

# The versatile transcription factor Oct-1: A crucial protein in embryonic development and A key component of the stress cellular response.

Inauguraldissertation

zur

Erlangung der Würde eines Doktors der Philosophie vorgelegt der Philosophisch-Naturwissenschaftlichen Fakultät Der Universität Basel

Von

### Mathieu Dalvai

Aus Strasbourg, (Frankreich)

Basel, 2008

### Genehmigt von der Philosophisch-Naturwissenschftlichen Fakultät

auf Antrag von

Prof. Markus Affolter und Prof. Witold Filipowicz

Basel, den 24/04/2007 (datum der Fakultätssitzung)

Prof. Dr Hans Peter Hauri

(Dekan)

### Acknowledgements

I would like to thank Pr. Patrick Matthias, my supervisor, who helped me a lot during this last four years by his theoretical and practical advices which allowed me to focus on what's essential in science and also in life in general. Thanks also to the other thesis committee members, Pr. Markus Affolter and Pr. Witold Filipowicz. Thanks to Pr. Hans Schöeler and Fatima Cavaleri for their collaboration. At least, thanks to all facilities members who help me a lot.

The thesis is a long period with some pleasant times but also (sometimes....) with some difficult moments where you think that science is not for you and you could be a good shell-necklace seller in a pacific island.....It's during this black moment that your friends, your family, are there to help you in many ways. I would like to thank all of them, especially my parents, my brothers and my sisters in law, but foremost my wife Florence who I love very much and of course my little boy Gaël which was a great moral support.

Special thanks to my colleagues, for all the things that make work in the lab great fun, So-Hee, Camille, Chun, Gabi (women first) and also Teppei, Fabien, Alain, Markus and Oliver. Thanks also to all the former lab members Ralph, Boris, Yu and finally Alexander with whom I had a lot of serious and interesting discussions and drank some beers.

### Contents

## Summary of the thesis

1. Introdu	ıction23
1.1 Cell cy	cle and regulation23
1.1.1 C	ell cycle phases23
1.1.2 C	yclin-dependent kinase (CDK)27
1.1.3 C	yclin dependent kinase inhibitor (CDKi)31
1.2 p27 is	a key negative regulator of G1/S checkpoint
1.3 Cyclin	D1, an important actor in G1/S transition37
1.3.1 A	ctivation and activity of cyclin D139
1.4 Recep	tor tyrosine kinase and signal transduction43
1.4.1 Th	he RTK subclass 1: ERBB receptors45
1.5 The R/	AS/ERK signaling pathway47
1.5.1 R	AS/ERK signaling and cell cycle progression51
1.6 The PE	SK/AKT pathway53
1.6.1 PI	BK/AKT signaling: cell cycle progression and proliferation control 55
1.7 Cell cy	cle response to DNA damaging agent57
1.7.1 pt	53 pathway57
1.7.2 p	.2161
1.7.3 B/	AX63
1.7.4 G	ADD4565
1.8 The ub	biquitous transcription factor Oct-169
1.8.1 O	ct-1: cell cycle and stress cellular response75

 Conditions.
 .79

 2. Results.
 .81

 2.1 Abstract.
 .81

 2.2 Introduction.
 .83

 2.3 Results.
 .87

 2.4 Additional data.
 .123

 2.5 Materials and Methods.
 .127

Chapter 2: Oct-1 is an essential factor in embryogenesis and

Chapter 1: Oct-1 is necessary for cell cycle arrest under stress

is required for a proper extra-embryonic	tissues	
formation	135	
1. Introduction	137	
1.1 Summary of mouse development	137	
1.2 Gastrulation	139	
<b>1.3</b> Cross-talks between extra-embryonic and embryonic tissues	141	
<b>1.4 Transcription factors: ES pluripotency and trophectoderm differentiation</b> .149		
1.5 Primary data and objective of research projects	151	
2. Results153		
3. Discussion163		
4. Materials and methods169		
References175		

### Abbreviations

Aa:	amino acids
BrdU:	bromodeoxyuridine
CDC:	cell division cycle
CDK:	cyclin dependent kinase
CDKI:	cyclin dependent kinase inhibitor
CFSE:	carboxyfluorescein diacetate succimidyl ester
ChIP:	chromatin immunoprecipitation
CRM1:	chromosome region maintenance
DNA:	desoxyribonucleic acid
EGF:	epidermal growth factor
ERK:	extracellular-signal regulated kinase
ES cells:	embryonic stem cells
ExE:	extra-embryonic ectoderm
F,f allele:	flox allele
FACS:	fluorescence-activated cell sorting
FGF:	fibroblast growth factor
GADD45:	growth arrest DNA damage
GDP:	guanosine diphosphate
GTP:	guanosine triphosphate
Gy:	gray
H&E:	hematoxilin and eosin
ICM:	inner cell mass
KO:	knock-out

MAPK:	mitogen activated protein kinase
MEF:	mouse embryonic fibroblast
NEBD:	nuclear envelope break-down
PCR:	polymerase chain reaction
PGC:	primordial germ cells
PKB:	protein kinase B
Rb:	retinoblastoma
RNA:	ribonucleic acid
RT:	reverse transcriptase
RTK:	receptor tyrosine kinase
SH2:	sarcoma homology domain
TE:	trophectoderm
TGF-β:	transforming growth factor- $\beta$
TOR:	target of rapamycin
TS cells:	trophoblastic stem cells
UV:	ultraviolet
Wt:	wild type

#### Summary of the thesis

#### Chapter 1

Oct-1 is a transcription factor belonging to the POU family (Clerc et al. 1988) (Herr et al. 1988) (Ryan and Rosenfeld 1997). The members of this family are involved in a broad range of biological processes like transcription of housekeeping genes (Oct-1), pluripotency of embryonic stem cells (Oct-4) or development of immune system (Oct-1, Oct-2) (Spaniol et al. 1996). The transcription factor Oct-1 is ubiquitously expressed in embryonic and adult tissues, and regulates the expression of a variety of genes. Previous studies described Oct-1 to be regulated at the protein level by phosphorylation in a cell cycle dependent manner. In addition, more recently it has been shown that Oct-1 is induced in response to DNA damage and modulates the activity of genes like *GADD45* important for the cellular stress response (Segil et al. 1991) (Zhao et al. 2000) (Jin et al. 2001) (Fan et al. 2002) (Tantin et al. 2005)..

Knockout and a conditional *oct-1* knockout alleles were created in our laboratory, and used to generate Oct-1 deficient mouse embryonic fibroblast (MEFs) and embryonic stem cells (ES). We used them as a model to study the cellular response to stress in absence of Oct-1. We have found that cells lacking Oct-1 were less sensitive to stress like  $\gamma$ -irradiation, deprivation of glucose or amino acids but not to serum starvation or H2O2 treatment. Under glucose starvation, this effect is in part mediated by activation of the signalling protein ERK and PKB which leads to maintenance of the level of cyclin D1 protein in KO cells

15

comparable to the level without treatment in heterozygous or rescue cells. These proliferative signals allow to KO cells to bypass the G1 cell cycle checkpoint and to proliferate better. Under stress conditions like glucose starvation, Oct-1 regulates the cell cycle by blocking cells in G1 phase, and by controlling the expression of the cyclin dependent kinase inhibitor p27.

Chapter 1 of this thesis provides a general introduction about the regulation of cell cycle and, more particularly, about key proteins and important pathways of the progression and transition of G0/G1 and S phases. It also gives some background about pathways like PKB and ERK, or other signaling pathways like p53, which plays determinant roles in cellular responses after stress induction. The role of the transcription factor Oct-1 in cell cycle regulation in response to cellular stress is discussed in the results part.

### Chapter 2

The POU transcription factors family was identified after the isolation of three mammalian transcription factors and a *Caenorhabditis elegans* developmental regulator: Pit-1, Oct-1, Oct-2 and Unc-86 (Ryan and Rosenfeld 1997). Oct-1 and Oct-2 proteins show a high degree of identity (90%) but differ in their expression pattern. Oct-1 is ubiquitously expressed, whereas Oct-2 is restricted to B cells, macrophages, hematopoietic cells, as well as cells of the central nervous system (Gerster et al. 1987) (He et al. 1989) (Staudt et al. 1988). Despite several studies, the exact role of Oct-1 and Oct-2 in immunoglobulin genes regulation and B-cell development is not well understood. In order to

17

investigate the functions of Oct-1 in B cells, our laboratory created a deficient mouse for Oct-1. Mice homozygous for *oct-1 (-/-)* were generated by crossing heterozygous +/- mice. Unfortunatly, these matings failed to produce any live born homozygous *oct-1 -/-* mutant, indicating that Oct-1 is essential for embryonic mouse development.

In this second project, the result part of chapter 2 will focus on the lethal phenotype observed during embryogenesis in absence of Oct-1 and gives an overview of the possible role of Oct-1 during mouse embryonic development. We showed that the embryos die around 8 dpc due to an arrest of the ectoplacental cone and giant trophoblastic cells development which occurs at 6.5dpc. This extraembryonic deletion leads to the formation of an impaired mesoderm.

In introduction the chapter 2 gives a general background about the different developmental embryonic stages in mouse and describes in more details the gastrulation and the different pathways and factors associated.