (locally weighted scatterplot smooth line) of the ratios between GFP-Myc levels in NDCs, PDCs, ML-NDCs and their respective neighbour means. E. Temporal evolution of Myc-GFP levels in NDCs and PDC neighbour means. F. GFP-Myc levels of ML-NDC neighbours synchronised to the initiation of periods in which their mean level is equal or higher to that shown by PDC neighbours before PDC death. A–C, E–F display SSLs. Pink areas between graph lines in A–C, E indicate p-values <0.01**; 0.001***; 0.0001****. Statistical analysis in A–C, E were performed by comparing 100-minute sections between the represented data sets, using a Wilcoxon matched-pairs signed rank test for comparing PDCs, NDCs and ML-NDCs with their neighbours and a Mann Whitney test for the rest of comparisons. Longitudinal changes of PDCs neighbours shown in B were studied using a Mann Whitney test to determine significant changes between consecutive 100-minute sections centred at 50-minute sliding positions. Transitions between temporal sections showing significant differences are shown by an arrowhead and indicate the p-value; p-val<0.05*; 0.01**, 0.001***.
5. Random contacts between cells with pre-existing discrepancy in Myc levels drive cell competition

Given that the first event anticipating a loser-cell fate is the increase of Myc level in its neighbourhood, we decided to study the nature of this increase. The results observed could correspond to replacement of neighbours by cells with higher Myc levels than their own, or to Myc upregulation in pre-existing neighbours. To analyse these options, we inspected individual GFP-Myc profiles of PDC neighbours and found that the increase in neighbour’s aggregate GFP-Myc levels normally correspond to the presence of only one neighbour at a time with very high GFP-Myc levels (Myc-highest neighbours, MHNs) (see examples in Figure 18 A). The total number of contacts contributing to the GFP-Myc plateau and the duration of these contacts were highly variable (Figure 18 A, B), indicating that MHNs do not engage in the interaction with PDCs. When we represented GFP-Myc level temporal profiles considering only MHNs we observed similar profiles to those including all neighbours (Figure 17 A-C), but with a wider discrepancy in GFP-Myc levels (Figure 18 C-E). In addition, we synchronised the complete temporal profile of PDC neighbours to the time of their first contact with PDCs and we found that Myc levels in neighbours are insensitive to these contacts (Figure 18 F, G). We also found no correlation between the frequency of the contacts and the cell cycle phase (Figure 18 B). Moreover, the distribution of the duration of PDC contacts with their MHNs was similar to that observed with their Myc-low neighbours (Figure 18 H), which indicates that there is no specific pattern of direct contacts between prospective winner and loser cells.

These results indicate that it is neighbour exchange and not Myc regulation what produces the increase in GFP-Myc levels that precedes cell death (see example in video 7); therefore, spontaneous cell competition is driven by apparently random contacts between ESCs that autonomously express highly discrepant Myc levels.
6. **iMOS-mediated induction of cell competition confirms the conclusions from the live analysis of spontaneous cell competition**

Our results suggest that Myc-low cell elimination takes place when accumulated contacts lead to sustained discrepancies for longer than 600-800 minutes and is preceded by a progressive decrease of Myc levels in the loser cell.

To test this model, we used the *iMOS* system (Claveria *et al.*, 2013) to experimentally induce cell competition by overexpressing Myc in a mosaic manner (see Material and Methods). We performed a time-lapse experiment in which we studied the temporal profile of EYFP-Myc cell induction and neighbour death events (Figure 19 A-C; video 8, 9; see Material and Methods). In accordance with the observations from the live-analysis of spontaneous cell competition, we found an increase over 3-fold of the cell death rate at 600 minutes from the onset of EYFP-Myc induction (Figure 19 D). Furthermore, when we quantified the death events that were not in contact with EYFP-Myc cells, the temporal profile did not exhibit any increase.
in the death rate, which remained linear during the observation time (Figure 19 D). In addition, this observation is in accordance with previous works reporting that cell competition is cell-to-cell contact dependent (Simpson and Morata, 1981; Moreno et al., 2002; Moreno and Basler, 2004; Li and Baker, 2007; Claveria et al., 2013).

Figure 19. *iMOS*-mediated induction of cell competition.

A, B. Confocal images showing the EYFP pattern in *iMOS*<sup>T1-Myc</sup> ESCs at two time-points during the time-lapse after conditional activation. EYFP reveals cells overexpressing Myc. Red arrowhead indicates negative cells and white arrowhead indicates EYFP-Myc cells. C. Selected frames from live analysis of *iMOS*-induced cell competition showing apoptosis of non-activated cells neighbouring activated *iMOS*<sup>T1-Myc</sup> cells. D. Cumulative temporal representation of EYFP activation events and cell death events in non-EYFP cells - in contact or not with *iMOS*<sup>T1-Myc</sup> cells - after *iMOS*<sup>T1-Myc</sup> activation. s, slope (events/hour). E, F. Confocal images showing the EYFP and GFP-Myc patterns in *iMOS*<sup>T1-Myc</sup> / GFP-Myc mixed ESC colonies at two time-points during time-lapse analysis after conditional activation. Widespread EYFP reveals cells overexpressing Myc and nuclear GFP reveals GFP-Myc cells. White arrowhead indicates EYFP-Myc cells, grey arrowheads indicate a PDC and red arrowheads indicate a NDC, both neighbours of EYFP-Myc cells. G. Cumulative temporal representation of EYFP activation events and GFP-MYC levels in PDC and NDC neighbours of EYFP-Myc cells in *iMOS*<sup>T1-Myc</sup> / GFP-Myc mixed ESC cultures following *iMOS* induction. D, H display locally weighted scatterplot smooth lines.
Next, in order to study the evolution of GFP-Myc levels in EYFP-Myc cell neighbours, we repeated the experiment but co-culturing the iMOS\textsuperscript{Tir}-Myc cell line with the GFP-Myc knock-in line (Figure 19 E-G; video 10). We then identified and traced the Myc levels of GFP-Myc cells that were neighbours of EYFP-Myc cells and died during the observation period. We found that they showed a sustained decreased in their GFP-Myc levels starting shortly after EYFP-Myc induction (Figure 19 H). On the contrary, GFP-Myc levels of EYFP-Myc neighbour cells that did not die during the tracking were stable during the total period of observation (Figure 19 H). These results confirm the conclusions from the live-analysis of spontaneous cell competition.

7. The competitive ability of ESCs correlates with Myc levels but not with proliferation

We next questioned what was the biological basis of different Myc levels in ESC sub-lineages. It has been previously shown that deletion of both Myc and Mycn or pharmacological inhibition of Myc activity leads to proliferation arrest by decreasing transcription, splicing and protein synthesis (Scognamiglio \textit{et al.}, 2016). We then studied the correlation between Myc levels and proliferation and its implication in cell competition. The live-imaging analysis allowed us to measure the proliferative activity of each cell sub-lineage. We observed that the Myc-Low population exhibited a much lower fraction of the population in S-phase than Myc-High cells (Figure 20 A) and a higher fraction of cells not dividing during tracking (Figure 20 B). This means that Myc-L cells proliferate less that the rest of the population. In contrast, we found no difference in the proliferative activity between Myc-M and Myc-L cell populations (Figure 20 B). Moreover, in cells that divided during the period of observation, GFP-Myc levels did no correlate with the frequency of mitosis (Figure 20 C). In addition, we were able to track cell cycle progression by synchronizing nuclear sizes to the moment of mitosis. This analysis showed that nuclear volume increased linearly during interphase and halved across mitosis (Figure 20 D), showing that the increase in nuclear size reports the S-phase of the cell cycle. These observations agree with the extremely short G1 and G2 phases in the ESC cycle (Savatier \textit{et al.}, 1994; Boheler, 2009). Study of the temporal profile of nuclear size increase indicated similar cycling speeds for Myc-H, Myc-M and Myc-L
cell populations (Figure 20 E). These results suggest that Myc levels over a certain threshold are required for ESCs to enter the proliferative status, but once in proliferation, Myc levels do not determine the cell cycle speed.

Interestingly, both dividing and non-dividing Myc-L cells showed substantially increased apoptosis rates (Figure 20 F) meaning that the proliferative status of Myc-L cells did not affect their chance to undergo apoptosis. This indicates that ES cells compete according to their Myc levels irrespective of their cycling activity.

**Figure 20. Myc levels and proliferation in ESCs.**

A. DNA content cytometry profiles. B. Percentage of dividing and non-dividing cells during time-lapse analysis in ESCs classified according to Myc levels. C. Average GFP-Myc levels in ESCs dividing once or twice during tracking. D. Display of individual (dots) and SSL of nuclear size tracking in dividing cells synchronized to cell division time (time 0). E. Similar data as in D, in ESCs classified by Myc levels. F. Frequency of apoptosis in Myc-L ESCs according to their proliferative behaviour during tracking. Boxplots in C show median, 25 and 75 percentiles (boxes), 10 and 90 percentiles (whiskers). Mann Whitney test was used for statistical analysis. ns, p>0.05.
8. Study of cellular dynamics and Myc levels in ESC colonies

We also used the tracking data to examine several parameters related with cell motility and its correlation with GFP-Myc levels. We compared those parameters between the Myc-H and Myc-L cell populations. We found no difference between both populations in cellular speed, total distance travelled and net displacement (Figure 21 A-C). These results indicate that differences in cell motility are not involved in the competitive behaviour, supporting the idea of random and passive contacts between winners and PDCs.

Figure 21. Myc levels and cellular dynamics in ESCs.

A. Measurement of cell speed between Myc-L and Myc-H populations. B. Measurement of total displacement between Myc-L and Myc-H populations. C. Measurement of net displacement between Myc-L and Myc-H populations. D. Measurement of cell roundness between Myc-L and Myc-H populations. E. Measurement of cell roundness variation (deformability) between Myc-L and Myc-H populations. Boxplots show median, 25 and 75 percentiles (boxes), 10 and 90 percentiles (whiskers). Mann Whitney test was used for statistics. ns, p-value>0.05; p<0.05*.
We also measured cell roundness and we observed no difference in this parameter between Myc-H and Myc-L cells (Figure 21 D). Remarkably, Myc-H cells did show higher variance of roundness over time than Myc-L cells (Figure 21 E), which indicates higher cell deformability associated with higher Myc levels.

9. Myc expression levels correlate with the pluripotency status

Since we wanted to study the biological origin of Myc-driven endogenous cell competition, we next performed a transcriptome analysis of mouse ESCs. We sorted three sub-populations of cells according to their Myc levels (Myc-Low, Myc-Medium and Myc-High) (Figure 22 A), we extracted RNA of the three sub-populations and performed RNAseq. These groups were comparable to those artificially generated in the computer workflow for the previous studies (Figure 13 A). After examination of the transcriptome, we found that the most significant difference between the populations was the regulation of the pluripotency network (Figure 22 B). We then decided to compare our transcriptome of ESCs classified by Myc levels with those of the naive and primed pluriplontent states. We used RNAseq analysis from previous works that described these pluriplontent states in naive ESCs and epiblast-like cells (EpiLCS) (Buecker et al., 2014; Sasaki et al., 2015). EpiLCS are cells in transition between naive and primed states that are accomplished in vitro (Hayashi et al., 2011).

Analysing principal components, we observed that Myc-L cells were closer to differentiation-primed EpiLCS, whereas the Myc-H population was more similar to naive ESCs and Myc-M cells were located at an intermediate position between both states (Figure 22 C). This observation was also supported by the examination of the main regulators of pluripotency (Martello and Smith, 2014) (Figure 22 D-E). The fact that the expression profile of the Myc-M population showed an intermediate position suggests that Myc levels correlate with changes in the transcriptome that occur in the transition from the naive to the primed primed pluripotential state.

Next, we examined whether Myc levels correlated with the activation of the two main signalling pathways that characterize the naive and primed pluripotency status. We found that in ESCs GFP-Myc levels negatively correlate with pERK expression (Figure 23 A, B), whose activation conveys acquisition and maintenance of the primed state (Kunath et al., 2007). Contrary, GFP-Myc levels correlate positively...
with pSTAT3 activation (Figure 23 C, D), which characterizes and promotes naive pluripotency state (Niwa et al., 1998; van Oosten et al., 2012). These observations further support that the differentiation status correlates with endogenous Myc levels in ESC culture and suggest that differentiation-primed lineages might be outcompeted by their naive neighbours.
Furthermore, these data agree with our experimental ESC culture conditions, (serum replacement and Leukemia inhibitory factor (LIF) without feeder fibroblasts), which allow heterogeneity in the pluripotency status.
DNA methylation (5mC) at CpG dinucleotides is an inhibitory epigenetic modification often associated with transcriptional silencing. Other roles of this modification in transposable element regulation, splicing and genome integrity have also been identified (Smith and Meissner, 2013; Hackett and Surani, 2014). DNA methylation mostly faithfully kept through cell divisions, stabilizing restriction of cellular identity, except during early development where 5mC is dynamically remodeled (Smallwood et al., 2011; Smith et al., 2012; Hackett and Surani, 2014). Naive pluripotent cells of the early epiblast show very low levels of methylation because of the activity of de-methylating enzymes (TET proteins) and global eradication of DNA methylation is needed to eliminate epigenetic barriers against the acquisition of naive pluripotency (Hackett et al., 2013). Previous works have reported that naive ESCs show globally hypomethylated DNA when derived in 2i/LIF. However, ESCs cultivated in serum exhibit increased levels of DNA methylation typically associated with the primed status and further differentiation into specific lineages typical of the late epiblast (Hackett and Surani, 2014). In fact, ESCs exhibit 3-fold higher 5mC in serum in comparison with 2i/LIF culture medium (Ficz et al., 2013; Habibi et al., 2013; Leitch et al., 2013). Therefore, ES cells maintained in 2i/LIF conditions show similar levels and distribution of 5mC to the naive epiblast cells, whereas cultures grown in serum contains not only naive cells but also cells that are closer to primed epiblast cells and show significant DNA methylation.
As reflected in our RNAseq analysis, our ESC culture conditions allow pluripotency heterogeneity, with some cells closer to naive status and others closer to differentiation-primed status, which associates with Myc expression levels. We thus tested whether experimental epigenetic modification of the methylation status would affect Myc levels. To address this question, we used the drug 5-Azacytidine, which
inhibits DNA methyltransferases, causing DNA hypomethylation. We observed that Myc levels of GFP-Myc ESCs did not show any variation after 10 hours of treatment (Figure 24 A). Notably, ESCs treated with Azacitidine for 24 hours did show an increase in their Myc-GFP levels (Figure 24 B; video 11). In addition, we measured Myc-GFP levels at 24 hours after treatment removal and observed a partial recovery of Myc expression levels in the absence of any treatment (Figure 24 C). These data suggest that Myc levels are controlled to a certain extent by DNA methylation and, together with the correlation between Myc and the pluripotency status, suggests that differentiation-associated methylation is one of the factors that determines the heritable modification of Myc levels in ES cell sub-lineages.

Next, we examined whether the increase in Myc levels mediated by 5-Azacitidine could trigger cell competition. To this end, we cultured GFP-Myc ESCs with 0.25 μM 5-Azacitidine for 24 hours and confronted them with membrane-tomato GFP-Myc ESCs kept in control medium for several days (Figure 24 D). We then analysed the proportions of both populations in coculture and found that the proportion of the treated cell population –exhibiting higher Myc levels- did no increase when confronted with a control cell population (Figure 24 E). This observation shows that increasing Myc level by exposure to the demethylating agent 5-Azacitidine is not sufficient to induce cell competition, suggesting that pleiotropic effects of the drug abrogate ESC competitive ability.

11. 2i culture conditions reduce Myc heterogeneity and blocks endogenous cell competition

As previously shown, our experimental ESC culture conditions allow heterogeneity in pluripotency status. In addition, since LIF limits the transition from naive to primed pluripotent states and the primed state is not reversible by changes in the signalling environment (Guo et al., 2009), our observations are also in accordance with the heritability of Myc levels.
Previous work reported that ESCs cultured with MAPK and GSK inhibitors (2i) maintain the naive state, giving rise to cultures expressing pluripotency markers in a more homogeneous manner (Ying et al., 2008). As we showed that cell competition takes place when cells with discrepant Myc levels are confronted and these levels correlate with the pluripotency status, we examined the effect of 2i culture conditions on Myc expression and the competitive behaviour of ESCs. To address this question,
we cultured GFP-Myc ES cells with or without 2i for three days. We found that Myc level heterogeneity was reduced in 2i conditions (Figure 25 A). Next, we studied whether the decrease in Myc heterogeneity led to a change in the competitive ability by measuring the frequency of apoptotic events and Myc levels of those apoptotic cells. Interestingly, the death frequency was 3-fold less in the 2i conditions with respect to serum+LIF culture conditions (Figure 25 B). We then measured Myc levels in non-apoptotic cells and in dying cells in the first phases of apoptosis, when they present active caspase-3 staining but still intact nuclei. As expected (Claveria et al., 2013), in serum replacement+LIF culture conditions Myc levels were higher in non-apoptotic cells than in early apoptotic cells (Figure 25 C). However, in 2i culture conditions there was no significant difference in Myc levels between apoptotic and non-apoptotic cells (Figure 25 D). These results show that, in similarity to the core pluripotency factors, Myc expression heterogeneity is reduced in ESCs cultured in 2i conditions, which further supports the idea of the correlation between Myc and the pluripotency status. Moreover, Myc-mediated spontaneous cell competition does not take place in naive ESC cultures where discrepancies in Myc levels between cells are reduced.

12. **Cell competition can be experimentally induced by confronting cells with different pluripotency status**

We showed in previous experiments that differentiation-primed cells might be eliminated by their naive neighbours. To directly test this hypothesis, we induced ES cells to differentiate by LIF withdrawal and confronted them with naive ES cells kept in LIF, plus 2i (Figure 26 A). We analysed the proportions of both populations over time and found that the differentiating ESC population was outcompeted by naive ES cells (Figure 26 B). These differentiating ES cells were otherwise viable when cultured alone, but the frequency of apoptosis increased almost 3-fold when co-cultured with naive cells (Figure 26 C), indicating that the elimination of the differentiating population occurs through cell competition. Moreover, in agreement with previous studies (Sumi et al., 2007) and our RNAseq analysis, the differentiating population exhibited lower GFP-Myc levels than the naive population in co-culture (Figure 26 D, E), which confirmed the expected correlation between Myc levels and competitive ability. These observations support that Myc levels associate with different states of
pluripotency in ESCs and that the confrontation of Myc-high naive cells with Myc-low differentiation-primed cells gives rise to the elimination of the latter through cell competition.

**Figure 26. Cell competition induced by confronting population with different pluripotency status.**

A. Procedure for confronting differentiating ESCs (Tomato⁺) and naive ESCs (Tomato⁻). B. Frequency of Tomato⁺ and Tomato⁻ cell populations after different periods of co-culture. C. Frequency of caspase3-positive cells in the differentiating Tomato⁺ cell population alone or in co-culture with naive ESCs. Data were collected by microscopic fields (N). D. Immunofluorescence study of Myc levels after 2-day co-culture of Tomato⁻ ESCs pre-cultured for 3 days with LIF+2i and Tomato⁺ ESCs pre-cultured for 3 days without LIF. E. GFP-Myc levels in Tomato⁺ and Tomato⁻ cell populations in co-culture. Column bar graph in C show mean and standard error of the mean (SEM). Boxplot in E show median, 25 and 75 percentiles (boxes), 10 and 90 percentiles (whiskers). Mann Whitney test was used for statistical analysis. p-value<0,05*; p<0,001***.
13. Blocking cell competition results in the accumulation of differentiation-primed cells

We next explored whether cell competition could be acting as a mechanism to prevent the accumulation of cells that are prone to differentiate in ESC cultures. To address this question we used the iMOS<sup>T<sub>1</sub>-p35</sup> ESC line (Claveria <em>et al.</em>, 2013), in which following tamoxifen addition, a mosaic of EYFP- and ECFP-expressing cells is induced and apoptosis is inhibited in the ECFP population because of the expression of the caspase inhibitor p35, whereas the EYFP population remains WT and is allowed to undergo apoptosis (Figure 27 A). ECFP protein was not well detected in the microscope, therefore we recognize ECFP positive cells by doing immunostaining with anti-GFP antibodies, which recognize both ECFP and EYFP, so that cells positive for anti-GFP staining and negative for EYFP were identify as ECFP p35-expressing cells (Figure 27 B).

We then compared the death-blocked population 48 hours after the induction (see Material and Methods) with the wild type neighbours and found that the rescued cells showed a higher proportion of active-pERK levels than the control population (Figure 27 B, C). This result suggests that cell competition might be acting in maintaining ESC cultures in a naive pluripotent state by restricting the differentiating population.
14. Differentiating-primed cells are eliminated from pre-gastrulation embryos through cell competition

14.1. Myc/pERK regulation in the epiblast

We then studied whether this observed mechanism applies to epiblast cells in the early mouse embryo. To this end, we first examined the pattern of pERK activation in early postimplantation embryos. We found that at E5.0 the epiblast did not contain any pERK positive cell (Figure 28 A). In contrast, E6.0 embryos showed around 1-4% of isolated cells positive for pERK in an apparently random pattern (Figure 28 B-D). This pERK positive cell population exhibited lower Myc levels than the pERK negative cells (Figure 28 C, D), suggesting a similar relationship in vivo as that found in ESCs.

Next, we studied the pattern of pERK expression in early gastrulating embryos. The embryo gastrulates by the recruitment of epiblast cells to the primitive streak. During this process, a coordinated loss of pluripotency occurs in the vicinity of the primitive streak for mesoderm formation. At this embryonic stage, we found a regional expression of pERK and downregulation of Myc close to the primitive streak region (Figure 28 E). These results suggest that Myc/pERK regulation in the epiblast correlates with the patterns found in ESC cultures.
Figure 28. Analysis of pERK activation in the epiblast and correlation with Myc levels.

A. Confocal detection (optical sections) of pERK immunofluorescence in an E5.0 mouse embryo showing no pERK+ cells in the epiblast. B. Confocal detection (optical sections) of pERK immunofluorescence in an E6.0 mouse embryo. White arrowheads show pERK+ cells. C. Co-localization by immunofluorescence of Myc and active pERK in the epiblast of E6.0 mouse embryos. White arrowhead shows a pERK+ cell exhibiting low GFP-Myc levels. D. Scoring of detected active pERK+ cells in the E6 mouse epiblast. E. Normalized Myc level distributions in active pERK+ and pERK- cells of the mouse E6 epiblast. F. Co-localization by immunofluorescence of Myc and active pERK in the epiblast of E6 and E6.5 mouse embryos. EE, extraembryonic; Epi, Epiblast; PS, primitive streak. Boxplot in E shows median, 25 and 75 percentiles (boxes), 10 and 90 percentiles (whiskers). Mann Whitney test was used; p-value<0.01**.
14.2. pERK positive cells are accumulated when blocking apoptosis in early embryos

We next studied whether the population of isolated pERK+ cells is restricted by cell competition in the pre-gastrulation embryo, as it does in ESC cultures. To address this question, we used the iMOS system to induce mosaics of p35-expressing cells and WT cells in embryos. We analyzed embryos at E5.5-6 where pERK-activated cells were scarcely present in the wild type population (Figure 29 A). Interestingly, we observed that pERK positive cells specifically accumulated in the death-blocked p35-expressing population (Figure 29 B, C). This observation suggests that differentiation-primed cells are eliminated through apoptosis from the pre-gastrulation epiblast.

**Figure 29.** Cell competition inhibition and pluripotency status analysed using the iMOS\textsuperscript{T2-p35} mosaic approach in the epiblast.

A. Co-localization of EYFP and active pERK+ cells in the epiblast of E5.5 iMOS\textsuperscript{T2-p35} mouse embryos recombined with Sox2Cre. B. Scoring of detected pERK+ cells in the epiblast of E5.5 iMOS\textsuperscript{T2-p35} embryos recombined with Sox2Cre. C. Frequency of EYFP-WT and ECFP-WT cells in the 10 embryos. Boxplot in C show median, 25 and 75 percentiles (boxes), 10 and 90 percentiles (whiskers). Mann Whitney test was used for statistical analysis; p-value<0.05*.
15. Myc directly defines the ESC competitive ability irrespective of the pluripotency status

We next studied the functional relationship between Myc and the pluripotency status. To this end we used the \textit{iMOS}^{Tr-Myc} cell line, in which we induced Myc overexpression in a random subset of cells (Figure 30 A). This \textit{iMOS} mediated Myc overexpression is enough to induce cell competition as it has been described (Claveria \textit{et al.}, 2013) and we showed above (Figure 19). We found that after Myc-overexpressing cells in mosaic cultures did not decrease pERK levels in comparison to their WT neighbors; in fact, they underwent a slight increase (Figure 30 A, B).

In addition, a compared transcriptome analysis by RNAseq between EYFP-Myc and control populations from mosaic cultures (C. Claveria, unpublished) not only did not show promotion of the naive pluripotency signature but rather a mild inhibition of this (Figure 30 C). Conversely, ESC differentiation by LIF withdrawal leads to GFP-Myc downregulation (Figure 26 D, E), in agreement with previous studies (Sumi \textit{et al.}, 2007).
These results, together with the induction of the winner phenotype by the Myc induction (Claveria et al., 2013), suggest a linear pathway in which Myc lies downstream the differentiation status and Myc directly determines the competitive ability of ESCs without modifying the pluripotency status.
DISCUSSION
In this doctoral thesis we have studied how Myc is regulated and how its heterogeneous expression and dynamics leads to spontaneous cell competition in ESCs. Additionally, we have explored the biological relevance of Myc-dependent endogenous cell competition both in ESCs and in early mouse development.

Despite the key role reported for both Myc and Mycn during embryogenesis (Charron et al., 1992; Trumpp et al., 2001; Dubois et al., 2008) and their redundant and cooperative function in maintaining the ESC proliferative program (Malynn et al., 2000; Scognamiglio et al., 2016), we showed that Mycn expression does not correlate with Myc expression in ESCs and, contrary to observations with Myc (Claveria et al., 2013), that endogenous Mycn levels do not correlate with the winner/loser ESC status. These observations, together with the absence of Mycl in ESCs, indicate that Myc regulation is the main driver of spontaneous cell competition in ESCs, with no contribution from other members of the proto-oncogene Myc family.

Given the major role of Myc in cell competition in ESCs, in this thesis we have developed new time-lapse microscopy and image analysis tools to track Myc levels during cell competition. To this end, we used an ESC line expressing a GFP-Myc fusion protein that faithfully reports endogenous Myc levels. To our knowledge, this new developed technique for tracking ESC lineages and endogenous Myc levels in 3D+t is the first described in the field of study. In a recently published work ESCs are tracked in 2D in conditions in which cell-to-cell interactions are inhibited and therefore it is not suitable for cell-interactive studies (Filipczyk et al., 2015). Our methodology allowed us to track ESC lineages through cell division and under entirely interactive conditions.

Furthermore, cell competition in ESCs and in the mouse epiblast is regulated by Myc expression heterogeneity among cells, however due to the lack of a strategy for isolating and tracking live cells according to Myc levels, the biological basis of Myc-low cells outcompetition from ESC cultures and early embryos was unknown. In this thesis, these limitations have been overcome by implementing this new microscopy and image analysis tool. Moreover, we also tracked the temporal evolution of Myc levels during cell proliferation, which has never been reported in any cell type. This approach allowed us to generate cell lineage trees that contain
information on the history of Myc levels for each cell, together with 3D information that allows to determine neighbourhood relationships between cells.

Using this tool we studied the stability of Myc levels in different subpopulations defined by Myc levels, both during the observation period and through cell divisions. We showed that Myc expression levels in ESCs are imprinted to a large extent, such they are conserved through mitosis. Additionally, retrospective clonal analysis experiments suggest that Myc levels display high heritability (0.8 per cell division). These data allow the emergence of a new concept that we called heritable fitness and consists on the heritable competitive ability of cells based on the transmission of Myc levels in ESCs sublineages coexisting in culture. The concept of Myc heritability is different from previous knowledge on the main factors contributing to Myc regulation. While Myc is generally found to be regulated by extracellular signals in somatic cells (Wierstra and Alves, 2008), both the cell-to-cell heterogeneous pattern and the heritability of MYC expression levels in ESCs indicate an important contribution of cell-intrinsic heritable features, in Myc regulation in ES cells.

The computer workflow established also allowed us to track Myc expression levels in individual cells and its neighbours, which allows the study of the dynamics of endogenous cell competition in a quantitative manner. Using this methodology, we have demonstrated that cell competition is driven by random contacts between cells with high discrepancies in Myc levels. Low-Myc cells are thus able to survive in ESC colonies as long as the differences in Myc expression levels between them and their neighbours are moderate or transient, however sustained discrepancies for longer than 600-800 leads to the elimination of the loser cells. We observed that the Myc expression profile during the 600-800 minutes preceding death of the loser shows a plateau shape. This observation indicates that loser cells measure time in exposure to neighbours with a discrepancy in Myc levels above a certain threshold, rather than accumulated Myc discrepancy. In case loser cells were measuring accumulated Myc discrepancies, then shorter exposures to higher Myc levels would also lead to cell competition and the shape of the Myc expression profile in neighbour of loser cells would show a continuous positive slope instead of a plateau.
Nevertheless, as the computer workflow only tracks and measures Myc levels of neighbour cells whose membranes are in direct contact with those of the loser cells, we only explored cell-to-cell contact dependent cell competition. Our initial hypothesis agrees with previous work reporting that direct contact is needed for competitive interactions (Simpson and Morata, 1981; Moreno et al., 2002; Moreno and Basler, 2004; Li and Baker, 2007; Claveria et al., 2013). Furthermore, since in our induced cell competition time-lapse experiments cell death rate is only increased in the population in direct contact with EYFP-Myc overexpressing cells, we obtained here further evidence of the need for cell-to-cell contact to induce apoptosis of loser cells.

We also observed, both in endogenous and in induced cell competition, a tendency in PDCs to reduce their Myc levels approximately 500 minutes before death and just after the increase in their neighbours’ Myc expression levels. The nature of this decrease is not known yet; however, in our time-lapse analyses using a caspase-3 reporter, we observed that caspase-3 is activated only few minutes before the cell death suggesting that Myc reduction in PDCs is not due to the activation of caspase-apoptotic pathway. Thus, competitive interactions might be operating on pathways triggering Myc downregulation for extended periods in loser cells.

Our analysis shows that in ESC cultures Myc-low cells are continuously eliminated and replaced by Myc-high cells. If this process were continuous, after cultures would continuously evolve towards higher and more homogeneous Myc levels; nevertheless, ESCs cultures always exhibit heterogeneous levels of Myc. Thus, we hypothesize that continuous and spontaneous generation of low-Myc cells and their elimination by fitter cells should maintain the equilibrium between ESC populations with different Myc levels in ESC colonies.

We also explored the different roles and dynamics of loser and winner cells during cell competition. On one hand, previous works have shown the involvement of increased cell mixing activity of winner cells in Myc-induced cell competition in Drosophila imaginal discs (Levayer et al., 2015), which contrast with our observations of any obvious active role of either winner or loser cells. In our model, interactions between loser and winner cells result from apparently random passive
contacts. Nevertheless, Drosophila imaginal discs cells are organized in a columnar epithelium where different sublineages mix poorly, whereas in ESC cultures there is a constitutive intense mixing, suggesting that the different environments may explain the differences observed between these two models.

On the other hand, we observed that Myc-high cells show higher variance of roundness over time that Myc-low cells. This result agrees with previous works in mammalian cells in which the winner phenotype presents higher deformability mediated by RhoA and actomyosin (Sun et al., 2014b). In addition, engulfment of losers by winner cells, which may involve differences in cytoskeleton dynamics, was also observed in mouse ESC and epiblast cells (Claveria et al., 2013). Engulfment was also reported to be the main system to eliminate loser cells in the Drosophila (Li and Baker, 2007); nonetheless, this observation contrasts with other recent works also performed in this model (Levayer et al., 2015).

Transcriptome analysis and the subsequent study of the main pathways characterizing naive and primed states allowed us to correlate Myc expression levels with the pluripotency status. A pool of pluripotent cells is maintained in early mammalian embryos for several days, however pluripotency is a highly unstable state, suggesting that specific mechanisms should be in place for pluripotency maintenance during this period. In this thesis we described a mechanism in which differentiation-primed cells are eliminated due to discrepancies in Myc levels with surrounding naive pluripotent cells. The fact that differences in pluripotency status may drive cell competition agrees with previous works reporting outcompetition of BMP-receptor-defective cells from ESC cultures and mouse epiblast (Sancho et al., 2013). BMP signalling collaborates with STAT3 in maintaining the naive pluripotency status, and therefore BMP receptor mutants may not be able to maintain the naive pluripotent status, leading to Myc downregulation and cell competition.

However, since not all Myc-low cells show differentiation-primed signatures, the difference in the pluripotent status may only explain part of the Myc-low population. Therefore, it would be important to identify additional cellular features characteristic of the Myc-low population to achieve a fully characterization of endogenous cell competition in the epiblast. Moreover, we only explored the
phenomenon of cell competition during one phase of the development and this mechanism could be also playing a role in other stages of the early mouse embryo. In fact, previous works already reported apoptosis in the inner cell mass of the blastocyst (Plusa et al., 2008), much before the stage in which this thesis is focused. At that stage the primitive endoderm differentiates and segregates from the inner cell mass of the blastocysts, which is the second event of differentiation after trophoderm differentiation (Stephenson et al., 2012). In a first step, primitive endoderm cell specification takes place in a spatially disorganized manner within the inner cell mass, so that cells of the PE lineage and those of the epiblast are intermingled. In a second step, cells specified as primitive endoderm segregate towards the blastocoel lumen. Interestingly, cells specified as primitive endoderm that fail to segregate from the pool of pluripotent cells of the inner cells enter in apoptosis (Plusa et al., 2008; Stephenson et al., 2012). Moreover, Myc levels are heterogeneously expressed also in the inner cell mass of blastocyst (Claveria et al., 2013) and is downregulated when cells differentiate to primitive endoderm (Smith et al., 2010). All these observations suggest that Myc-driven cell competition triggered by discrepant differentiation status may not only occur in the context described here, but may function as a more general mechanism affecting other developmental stages.

Additionally, recent work reports an apoptosis-dependent selection process during the colonization of the blastocyst inner cell mass by injected ESCs. In this model, naive ESCs preferentially colonize the inner cell mass, while primed ESCs are preferentially eliminated from the embryo by apoptosis (Alexandrova et al., 2016). This observation agrees with the model described here, however, in this work, ESC elimination does not correlate with Myc levels in the inner cell mass cells, which contrasts with the direct role of Myc in cell competition in the early epiblast and ESCs.
In ESCs differentiation-primed Myc-low cells are continuously outcompeted and replaced by naive Myc-high cells. However, there is a spontaneous generation of these primed low-Myc cells that maintains equilibrium in ESC colonies.

In the embryo cell competition is preserving the naive pluripotent status of the epiblast by eliminating cells in transit to differentiation before the time is right for differentiation. In contrast, when epiblast gastrulates a coordinated differentiation occurs in the posterior part of the epiblast and Myc is simultaneously downregulated, which allow controlled differentiation with no competitive interactions.
Myc has a role during induced pluripotent stem cell generation in promoting cell reprogramming of somatic cells in cooperation with transcription factors of the core pluripotency network (Takahashi and Yamanaka, 2006); in this thesis we described a mechanism that could help to better understand Myc implication in this process. First, we show that enhanced Myc activity does not promote the naive pluripotency signature, which fits results from other studies (Kim et al., 2010; Gu et al., 2016; Scognamiglio et al., 2016). Second, we observe that pluripotent cell proliferation can be efficiently driven by low Myc activity suggesting that Myc expression levels would be hardly relevant for stimulating proliferation during reprogramming. Indeed it has been reported that reprogramming of somatic cells can take place at lower efficiency in the absence of Myc overexpression (Nakagawa et al., 2008), which is compatible with our findings suggesting Myc’s role in reprogramming relates to promotion of pluripotent cell selection rather than to direct stimulation of pluripotency or proliferation.

Taken altogether, we propose a model in which cell competition maintains the naive pluripotency status in mouse pluripotent stem cell populations by eliminating cells in transit to differentiation (Figure 31). We provide evidence of a linear pathway in which Myc levels are determined by the pluripotency status and set ESC competitive ability without affecting the pluripotency status. This mechanism provides protection of naive pluripotent cell pools from premature differentiation by eliminating differentiation-prone cells; therefore cell competition operates as a surveillance mechanism of the stem cell pools in the mouse epiblast and ESC groups. When isolated cells spontaneously start to differentiate in a context of predominant naive cells, the contrast in Myc levels between the differentiating cell and its neighbours induces their elimination. In the early epiblast, this mechanism controls the naive status of the epiblast cell pool before the time is right for differentiation. In contrast, when the embryo starts to gastrulate, coordinated differentiation occurs and Myc levels and differentiation priming are homogeneously regulated in the posterior area of the epiblast. We therefore hypothesize that at this stage, the homogenous decrease in Myc levels avoids differences between cells and programmed differentiation can take place without induction of cell competition (Figure 31).
1. Myc regulation is the main driver of spontaneous cell competition in ESCs, with no involvement of other members of the Myc family.

2. The GFP-Myc knock-in line faithfully reports endogenous Myc levels and the imaging tool established provides a complete and reliable set of biological data during Myc-dependent cell competition in ESCs.

3. Myc expression levels are depleted by specific degradation of the Myc protein during ESC mitosis.

4. Myc levels are largely imprinted in ESCs, being mostly regulated by cell-intrinsic heritable factors.

5. Passive contacts between cells with high discrepancies in Myc levels that extend longer than ~10 h trigger cell competition in ESCs.

6. The proliferative state of ESCs requires Myc levels over a certain threshold, however, Myc levels do not modulate the speed of proliferation in cycling ESCs. The competitive ability of ESCs is dictated by Myc levels irrespective of the proliferation rate.

7. Myc levels correlate with the differentiation status and are controlled by epigenetic control in ESCs.

8. Endogenous cell competition maintains the naive pluripotent status of ESC cultures by eliminating differentiating cells.

9. Myc lies downstream the differentiation status and directly determines the ESC competitive ability.

10. Myc-driven cell competition triggered by discrepant pluripotency status also occurs in vivo, where differentiation-primed cells are eliminated from the pre-gastrulation epiblast.
1. La regulación de Myc es el principal factor desencadenante de la competición celular espontánea en CMEs, sin implicación de otros miembros de la familia de genes *Myc*.

2. La línea de ratón *knock-in* GFP-Myc reporta fehacientemente los niveles de expresión endógenos de Myc y la herramienta de imagen desarrollada proporciona un completo y fiable conjunto de datos biológicos durante la competición celular dependiente de Myc en CMEs.

3. Los niveles de expresión de Myc disminuyen por degradación específica de la proteína Myc durante la mitosis en CMEs.

4. Los niveles de Myc son en gran parte “grabados” en CMEs, siendo en general regulados por factores heredables intrínsecos.

5. En CMEs la competición celular se desencadena por contactos pasivos entre células con grandes discrepancias en sus niveles de Myc que se extienden más de ~10 horas.

6. El estado proliferativo de las CMEs requiere unos niveles de Myc por encima de un cierto umbral, sin embargo, éstos no modulan la velocidad de proliferación de células en mitosis. La capacidad competitiva de las CMEs está determinada por los niveles de Myc independientemente de la tasa de proliferación.

7. Los niveles de Myc correlacionan con el estado de diferenciación y son controlados mediante regulación epigenética en CMEs.

8. La competición celular endógena mantiene el estado de indiferenciación de los cultivos de CMEs mediante la eliminación de células en proceso de diferenciación.

9. Myc se encuentra por debajo del estado de diferenciación y determina directamente la capacidad competitiva de las CMEs.
10. La competición celular dependiente de Myc y desencadenada por diferencias en el estado de pluripotencia también tiene lugar in vivo, donde las células en proceso de diferenciación son eliminadas del epiblasto en estadio de pre-gastrulación.


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**Cover page.** Confocal image of an ESC culture showing GFP-Myc levels (green) and tdTomato reporting the cell membrane (red).

**Back page.** Confocal image of an E6.0 mouse embryo showing tdTomato reporting the cell membrane (red), DAPI staining the nuclei (blue) and pERK immunofluorescence (green).