Coordination of gene expression programs

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ABSTRACT

Most cellular processes depend on the activity and interactions of proteins. The proteome, i.e. the entire set of proteins in a specific condition, is shaped by regulation of transcription, mRNA-degradation, -processing, -storage, -translation and protein degradation. Cancer cells are known to highjack gene expression processes, including the translation machinery, for their growth and survival. This occurs as a result of converging oncogenic signaling pathways which impinge on translation factors to selectively modulate synthesis of cancer-related proteins.

Our understanding of mechanisms by which oncogenic pathways dynamically control their targets' translational activity is limited and could be extended by transcriptome-wide studies of changes in translation efficiency. In Paper I, we developed anota2seq which allows for statistical analysis of such data. Using a simulation approach, we showed that anota2seq constitutes an improvement compared to other methods for identification of genes under translational regulation.

The relative contribution of transcriptional and translational regulation to proteome modulation has been extensively debated. This raises the interest in studies integrating data on multiple levels of gene expression regulation. In Paper II, we study the role of estrogen receptor alpha (ERα), a transcription factor that is commonly targeted in hormone-dependent cancers, in coordinating transcriptional alterations with control at the level of translation. Upon ERα depletion in a prostate cancer model, we observed massive translational offsetting whereby the translational output remains unchanged despite changes in mRNA levels. To characterize mechanisms underlying translational offsetting, we extended the scope of the anota2seq method (Paper I) to also identify genes regulated by this underappreciated mode of gene expression regulation. Next, our detailed mechanistic study revealed that upon ERα depletion, mRNAs whose levels are reduced but translationally offset have less structured 5’UTRs and are devoid of miRNA target sites and thus cannot be influenced by such translational repressors. In contrast, transcripts which were upregulated but offset at the level of translation are enriched in codons requiring U34-modified tRNAs for their translation. We finally demonstrated that ERα impacts the levels of such modified tRNAs.

Cancer is a highly heterogeneous disease. In our studies of translational control, we are reaching the limits of reasonable inference when extending conclusions from experiments in cell lines into clinical settings. However, experimental methods to quantify translomes such as polysome-profiling, are not suitable for samples with low RNA input such as tissue samples from cancer patients. Paper III presents an optimization of the polysome-profiling method, compares it with the classical approach and validates that this new approach is suitable to study novel mechanisms regulating mRNA translation in large collections of tissue samples.
LIST OF SCIENTIFIC PAPERS

I. Oertlin C, Lorent J, Murie C, Furic L, Topisirovic I§, Larsson O§.

Generally applicable transcriptome-wide analysis of translation using anota2seq.
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Translational offsetting as a mode of estrogen receptor α-dependent regulation of gene expression.
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III. Liang S#, Bellato HM#, Lorent J§, Lupinacci FCS, Oertlin C, van Hoef V, Andrade VP, Roffé M, Masvidal L§, Hajj GNM§, Larsson O§.

Polysome-profiling in small tissue samples.
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# Equal contributions
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**IL-15 activates mTOR and primes stress-activated gene expression leading to prolonged antitumor capacity of NK cells.**


**Cancer as an ecomolecular disease and a neoplastic consortium.**


**eIF4E-binding proteins 1 and 2 limit macrophage anti-inflammatory responses through translational repression of IL-10 and cyclooxygenase-2.**


**The protozoan parasite toxoplasma gondii selectively reprograms the host cell translatome.**

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List of abbreviations

4E-BP  eIF4E binding protein.
a.u.  arbitrary units.
ALKBH8  alkB homolog 8, tRNA methyltransferase.
ATP  adenosine triphosphate.
AUC  area under the receiver operating characteristic curve.
BCL-XL  B-cell lymphoma extra large.
BRAF  B-Raf proto-oncogene, serine/threonine kinase.
CCNA2  cyclin-A2.
CCND3  cyclin D3.
CDK5  cyclin-dependent-like kinase 5.
CTU  cytosolic thiouridylase subunit.
DEK  DEK Proto-Oncogene.
DNA  deoxyribonucleic acid.
eEF  eukaryotic elongation factor.
eIF  eukaryotic initiation factor.
ELP3  elongator acetyltransferase complex subunit 3.
EMT  epithelial-to-mesenchymal transition.
ER  estrogen receptor.
eRF  eukaryotic peptide chain release factor.
ERK  extracellular signal-regulated kinase.
ERα  estrogen receptor alpha.
ERβ  estrogen receptor beta.
FC  fold change.
FDR  false discovery rate.
GCN2  general control non-derepressible 2.
GDP  guanosine diphosphate.
GLM  generalized linear model.
GTP meaning guanosine triphosphate.
HER2 denotes human epidermal growth factor receptor 2.
HRI stands for heme-regulated inhibitor.
HuR represents human antigen R.

m7G is 7-methylguanosine 5’-cap.
MAPK stands for mitogen-activated protein kinase.
MAPKKK signifies mitogen-activated protein kinase kinase kinase.
mcm5s2U denotes 5-methoxycarbonyl-methyl-2-thiouridine.
MDM2 refers to Mouse Double Minute 2.
MEF represents mouse embryonic fibroblast.
MEK signifies MAPK/ERK kinase (also known as MAPKK).
MET is mesenchymal epithelial transition factor.
miRNA means microRNA.
MNK stands for MAPK-interacting kinase.
mRNA means messenger RNA.
mRNP stands for messenger ribo-nucleoprotein particle.
mTOR denotes mammalian/mechanistic target of rapamycin.
mTORC stands for mTOR complex.
MYC means MYC proto-oncogene.

p53 refers to tumor protein p53.
PABP represents poly(A)-binding protein.
PDC4 means programmed cell death 4.
PDK1 signifies 3-phosphoinositide-dependent protein kinase 1.
PERK stands for PKR-like endoplasmic reticulum kinase.
PI3K refers to phosphatidylinositol-3-OH kinase.
PIC means pre-initiation complex.
PIP2 stands for phosphatidylinositol-4,5-bisphosphate.
PIP3 denotes phosphatidylinositol-3,4,5-trisphosphate.
PKR indicates double-stranded RNA-dependent protein kinase.
Pol II stands for RNA polymerase II.
Pol III represents RNA polymerase III.
pRB refers to retinoblastoma protein.
PTEN means phosphatase and tensin homologue.

RBP signifies RNA-binding protein.
RFM represents ribosome flow model.
RHEB stands for RAS homologue enriched in brain.
RNA ribonucleic acid.
RNase ribonuclease.
RNAseq RNA sequencing.
RPF ribosome-protected fragment.
rRNA ribosomal RNA.
RT-qPCR quantitative reverse transcription polymerase chain reaction.

S6K ribosomal S6 kinase.

TC ternary complex.
TE translational efficiency.
TOP terminal oligopyrimidine.
tRNA transfer RNA.
TSC tuberous sclerosis.
TSS transcription start site.

U34 Uridine 34.
uORF upstream open reading frame.
UTR untranslated region.

VEGF vascular endothelial growth factor.

WB Western blotting.
1 Prolegomenon

1.1 Translational control

1.1.1 Introduction about the central dogma of molecular biology

An organism’s genetic information is encoded in the deoxyribonucleic acid (DNA) located in its cells’ nuclei. Depending on its cell type and on which state a cell is in, specific locations (genes) along the DNA will be transcribed (expressed). This entails synthesizing a necessary amount of temporary copies of the DNA region called transcripts. About 62% of the human genome is transcribed and processed further (Djebali et al. 2012). Of these transcripts, 2-5% are messenger RNAs (mRNAs) which encode proteins (Carninci et al. 2005). The process of "converting" an mRNA template into a protein is called mRNA translation (Figure 1); this thesis will mainly be focused around studying regulation of gene expression at the level of mRNA translation.

![Figure 1: Main steps of gene expression regulation. This figure illustrates, for one gene, the processes of the central dogma of molecular biology. The protein level (p) is shaped by regulation of the rates of different steps including rate of transcription and processing (βm), of mRNA-decay (δm), -storage, -translation (βp), and protein degradation (δp). In the figure, m is the mRNA abundance. At a given time t, p(t) = (βm(t)−δm(t))βp(t)−δp(t). When measuring translational efficiency, some methods such as polysome-profiling (see section 1.1.4.1) estimate the proportion of efficiently translated mRNAs (mRNAs associated with many ribosomes, colored in blue) among all mRNAs. Other methods, such as ribosome-profiling (section 1.1.4.2), count the number of ribosome footprints (colored in yellow).](image)

A cell’s activity is characterized by quantity and interactions of proteins. The abundance of a protein is determined by the rate of transcription of its corresponding gene,
translation of the mRNA intermediate as well as rate of mRNA and protein degradation (Figure 1). These processes constitute the so-called central dogma of molecular biology (Crick 1970). At steady state, from one gene product to another, the range of transcription and translation rates is wide. Between 0.1 and 100 mRNAs are synthesized every hour while translation rates range from 10 to 10 000 proteins per mRNA per hour (Schwanhäusser et al. 2011; Li et al. 2014; Hausser et al. 2019). mRNA and protein median half-lives have been measured to be around 11 and 35.5 hours respectively and vary over a 10-fold range (Cambridge et al. 2011; Gregersen et al. 2014; Hausser et al. 2019). In terms of cellular energy consumption, mRNA translation is the most demanding step (~28% of the total adenosine triphosphate (ATP) production) in normal proliferating cells (Rolfe and Brown 1997). In uncontrolled proliferation contexts such as in cancer cells, control over the translational machinery therefore becomes essential (Robichaud et al. 2018).

1.1.2 Translation of an mRNA

Once transcribed, pre-mRNAs are processed (5’capping, 3’ polyadenylation, intron splicing) and exported from the nucleus to the cytoplasm of the cell. During translation, mRNA messages consisting of nucleotide sequences are converted into protein sequences consisting of amino acid sequences (Jacob and Monod 1961). On an mRNA, the part which codes for the protein is called coding sequence and is enclosed between the 5’ and the 3’ untranslated regions (UTRs). These regions typically include regulatory sequences which are used for translational control as will be described later.

Before protein synthesis starts, the ribosome which will translate the mRNA and allow for the assembly of the amino acid sequence, scans the 5’ UTR until recognition of the beginning of the coding sequence (Kozak 1989). This step is called translation initiation. Each triplet of nucleotides (codon) along the coding sequence will then be paired to the anticodon of a transfer RNA (tRNA) which will have previously been charged with the corresponding amino acid to be added to the polypeptide chain (Crick 1958; Chapeville et al. 1962). This occurs from the AUG start codon until the recognition of a stop codon (UAG, UAA or UGA) which terminates translation.

1.1.2.1 Translation initiation

More specifically, for translation to be initiated, several translation factors have to coordinate and the two subunits of the ribosome have to assemble at the start codon (Figure 2). On one hand a TC consisting of eIF2, initiator tRNA and GTP is formed. On the other hand, eIF4E binds the mRNA-cap and together with eIF4G and eIF4A form the eIF4F complex. This allows for recruitment of the 43S pre-initiation complex (40S small ribosome subunit, TC and additional initiation factors including eIF3) to the mRNA template. 5’ UTR scanning starts towards the
Figure 2: Translation initiation. Initiation is the rate-limiting phase of translational regulation. The formation of the 43S pre-initiation complex (PIC) assembles the 40S ribosomal subunit with eukaryotic initiation factor (eIF)1, eIF1A, eIF3, eIF5 and the ternary complex (TC) (eIF2 (containing α-, β- and γ-subunits), initiator methionyl tRNA and guanosine triphosphate (GTP)). eIF4E binds the 5’-cap of mRNAs and associates with eIF4G, a large scaffolding protein and eIF4A, a DEAD box ribonucleic acid (RNA) helicase. Assembly of such an eIF4F complex (eIF4E, eIF4G, eIF4A) facilitates the recruitment of ribosomes on the mRNA and forms a 48S PIC. Circularization of mRNAs via interaction between eIF4G and the poly(A)-binding protein (PABP) helps stability of the complex and increases translation efficiency.


start codon where the 60S big ribosomal subunit joins (Gingras et al. 1999; Jackson et al. 2010; Hinnebusch and Lorsch 2012; see also Figure 2).
1.1.2.2 Translation elongation  Once the 80S ribosome is assembled, translation elongation and the formation of the polypeptide chain can start. For each codon, eukaryotic elongation factor (eEF)1A:aminoacyl tRNA:GTP are presented to the ribosome to its A site (site for incoming amino-acid charged tRNA). Upon codon recognition, GTP is hydrolyzed, the tRNA is accommodated to the A-site and a peptide bond is formed between the amino acid of the A-site tRNA and the amino acid of the tRNA in the peptidyl (P) site. The polypeptide chain is transferred to the A-site before translocation is assured by eEF2 binding and GTP hydrolysis (Dever and Green 2012). The uncharged tRNA which was in the P-site is then moved to the exit E-site and the charged tRNA and polypeptide chain are moved to the P-site.

Before tRNAs can be efficiently used in translation, they undergo a tightly controlled biosynthesis involving multiple steps: RNA polymerase III (Pol III) transcription, removal of the 5’ leader and 3’ trailer sequences, addition of CCA on the 3’ end, splicing, modification and aminoacylation (Phizicky and Hopper 2010). tRNAs are ubiquitous and very abundant (15% of the total RNA) which facilitated their identification (Hoagland et al. 1958) even before understanding the concept of mRNA and the machinery of protein synthesis (Brenner et al. 1961). Notwithstanding major advances in the characterization of many RNAs notably thanks to breakthrough in sequencing technologies, a complete map of heavily modified RNAs such as tRNAs is still lacking (Juhling et al. 2009; Czerwoniec et al. 2009). Among the important tRNA modifications which are currently extensively studied, are the modifications which enable wobbling (non Watson-Crick base pairing between the first position of the tRNA anticodon (position 34) and the third position of the codon). Because of the wobbling rules, the same tRNA can decode several codons. In theory, a minimum of 30 tRNAs would be absolutely required. Yet, in human, tRNAs with 49 different anticodons are represented; these tRNAs are encoded by 513 genes (Chan and Lowe 2009). Not all tRNA genes are expressed at the same level and ribosomal speed at a specific codon will depend on abundance of the corresponding tRNA(s), the cellular demand for this/these tRNA species as well as the nature of the pairing (Watson-Crick or not). Along an mRNA, the ribosome transiently pauses when low abundance tRNAs are required which facilitates co-translational folding (Zhang and Ignatova 2011). The tRNA abundance correlates strongly with codon usage in prokaryotes and unicellular eukaryotes but not in higher eukaryotes (Novoa and Ribas de Pouplana 2012; Plotkin and Kudla 2011; Novoa et al. 2012).

New technological and analytical findings are allowing further understanding of the essential role of tRNAs and their modifications in translation (El Yacoubi et al. 2012; Sarin et al. 2018; Cozen et al. 2015). Concomitantly, more and more evidence have shown that dysfunctional tRNAs associates with the development of specific dis-
eases (Abbott et al. 2014; Torres et al. 2014; Kirchner and Ignatova 2015). In this
thesis, the effect of the modulation of specific tRNA-modifying enzymes will be stud-
ied in prostate and breast cancer cell lines (Paper II).

1.1.2.3 Translation termination When a stop codon enters the A site, the eukary-
optic peptide chain release factor (eRF)\textsubscript{1}:eRF\textsubscript{3}:GTP complex binds, GTP is hydrolyzed
and the polypeptide is released leading to ATP hydrolysis and subunit dissociation
(Zhouravleva et al. 1995). Recycling can occur by ribosome splitting, release of tRNA
and mRNA for reuse in synthesis of additional proteins (Hellen 2018).

1.1.3 Efficiency of translation of an mRNA

For a specific protein, the synthesis rate will depend on the amount of available
corresponding mRNA and on the rate of translation of each of these mRNA molecules. It has been known since the 1960s that protein synthesis occurs in polysomes meaning that mRNAs undergoing translation are typically associated with multiple ribo-
somes (Warner et al. 1963). Several models exist to describe ribosomal dynamics along an mRNA but the basic principle is illustrated in Figure 3.

The factors which can influence rate of initiation, elongation or termination include the number of available ribo-
somes, aminoacylated tRNAs, translation factors in the cell, the presence of specific structure or binding sites of the mRNA, codon usage along the transcript
and amino acid charge of the growing polypeptide chain (Tuller 2014; Figure 4). Regulation of global players (ribo-
somes, tRNAs, translation factors) can impact translational efficiency of all mRNAs in the cell. In contrast, reduction or

![Figure 3: Dynamics of translational efficiency regulation. The efficiency at which an mRNA is translated will depend on the initiation rate $\lambda$ (rate at which ribosomes start elongating), the transition rate at each codon ($\lambda_1, \lambda_2, ..., \lambda_n$ for an mRNA with $n$ codons) and the termination rate. In yeast, initiation rates have been estimated to be around 0.01-1.9 ribosomes per second while elongation rates would range from 1 to 20 codons per second (Riba et al. 2019). The average elongation rate in mouse embryonic stem cells was approximated around 5.2 codons per second (Sharma et al. 2019). More recent methods have included in this model the rate and position of ribosome drop-off (Bonnin et al. 2017).](image)
overexpression of some translation factors are known to affect specific subsets of mRNAs more than others (Koromilas et al. 1992; Rubio et al. 2014; Wolfe et al. 2014). Moreover, even at steady-state and in the absence of any specific stress or treatment, the range of translational efficiencies from one mRNA to another is widespread (Mathews et al. 2007; Hauser et al. 2019). Finally, translation of mRNAs holding particular RNA or protein-binding sites may be affected according to the level, availability or likelihood of the interaction with corresponding partners (Gebauer et al. 2012; Hentze et al. 2018). As such, inherent characteristics of translational efficiencies and differential sensitivities are encoded in the mRNA’s sequence and structures (cis-regulatory elements) while modulation of trans-acting factors (e.g. microRNAs (miRNAs), RNA-binding proteins (RBPs)) will mediate translational changes on the mRNA subsets that they target (Figure 4; Hinnebusch et al. 2016; Leppek et al. 2018). The interplay between RNA elements and trans-acting factors will be further described in section 1.2 about Coordination of gene expression programs.

**Figure 4: Cis- and trans-regulation of mRNA translation.** Primary and higher-order structures along the mRNA influence the efficiency at which it is translated. For instance, complex structures in the 5’UTR, presence of upstream open reading frames (uORFs) or miRNA binding sites are mostly associated with down-regulation of translation efficiency of the main open reading frame; specific RNA modification such as m6A may help ribosome scanning while RBP can act either as enhancers or repressors of translation. Finally, presence of suboptimal codons may reduce the speed of elongation especially in relation to availability of their corresponding tRNAs.

Initiation is the rate limiting step of translation in most conditions (Sonenberg and Hinnebusch 2009). This implies that small changes in elongation or termination rates are likely to have limited effect on the rate of protein output whereas modulation at the initiation step typically have more impact. Accordingly, regulatory mechanisms will affect assembly of initiation complexes or scanning along the 5’UTR more often than other steps of the translation process.
1.1.4 Experimental methods to measure differential translational efficiency

In cases where translation initiation is rate-limiting (which is the most common scenario), the number of ribosomes associated with mRNAs is a good proxy for efficiency of translation. mRNAs associated with few ribosomes are then considered inefficiently translated while heavy polysomes (mRNAs associated with many ribosomes) would have higher protein output per mRNA molecule and time unit.

1.1.4.1 Polysome-profiling  The polysome-profiling method is based on the principle that heavier polysomes will contain more efficiently translated mRNAs. A gradient of sucrose is prepared with density increasing linearly from 5% to 50%. Treatment with an inhibitor of translation elongation such as cycloheximide immobilizes ribosomes on mRNAs and cytoplasmic lysates are loaded on the gradient. This allows, after ultracentrifugation, for differential sedimentation of mRNAs depending on the number of ribosomes they are associated with (Gandin et al. 2014; Figure 5A).

Polysome-profiling can be used to assess the overall quantity of translating mRNAs under different conditions. A decrease in global translation, i.e. in translation of most mRNAs, would be identified by a higher 80S peak and lower polysome peaks as exemplified in Figure 5B upon inhibition of mTOR translation by torin1 (pink tracing) as compared to insulin-stimulated MCF-7 cells (orange tracing). Quantification of specific transcripts along the gradient fractions allows for assessment of transcript-level regulation of mRNA translation. This potentially permits identification of subsets of transcripts sharing common features leading to their co-regulation at the level of translation. For a specific mRNA, in conditions where it is efficiently translated, its ribosome occupancy mode (most common value) will typically be higher than 3 while in inefficient translation conditions, it would usually be associated with fewer than 3 ribosomes. This "3-ribosome-cutoff" (Figure 5A, orange dotted line) for differentiation between efficient and inefficient translation applies to most mRNAs (Larsson et al. 2013; Gandin et al. 2016). Therefore, for transcriptome-wide analysis of differential translation, a cutoff is set at 3 ribosomes and fractions corresponding to mRNAs with more ribosomes than this cutoff are pooled (later referred to as polysome-associated mRNA). This efficiently translated mRNA pool is then extracted and quantified using DNA microarrays or RNAseq. At the single gene level, a change in abundance within the polysome-associated mRNA pool can either be caused by a shift in the number of bound ribosomes for that mRNA (i.e. change in translational efficiency, Figure 5B) or a change in the steady state mRNA level (i.e. change in transcription and/or mRNA stability, Figure 5C). To distinguish between changes in translation and changes in steady state mRNA levels, cytoplasmic mRNA is collected and quantified in parallel. In the absence of regulation via translation, changes in cytoplasmic mRNA levels should be reflected in corresponding changes in polysome-associated mRNA.
Figure 5: Examples of regulation by mRNA translation or abundance measured by polysome-profiling. After starvation, MCF7 cells were stimulated with insulin or insulin in the presence of torin1, an inhibitor of mammalian/mechanistic target of rapamycin (mTOR) translation (see also section 1.2.2.2.1). Polysome-profiling was performed on insulin and insulin+torin1 treated cells. (A) The profile illustrates separation of ribosomal subunits 40S and 60S, the 80S monosome peak as well as peaks corresponding to mRNAs associated with 2, 3, 4, ribosomes etc. For gene level quantification by DNA-microarrays or RNA sequencing (RNAseq), fractions corresponding to mRNAs associated to more than 3 ribosomes are pooled and quantified in parallel of the cytoplasmic mRNA input. (B) A known torin1-sensitive mRNA (cyclin D3 (CCND3)) was quantified in each fraction by quantitative reverse transcription polymerase chain reaction (RT-qPCR) and illustrates a shift in translational efficiency towards lighter fractions upon inhibition of mTOR translation (blue and green lines). (C) In contrast, suppression of the cytoplasmic level of an mRNA i.e. by down-regulation of its synthesis or stability, leads to a vertical shift without reduction of the average number of associated ribosomes. However, polysome-associated mRNA is reduced between the green and blue tracings both in (B) and (C).

Modified from Gandin, V et al. (2016). "nanoCAGE reveals 5’ UTR features that define specific modes of translation of functionally related MTOR-sensitive mRNAs." In: Genome research 26(5), pp. 636-648. doi: https://doi.org/10.1101/gr.197566.115. ©2016 Gandin et al.; Published by Cold Spring Harbor Laboratory Press. This article, published in Genome Research, is available under a Creative Commons License (Attribution 4.0 International), as described at http://creativecommons.org/licenses/by/4.0/

This standard polysome-profiling protocol is often adapted to specific cases: when studying specifically short mRNAs, a cutoff at 2 ribosomes can be deemed more ap-
propriate (Aspden et al. 2014); when assessing usage of alternative isoforms, more fractions and deeper sequencing may be required (Floor and Doudna 2016). **Paper III** of this thesis will present an optimization of the polysome-profiling technique allowing to expand its applicability to samples with low RNA-amounts such as most tissue samples.

In comparison with other methods measuring differential translation (see 1.1.4.2 Ribosome-profiling below), the polysome-profiling technique has the advantage to directly assess, on each mRNA molecule, whether it was associated with low or high amounts of ribosomes. However, it does not provide information about ribosome positioning along the mRNA.

### 1.1.4.2 Ribosome-profiling

Ribosome-profiling was developed more recently than polysome-profiling. It entails blocking translation elongation before degrading RNA not protected by ribosomes using ribonuclease (RNase) (Ingolia et al. 2009). Sequencing libraries are built based on the remaining ribosome-protected fragments (RPFs) providing a quantitative profile of ribosome occupancy after alignment to the transcriptome. For reasons already mentioned in the previous section, cytoplasmic mRNA input (which can be randomly fragmented in order to obtain RNA fragments of similar sizes as RPFs) is collected and quantified in parallel. On one hand, because this method is based on RNase digested samples, information about the number of ribosomes associated with each mRNA is lost. On the other hand, due to its ability to locate ribosomes, it has been providing valuable insights into regulatory mechanisms that control elongation speed, ribosome pausing and translation of alternative open reading frames (Ingolia et al. 2018). Therefore, polysome- and ribosome-profiling methods are complementary.

Ribosome-profiling has however been suffering from experimental artefacts. For instance, it has been shown that cycloheximide, which can be used to inhibit translation elongation, may modify ribosome distribution along the mRNA near specific sequences leading to spurious footprints (Brar and Weissman 2015). Furthermore, the RNase treatment which is used in order to digest RNA between translating ribosomes, also tends to digest ribosomal RNA (rRNA) from ribosomes leading to partial loss of the footprints. However, tools to better assess the quality of ribosome-profiling data, alternative ribonucleases (Gerashchenko and Gladyshev 2017) and in silico methods to flag spurious RPFs have been made available (O’Connor et al. 2016; Brar and Weissman 2015; Kiniry et al. 2019).

One limitation which is shared by polysome- and ribosome-profiling arises in the context of specific mRNAs and/or conditions where translation elongation would be
the rate-limiting step, instead of initiation. Indeed, even if initiation speed would typically control protein synthesis rate, assuming that it is the case in any condition and for any gene may be oversimplifying. In occasions where elongation would be rate-limiting, an increased number of ribosomes along an mRNA would indicate a reduced rate of elongation and, provided that translation initiation remained unchanged, a reduced efficiency of translation. As such, when it is unknown which step is rate-limiting, a change in polysome-associated mRNA or RPF, even in the absence of corresponding change in cytoplasmic mRNA, cannot be attributed with certainty to a change in the same direction of the translational efficiency (Mathews et al. 2007). In order to overcome this limitation, other methods to measure translational output, for instance by labeling of newly synthesized polypeptides can be used. Puromycin incorporation-based methods are typically used in this context (Iwasaki and Ingolia 2017) and can be combined with protein-specific antibodies in order to assess translation output of a given protein (Söderberg et al. 2006; Tom Dieck et al. 2015). In order to assess in greater details the translational metrics, such as initiation and elongation rates, one would need to calculate them from the polysome size and ribosome transit time (defined as the time for the ribosome to traverse the mRNA) which can be measured based on radioactivity kinetics from nascent polypeptides to released polypeptides (Fan and Penman 1970; Gehrke et al. 1981). Even if they do not allow the level of precision offered by these methods, polysome- and ribosome-profiling have the advantage to provide a transcriptome-wide perspective of translational control.

1.1.5 Analytical methods for transcriptome-wide analysis of differential translation

Development and improvements of experimental methods highlighted the need for appropriate statistical methods to detect genes showing differential translational efficiencies within transcriptome-wide pools. This section will focus on reviewing analytical methods testing for regulation at the level of translation between 2 or more conditions. Such results can be obtained from both polysome-profiling data (provided that both efficiently translated mRNA and cytoplasmic mRNA were quantified) and ribosome-profiling data (providing quantification of RPFs and cytoplasmic mRNA). The main challenge of such methods is to accurately "correct" polysome-associated mRNA or RPF changes for modulations in steady-state mRNA. As mentioned previously, for one gene, a difference between conditions in its polysome-associated mRNA expression or its RPF expression, can mirror a similar difference of expression observed in its cytoplasmic mRNA expressions which were quantified in parallel; this would primarily be the case of genes under regulation by transcription or mRNA degradation. Translational regulation would then be defined as changes in polysome-associated mRNA or RPF which are independent from fluctuations in cytoplasmic mRNA levels.
It is important to note that even though similar computational methods are typically used downstream of polysome- or ribosome-profiling, data coming from these methods are inherently different. Indeed, while in polysome-profiling data, gene-level differences in translational efficiencies will be measured as horizontal shifts (exemplified in Figure 5B), they will be measured as differences in total numbers of ribosomes (coming from any mRNA synthesized from this gene) in ribosome-profiling experiments (see also Figure 1). The relationship between the log number of ribosomes associated with an mRNA and the sedimentation distance along a sucrose gradient is very robust (as observed in Gandin et al. (2016) and Paper III). However, on a gene-level, differences in RPFs (which "counts" ribosomes) can be very distinct from differences in polysome-associated mRNA (which quantifies copies of efficiently-translated mRNAs). As such, even when studying similar biological mechanisms, studies using polysome- and ribosome-profiling data have proven to sometimes lead to conflicting conclusions (Larsson et al. 2012; Hsieh et al. 2012; Thoreen et al. 2012; Gandin et al. 2016; Masvidal et al. 2017) and may require deviating interpretations of their results.

Most of the methods which will be described below have been designed for ribosome profiling data while Paper I involves a method which was developed for polysome-profiling. However, in theory, all these tools could be applied to any data source where the intention is to identify changes in an RNA subset that is independent of a background (e.g. total or cytoplasmic RNA). For this reason, in the next paragraphs, "translated mRNA" will be used as a generic term referring to polysome-associated mRNA or RPFs.

When most mRNA quantification was performed on DNA-microarrays, few methods for analysis of changes in translational efficiencies were available. They have been described and reviewed previously (Larsson et al. 2010; Larsson et al. 2013). Typically, statistical methods which were applicable to DNA-microarrays data cannot be directly used on RNAseq data because of inherent differences in the properties of these two data types (Robinson and Smyth 2008). DNA-microarrays quantify gene expression by measuring intensities or intensity summaries whereas RNAseq quantifies it by measuring numbers of short reads. Empirically, this results in data from DNA-microarrays which, after log-transformation, fulfils normality requirements of most statistical frameworks and data from RNAseq remaining non-normal even after classical transformations and rather following negative binomial-like distributions (the variance of read counts between replicates is generally higher than their mean).

Babel was one of the first method specifically developed for RNAseq based measurements of translation (Olshen et al. 2013) and was designed for data from ribosome-
profiling. It took into account the count nature of the data and used an errors-in-
variables regression model between RPF and cytoplasmic mRNA in order to account
for the fact that gene expression can only be measured with some level of uncertainty.
This is particularly the case of non-replicated experiments which are not uncommon
and for which Babel can be used. However, in later benchmark studies, this method
showed to perform relatively poorly in terms of control of type I error (Xiao et al.
2016).

eXgeR (Robinson et al. 2010) and DESeq2 (Love et al. 2014) are two methods
which were initially developed for identification of differential gene expression from
RNAseq experiments where only cytoplasmic mRNA was considered. These two meth-
ods differ in the specific methods for dispersion estimation, information sharing across
genes and normalization but they share similar principles. They are both parametric
methods assuming a negative binomial distribution for the read counts and use gen-
eralized linear models (GLMs) for differential expression testing. The use of GLMs
makes these methods very flexible and allow for analyzes of complex study designs.
As such, they can easily be adapted towards analysis of differential translation. Reads
counts $K_{gj}$ are modelled as negative binomial distributions with mean $\mu_{gj}$ and dis-

tersion $\sigma_g$ for gene $g$ and sample $j$. The mean is considered as the product of $q_{gj}$
(parameter proportional to the expected true concentration of fragments from gene $g$
for sample $j$) by a size factor accounting for differences in sequencing depth between
samples. For classical differential expression analysis between, for instance, a treated
and a control condition the GLM would be as follows:

$$
\log(q_{gj}) = \beta_g^{Trt} X_{j}^{Trt} + \epsilon_{gj}
$$

with coefficients $\beta_g^{Trt}$ representing the log$_2$ fold changes for gene $g$ for each col-

umn of the model matrix $X$ and $\epsilon_{gj}$ the error term. For analysis of differential transla-
tion, two additional parameters can be incorporated into the model: a parameter for
the RNA type (i.e. cytoplasmic or translated mRNA), and an interaction parameter
between treatment and RNA type.

$$
\log(q_{gj}) = \beta_g^{Trt} X_{j}^{Trt} + \beta_g^{RNAtype} X_{j}^{RNAtype} + \beta_g^{RNAtype:Trt} X_{j}^{interaction} + \epsilon_{gj}
$$

The interaction term can be interpreted as the differential effect of treatment in
translated mRNA compared to cytoplasmic mRNA i.e. the translational control effect.
This "GLM with interaction term" is the principle used by several recent methods in-
cluding Riborex (Li et al. 2017b), Ribodiff (Zhong et al. 2017) and deltaTE (Chothani
et al. 2019). Riborex is directly available either as "DESeq2 with interaction term"
or "edgeR with interaction term" whereas Ribodiff uses slightly different estimation
methods and allows for different dispersion estimations from cytoplasmic and translated mRNA data (Zhong et al. 2017). Although estimating different dispersions for data coming from different experimental procedures seems relevant, Ribodiff seemed to underperform in simulation studies compared to Riborex (Li et al. 2017b). In the implementation of Riborex as an R package, its use is restricted to simple designs whereas the deltaTE implementation of the same methods maintains their availability to more advanced applications including the possibility to correct for batch effects in the model (Chothani et al. 2019). Xtail (Xiao et al. 2016) uses DESeq2 for estimation of mean and dispersion while allowing for analysis of experiments with only one replicate per condition (which only Babel allowed so far). Xtail outperforms other methods in terms of specificity and sensitivity when the sample size is very low (1 or 2 replicates) but further analyzes included in Paper I show that this is at the cost of detection of high amounts of non-differentially translated genes when tested under a NULL model (no true differences in gene expression) especially when the sample size is low. Xtail was also shown to be strongly impacted when a batch effect was added to the data even though systematic differences between replicates are commonly observed in translatome data (Chothani et al. 2019).

The importance of dysregulated translation for cancer progression has been increasingly recognized over the last 30 years (Lazaris-Karatzas et al. 1990; De Benedetti and Graff 2004) such that the translation initiation apparatus is presently a target in oncology (Chu and Pelletier 2018). It has also been associated with other diseases such as neurodegenerative diseases (Moreno et al. 2012) or metabolic disorders (Shi et al. 2003). Consequently, efforts have been put to understand mechanisms dysregulating mRNA translation and specifically to improve experimental methods for quantification of the translatome as well as computational methods to detect differential translational control. Two of the constituent papers of this thesis (Paper I and Paper III) are extensions of classical methods: Paper III presents an optimization of polysome-profiling allowing it to become scalable to large sets of small samples (low RNA input as in biobanked tissue samples) and Paper I provides an improvement of the computational method anota (Larsson et al. 2010) which was only available for quantification by DNA-microarrays data. Anota2seq (Paper I) extends its use to RNAseq-quantified polysome- or ribosome-profiling data. Furthermore, we demonstrated that anota2seq shows appropriate control of type I error, can include batch-effect correction and allows identification of genes whose mRNA levels are buffered at the level of translation (this gene expression mode of regulation will be presented in the next sections).
1.2 Coordination of gene expression programs

In general, the objective of omics studies (e.g. transcriptomics which measure the full set of transcribed RNA molecules, proteomics which measure the full set of proteins, metabolomics measuring any metabolites, translatomics measuring translated mRNAs, etc.) is to understand which cellular pathway or mechanism is influencing a specific phenotype. For instance, in an attempt to study which regulatory events or mechanisms lead to the development of resistance to BRAF inhibitors in melanoma (a classical treatment in this disease which is known to be efficient for a certain time until the patient stop responding and the tumor relapses), one would extract biomolecules of interest (RNA, proteins, metabolites, translated mRNAs, etc.) from a cell line which is sensitive to the treatment and one which has developed resistance. From inferred differences in abundance of specific species of these biomolecules, one would generate hypotheses regarding essential cellular events driving the resistance.

Because proteins are arguably the most important players in many biological processes, a majority of such research projects would favor proteomics over other omics approaches. Transcriptomics may be considered a less expensive approach which benefits from well-established technologies and which, in some specific contexts, would be deemed to provide an appropriate proxy for protein levels. The question of whether RNA and protein levels strongly correlate, of whether RNA levels is a good surrogate for protein levels is controversial. In other words; among transcription, mRNA translation or decay, which step has the most important contribution to shaping the proteome has been extensively debated (Schwanhäusser et al. 2011; Vogel and Marcotte 2012; Jovanovic et al. 2015; Li and Biggin 2015; Li et al. 2017a; Liu et al. 2016). A consensus has nonetheless been reached on one aspect of this discussion: the dynamic contribution of each process between conditions is context- and biological system-dependent with mRNA translation having a high contribution upon severe stress (Liu and Aebersold 2016) such as endoplasmic reticulum stress (Baird et al. 2014; Guan et al. 2017; Cheng et al. 2016). Thus, changes in mRNA levels cannot be generally considered as a good proxy for changes in protein levels.

Notwithstanding major improvements in coverage and precision of recently developed proteomics technologies (Branca et al. 2014; Orre et al. 2019), observing changes in protein levels upon a specific perturbation only informs on the impact on the output product of the gene expression pathway. In systems where it could be assumed that the intervention directly affects this output (i.e. the protein abundance), this would provide sufficient evidence for a broad understanding of the mechanisms affecting the phenotype. However, cellular pathways, including those active in cancer, are often very complex containing feedback loops and alternative branches. This is a
reason for the increased interest in multiple omics studies (i.e. where several levels of gene expression are measured on the same samples). In the next section, control at the translational level will be considered in the general context of gene expression regulation.

1.2.1 Regulation of gene expression at multiple levels

In eukaryotes, because mRNA translation occurs in the cytoplasm and transcription in the nucleus, these processes are often thought to be controlled independently. However, this view is probably over-simplistic and mechanistic examples of crosstalk between separate gene expression processes are common. Transcription and splicing are predominantly coupled (Beyer and Osheim 1988) and it has been hypothesized that co-transcriptional splicing could impact the integrity of transcription (Komili and Sil- ver 2008). The development of methods to efficiently sequence long nascent RNA molecules will soon unravel dynamics and order of intron removal (Drexler et al. 2019). In the cytoplasm, advances in other sequencing-based technologies allowed 5’ to 3’ mRNA co-translational decay to be characterized by observing 3-nucleotide periodicity in mRNA degradation intermediates. These were interpreted as products of exonucleases following the last translating ribosome (Pelechano et al. 2015). Co-translational decay is general and conserved which indicates that interplay between mRNA translation and decay is more complex than the classical view stating that translating mRNAs are protected from decay (Roy and Jacobson 2013).

Concerning crosstalks between different cellular localizations, Haimovich et al. (2013) have introduced the concept of synthegradosome where mRNA synthesis is linked to degradation by decay factors that were shown to shuttle from the cytoplasm to the nucleus to associate with transcription start sites and regulate transcription initiation. Furthermore, few examples exist of coordination between transcription and translation. For instance, Rpb4p and Rpb7p which are subunits of the yeast RNA polymerase II (Pol II) and also mediate mRNA decay (Choder 2004; Lotan et al. 2005; Lotan et al. 2007) have later been associated with regulation of translation initiation by interacting with eIF3 (Harel-Sharvit et al. 2010).

Interestingly, when conducting ribosome- and polysome-profiling experiments, cytoplasmic mRNA samples always need to be quantified in parallel of translated mRNA. Thus, data from these measurements provide more information than only translational regulation. Differences in mRNA abundance and potentially coordination between different layers of gene expression regulation can be also be analyzed. This could unravel whether the mRNA synthegradosome (transcription and mRNA decay) act in concert with mRNA translation or if translation seems to compensate or counteract modula-
1.2.1.1 When mRNA syntheagradosome and translation act in concert  Gingold et al. (2014) gave an interesting example where regulation at the level of transcription/mRNA decay and mRNA translation may be coordinated to both enhance or both repress cellular functions. Namely, in proliferating or cancer cells vs. differentiated or normal cells, opposing tRNA signatures are expressed (Figure 6). Notably, tRNAs translating different codons for the same amino acid showed opposite trends. Furthermore, when analyzing the codon usage of genes expressed in proliferation and differentiation, this study demonstrated that proliferation genes are enriched in codons (Figure 6, red colored codons) matching the "proliferation-tRNA-signature" (Figure 6, red colored anticodons) and inversely differentiation mRNAs require the "differentiation-signature" tRNAs for their translation (Figure 6, blue). They conclude that if in translation, the codon usage of an mRNA is the demand and the available tRNAs correspond to the supply; proliferation and differentiation are cellular states in which supply and demand are well matched.

Figure 6: Coordination between tRNA supply and demand in proliferation and differentiation. Genes belonging to proliferation-, respectively differentiation-, related ontology genesets are enriched for a specific set of codons (red, respectively blue). Red codons preferably have, at their third nucleotide position, A or U while blue codons preferably have C or G at this position. Interestingly, method development in tRNA quantification revealed differences in tRNA pools between cell models of proliferation vs. differentiation as well as between cancer and normal tissue samples. Gingold et al. (2014) observed in each cellular state, a match between tRNA supply and codon usage.


This interpretation has however been challenged. Rudolph et al. (2016) has argued that although codon-driven differences in translational efficiency between conditions have been extensively observed in prokaryotes and single-cell eukaryotes (Man
and Pilpel 2007; Drummond and Wilke 2008), the level of evidence for similar mechanisms in mammals remains low. When studying several steps of tissue development in mice, they did not detect overall differences in tRNA supply nor demand (Schmitt et al. 2014). Deciphering such contradicting results is complex: established methods to measure modified and charged tRNAs are lacking as well as trustworthy quantification of global differences in tRNA levels; assessing whether a specific tRNA pool can equally well translate different transcriptome-weighted codon usages is difficult; translational efficiency correlates to some extent with codon usage and with transcriptomic GC content but the causality between these events is less trivial to address. However, the conclusion which has a high level of confidence is that mammalian systems do not depend on codon bias as a translational regulatory mechanism as much as prokaryotes (Rudolph et al. 2016).

1.2.1.2 When mRNA translation compensates modulations at the mRNA level

Situations where a gene is strongly up-regulated at the mRNA level while the corresponding protein is strongly down-regulated are uncommon and sometimes attributed to data normalization issues (Albert et al. 2014). However translational buffering where protein levels remain stable between conditions despite alterations in the RNA levels has been described in several organisms (Lalanne et al. 2018; McManus et al. 2014; Artieri and Fraser 2014; Cenik et al. 2015). For instance, Lalanne et al. (2018) compare mRNA levels and protein synthesis rates between divergent bacteria species which have evolved independently for several billion years. They take the example of 4 genes (rpsP, rplS encoding two ribosomal proteins, rimM an rRNA-maturation factor and trmD a tRNA modification enzyme) which are expressed as a polycistronic mRNA in E. coli whereas in B. subtilis, they have a wide range of expression levels. Three of these genes show big differences in expression between the two bacterial species (from 5 to more than 30 fold). However, the function of these homologous proteins are similar and the relative levels of RPFs between the species are similar (less than 2-fold differences for all 4 mRNAs). Thus, transcriptional differences between divergent bacterial species are translationally compensated. Mechanistically, this is achieved by translationally suppressive mRNA secondary structures which are present in E.coli and absent in B. subtilis. Consequently, while regulation of these genes occurs at the transcriptional level in B. subtilis, it occurs by mRNA translation in E. coli.

Other mechanisms mediating translational buffering have been characterized in transcriptome-wide studies across yeast species (McManus et al. 2014; Artieri and Fraser 2014) and between human patients (Cenik et al. 2015).

Notably, several definitions of "translational buffering" co-exist in the literature. First of all, in some publications, steady-state co-expression of proteins encoded by
non co-expressed mRNAs (or vice versa) is also termed translational buffering or translational compensation (Kustatscher et al. 2017; Dassi et al. 2015) but this is not the definition used herein. Indeed, this thesis will focus on changes in gene expression observed under dynamic settings (i.e. between conditions) and therefore needs to be distinguished from when mRNA and protein levels are compared under a single condition. The different modes of regulation of gene expression which can be analyzed from transcriptome-wide analyzes of differential translation (i.e. from polysome- or ribosome-profiling data for instance) are described in Figure 7 between 2 hypothetical conditions (treatment T and control C). As explained earlier, a change in translated mRNA between conditions can be due to a change in mRNA abundance (Figure 7A top left; Figure 7B bottom) or a change in translational efficiency (i.e. a difference in ribosome occupancy resulting in a shift along the polysome gradient; Figure 7A top right; Figure 7B top; see also Figure 5). Finally, translational buffering is defined as a change in cytoplasmic mRNA levels which is not reflected in a change in translated mRNA (Figure 7A bottom left; Figure 7B middle).

It remains to be fully characterized whether specific mechanisms or specific biological processes are associated with situations where mRNA level modulations are buffered or not at the level of translation. An example of such a mechanism will be discussed in Paper II. However, this is largely unexplored area of research and to further the knowledge of mechanisms coordinating gene expression, availability of analytical methods distinguishing between changes in translational efficiencies leading to altered proteins levels or buffering is essential. This is an important feature of the anota2seq methodology which will be presented in Paper I of this thesis.

### 1.2.1.3 Role of estrogen receptor alpha in breast and prostate cancer

Most studies analyzing transcriptome-wide changes in translational efficiencies are focused on the role of translation factors or on mechanisms known to impact mRNA translation such as stress responses. In contrast, in Paper II, we studied the role of a transcription factor, namely estrogen receptor alpha (ERα), in coordinating transcriptional and translational output. Indeed, in addition to its role as a transcription factor, ERα has been shown to potentially have non-nuclear functions and to affect the mTOR and mitogen-activated protein kinase (MAPK) pathways which impinge on mRNA translation (see section 1.2.2.2). Further details about ERα-dependent gene expression regulation will be provided in Paper II while this section will expose some background information regarding ERα’s position as therapy target in breast cancer and co-driver of tumorigenesis in prostate cancer.

ERα is mainly studied in breast cancer which is the most common cancer type in women accounting for almost one in 4 cancers. In March 2019, the reported world-
wide age standardized incidence was 46.3 per 100 000 with a mortality rate of 13/100 000 (Cancer fact sheets - Breast). In this disease, ERα is both a prognostic marker, i.e. a characteristic of the disease which is associated with patients’ survival independently of treatment, and a predictive factor for endocrine therapies, i.e. it characterizes how responsive to this treatment tumors are. Namely, patients whose breast tumor expresses the receptor (estrogen receptor (ER)+), have a better prognosis than ER- patients, at least in the first 5 years after their cancer diagnosis (Knight et al. 1977). Furthermore, ER+ tumors have been shown to benefit from endocrine therapy (Tamoxifen) whereas little to no such effect was seen in ER- patients (EBCTCG 1998).

Hormonal treatment is also the most common strategy against prostate cancer but in this case targeting the androgen signaling or receptor. Prostate cancer is the second most common cancer in men worldwide. Its age standardized incidence and mortality rates are 29.3 and 7.6 per 100 000 respectively (Cancer fact sheets - Prostate). Prostate cancer is thought to be an androgen-dependent disease making androgen-deprivation therapy its most classical treatment strategy (Parker et al. 2015). However,
evidence that androgens alone could induce sustained proliferation and carcinogenesis in prostate cells is remarkably lacking (Morgentaler and Traish 2009). In combination with other carcinogens or upon chronic treatment of high doses of testosterone (androgens), prostate cancer could be induced (Pour and Stepan 1987; Bosland 2013). Strikingly, estrogens, in combination with testosterone result in prostate cancer in 90 to 100% of cases in animal models (Bosland et al. 1995); they can also stimulate cancer progression and metastasis (Ricke et al. 2006). Estradiol (estrogens) are synthesized from testosterone by the aromatase enzyme which has been shown to be expressed in the prostate (Ellem et al. 2004). A landmark study showed that aromatase knockout mice which were treated with androgens and estrogens developed pre-malignant prostate disease whereas androgen alone did not result in such development. Consistent with a leading role for ER signaling, the combination of androgens and estrogens did not lead to the disease in ERα knockout mice (Ricke et al. 2008). Understanding of the role of estrogens in prostate cancer is further complicated by the fact that estrogen receptor beta (ERβ) may, in contrast to ERα, have beneficial effects on cancer development, although this is under debate (Nelson et al. 2014). In the study mentioned above (Ricke et al. 2008), knockdown of ERβ led to development of pre-malignant prostate cancer upon treatment with both hormones.

In humans, even though aromatase inhibitors have proven their efficacy in post-menopausal breast cancer (i.e. in women whose ovaries no longer produce significant amounts of estrogens but in whom estrogens are still produced from androgens; Dowsett et al. 2010, Cardoso et al. 2019), clinical trials of similar drugs concluded that they should not be advised for prostate cancer treatment (Smith et al. 2002; Santen et al. 2001).

Although androgen-deprivation therapies are efficient against most prostate tumors, resistance to this treatment is not uncommon and more than 15% of men initially diagnosed with prostate cancer die from advanced incurable disease. A recent study showed that ERα is specifically expressed in the epithelium of advanced prostate tumors (Takizawa et al. 2015). The role of this receptor is still poorly understood in this disease but advances in this field could lead to promising therapeutic strategies. As mentioned previously, ERα is an extensively studied transcription factor in breast cancer; yet, its transcriptional targets can be tissue-specific which extends the need for transcriptome-wide studies of ERα-dependent gene expression in prostate tissue. Furthermore, Simoncini et al. (2000) showed that ERα can associate with phosphatidylinositol-3-OH kinase (PI3K) which is upstream of important regulators of mRNA translation. Thus, both transcriptional and translational outputs of ERα will be investigated in Paper II.
1.2.2 Coordinated regulation of sets of mRNAs

A well-accepted perspective is to consider that mRNAs are not linear nucleic acids but rather, in addition to harboring secondary structure and carrying multiple modifications, exist most often in complexes with proteins and other RNAs, so called messenger ribo-nucleoprotein particles (mRNPs) (Gebauer et al. 2012). If each mRNA should be viewed in regard to the activity of each of its interacting partners, each trans-acting factors, e.g. RBP or miRNA, typically binds to multiple mRNA targets thus coordinating entire RNA operons (Keene 2007). These many-to-many relationships define complex interactomes. Furthermore, multiple post-transcriptional regulatory features typically co-occur in mRNAs with the activity of one RNA feature also potentially depending on the proximity of other features (Cottrell et al. 2018). Recent advances in technology have facilitated better characterization of RNA features and trans-regulatory elements required for selective post-transcriptional regulation (Truitt and Ruggero 2016). Among others, experimental challenges have been resolved to better assess which mRNAs are selectively translated (Brar and Weissman 2015), binding and activity of RBPs (Van Nostrand et al. 2016), regulatory patterns of miRNAs (Katchy and Williams 2016) and tRNAs (Sarin et al. 2018), usage of alternative transcription start sites (Poulain et al. 2017; Gandin et al. 2016; Masvidal et al. 2017) but additional studies of the translatome under multiple conditions will be essential before a complete understanding of how post-transcriptional regulation is manifested can be reached.

As mentioned in section 1.1.5, computational methods for transcriptome-wide analysis of differential translation will distinguish genes regulated at the translational level from changes in mRNA abundance. Previously annotated sets of mRNAs co-regulated at the translational level constitutes essential tools helping to extract relevant biological findings from lists of genes regulated at different steps of the gene expression pathway in new studies. Let us consider the following example: eIF4A is a DEAD-box RNA helicase which helps unwinding secondary structures along 5'UTRs (Parsyan et al. 2011). Consequently, upon treatment with an eIF4A-inhibitor, the set of mRNAs harboring long and highly structured 5'UTRs will show reduced translational efficiency (Wolfe et al. 2014; Rubio et al. 2014; Gandin et al. 2016). It has also been specified recently that the position of the structures along the UTR impacts the sensitivity to eIF4A (Waldron et al. 2019). Therefore, if, in a new study, it can be proven that mRNAs having complex structures close to the start codon are specifically downregulated at the translational level, a reasonable hypothesis could be that eIF4A suppression may drive the studied phenotype. Similarly, as a second example, if in another study, enrichment for a specific RNA-element is observed among translationally modulated genes, one might deduce that a factor binding to this element may be
regulated between the studied conditions. Several examples of known mechanisms by which subsets of mRNAs may be translationally co-regulated will be described in the next paragraphs.

1.2.2.1 Regulatory RNA elements and trans-acting factors

1.2.2.1.1 microRNA and RNA-binding proteins Regulatory mechanisms of translation commonly involve interactions between specific structure or sequences along the UTRs and RNA or proteins which bind to it (Figure 4). miRNAs for instance typically bind to 3'UTR of specific subsets of mRNAs thereby inhibiting their translation or inducing their degradation (Jonas and Izaurralde 2015). RBPs commonly affect mRNA stability, translation as well as processing or modification (Hentze et al. 2018). About 1400 RBPs have been identified in humans; some of them, like PABPs target widespread RNA sequences while others act on small groups of mRNAs. For instance, human antigen R (HuR) is a well-characterized RBP which binds U- or AU-rich sequences on 3'UTRs (Grammatikakis et al. 2017). This RBP has attracted special interest because many of its targets are involved in cellular functions such as proliferation, apoptosis and differentiation (Abdelmohsen and Gorospe 2010). HuR function is mainly regulated by posttranslational modifications. For instance, phosphorylation of HuR by cyclin-dependent-like kinase 5 (CDK5) was shown to specifically prevent binding to cyclin-A2 (CCNA2) mRNA, one of its target leading to its reduced translation and to cell cycle arrest (Filippova et al. 2012).

1.2.2.1.2 5'UTR elements 5'UTR length, presence of secondary structure or of uORFs play critical roles in translational control (Gebauer et al. 2012; Hinnebusch et al. 2016; Gandin et al. 2016). Studies by Kozak in the 80s suggested that an optimal 5'UTR length for efficiency scanning and translation initiation could be found around 80 nucleotides implying that both shorter and longer UTRs would be less efficiently translated (Kozak 1987). Subsets of long and structured 5'UTR mRNAs have been shown to be particularly sensitive to eIF4E availability i.e. reduction of eIF4E primarily affect these so-called "weak mRNAs" (Graff et al. 2008). They represent a classical example of oncogenic translational control. Indeed, these weak mRNAs are enriched in cellular functions such as survival, proliferation, angiogenesis (e.g. MYC proto-oncogene (MYC), Survivin, Cyclin D1, vascular endothelial growth factor (VEGF)). Most cancers overexpress eIF4E thereby enhancing translation of these oncogenic signatures (De Benedetti and Graff 2004).

Another example where translation is selectively controlled by differential sensitivity to a translation factor was described above (eIF4A-sensitive subset); in this case, the regulation was also mediated by structural features in the 5'UTR.
1.2.2.1.3 Codon-dependent elongation rates and the role of tRNA availability

In addition to modulation of translation impinging on scanning and initiation, dysregulated translation depending on codon composition of mRNAs was recently described during tumorigenesis and therapy resistance (Rapino et al. 2018). In this context, tRNAs can be thought of as trans-acting factors whose regulation may impact translational elongation speed. In case of a global reduction of the levels of all tRNA species, the potential impact on protein synthesis rate is predicted to also be general. tRNA synthesis is for instance known to be stimulated by mTOR via suppression of the Pol III-inhibitor MAF1 (Michels et al. 2010). However, as already discussed, it has been debated whether regulation of specific tRNAs may have an impact on translation in a codon-dependent manner.

In the example of the study by Gingold et al. (2014) described above (Figure 6), differential expression of tRNAs corresponded to differences in codon usage between proliferation and differentiation conditions. Such codon-specific regulation may however occur at any level of tRNA biogenesis. Alterations of tRNA function by post-transcriptional modifications has recently raised interest (El Yacoubi et al. 2012). The 5-methoxycarbonyl-methyl-2-thiouridine (mcm\textsuperscript{5}s\textsuperscript{2}U) modification will, for instance, be of particular interest in Paper II. mcm\textsuperscript{5}s\textsuperscript{2}U strengthen the interaction between tRNA\textsubscript{UUC}\textsuperscript{Glu}, tRNA\textsubscript{UUU}\textsuperscript{Lys}, tRNA\textsubscript{UUG}\textsuperscript{Gln} and the codons they can pair with, namely GAA and GAG (coding for Glutamic acid), AAA and AAG (coding for Lysine), CAA and CAG (coding for Glutamine). This modification is necessary for wobbling (decoding GAG, AAG and CAG) but the cognate tRNAs of these G-ending codons (tRNA\textsubscript{CUC}\textsuperscript{Glu}, tRNA\textsubscript{CUU}\textsuperscript{Lys}, tRNA\textsubscript{CUG}\textsuperscript{Gln}) are also expressed in humans. Ribosome-profiling studies performed in yeast upon loss of the mcm\textsuperscript{5}s\textsuperscript{2}U modification showed increased ribosome occupancy at AAA, CAA and, to a lesser extent, GAA codons which can be interpreted as a reduction of elongation rates at these codons (Zinshteyn and Gilbert 2013; Nedialkova and Leidel 2015). Interestingly, the increases in occupancy at AAG, CAG and GAG codons were unconvincingly small compared to those of A-ending codons. Therefore, mcm\textsuperscript{5}s\textsuperscript{2}U modification impact translation of A-ending codons to a higher extent than G-ending codons. The estimated differences in ribosome dwell time at A-ending codons upon loss of the modification did not lead to a significant reduction of global protein synthesis (indicating that translation initiation remains rate-limiting in these conditions; Zinshteyn and Gilbert 2013). When focusing on the set of proteins encoded by mRNAs which are enriched for AAA, CAA and GAA codons, Rezgui et al. (2013) observed a significant decrease in protein abundance upon loss of the modification. However, in the study by Nedialkova and Leidel (2015), these differences were also observed on cytoplasmic levels of AAA, CAA and GAA enriched mRNAs thus disproving the hypothesis that such regulation would be due to reduced translation elongation rate at specific codons. Instead, they showed that increased ribosomal
pausing perturbs co-translational folding and solubility of proteins.

The mcm\(^5\)s\(^2\)U modification and the tRNA species it impacts is well defined in yeast (Johansson et al. 2008) but remains to be fully characterized in mammalian systems. In melanoma, it has recently been shown that modulating expression and/or activity of the enzymes catalyzing this modification (so called Uridine 34 (U34) modifying enzymes; (Figure 8)) promotes survival and resistance to therapy (Rapino et al. 2018). Similarly, these enzymes has been described to have a key role in breast cancer metastasis (Delaunay et al. 2016) and Wnt-driven intestinal cancers (Ladang et al. 2015). These phenotypes were proposed to rely on differential requirement for tRNAs modified at the U34 position across mRNA-species. Thus, a higher level and/or activity of U34 modification enzymes would allow more efficient translation of mRNAs with high frequency of codons requiring U34 modified tRNAs.

![Figure 8: U34 tRNA modification pathway.](image)

1.2.2.2 Regulatory pathways impinging on translational outputs of subsets of mRNAs Translational co-regulation of subsets of mRNAs also typically occurs downstream of specific regulatory pathways. A few examples will be explained below. The features allowing co-regulation can be found, for instance, along their 5’UTRs (presence of terminal oligopyrimidine (TOP) motifs, complex secondary structures (see section 1.2.2.2.1)).

1.2.2.2.1 mTOR-sensitive mRNAs Activation of mTOR, more specifically of its complex 1 (mTOR complex (mTORC)1), leads to phosphorylation of eIF4E binding proteins (4E-BPs) which are inhibitory proteins for translation initiation. Phosphorylated 4E-BPs cannot bind eIF4E freeing the cap-binding effector for assembly of the
Figure 9: Regulation of selective translation by oncogenic pathways. Hyperactive PI3K-mTOR and MAPK pathways impinge on eIF4F complex assembly.


eIF4F complex and initiation of translation (Figure 9; Brunn et al. 1997). Specific sets of mRNAs are known to be more sensitive than others to alterations in the mTOR pathway. For instance, mRNAs harboring a TOP motif (cysteine after the 5’cap followed by 4-15 pyrimidines), corresponding mainly to mRNAs involved in the translational...
machinery itself such as ribosomal proteins, show coordinated drastic reduction in their translational efficiency upon mTOR inhibition (Hsieh et al. 2012; Thoreen et al. 2012; Larsson et al. 2012). Another subset of mRNAs, those having long and structured 5’UTRs, shows specific sensitivity to mTOR inhibition (Hinnebusch et al. 2016; Larsson et al. 2012; Gandin et al. 2016). This effect depends on signaling via 4E-BPs.

mTOR is a regulator of responses to nutrient and growth factor stimuli. It can be part of several protein complexes including mTORC1 which regulates cell growth and proliferation (by phosphorylating ribosomal S6 kinase (S6K) and 4E-BPs) and mTORC2 which regulates survival mainly via the AKT (also known as Protein kinase B) pathway (Dowling et al. 2010; Sarbassov et al. 2005). mTOR is commonly hyperactive in human cancers (Vivanco and Sawyers 2002; Taylor et al. 2010) and attempts to selectively and efficiently target its pathway as an oncological therapy are still ongoing (Rodrik-Outmezguine et al. 2016; Fan et al. 2017).

ERα has been shown to associate with PI3K, an upstream activator of mTOR and directly or indirectly influence this oncogenic pathway in several tissues (Simoncini et al. 2000; Levin 2009). A recent study linked resistance to estrogen receptor antagonists with hyperactivation of mTOR as well as increased phosphorylation of eIF4E by MAPK-interacting kinase (MNK)1 (Geter et al. 2017).

1.2.2.2.2 Regulation of selective translation by MAPK signaling and eIF4E phosphorylation
eIF4E activity and initiation of cap-dependent mRNA translation can also be modulated by phosphorylation of eIF4E at Serine 209 (Furic et al. 2010). This is mediated via MAPK or MAPK/ERK kinase (also known as MAPKK) (MEK)-extracellular signal-regulated kinase (ERK) signaling (Figure 9). The role of eIF4E phosphorylation is still poorly understood but has been associated with preferential translation of selected mRNAs (Robichaud et al. 2015; Furic et al. 2010). In mouse embryonic fibroblasts (MEFs), translational activation through increased phosphorylation of eIF4E is necessary for epithelial-to-mesenchymal transition (EMT) induction. Therefore, targets of the MAPK pathway constitutes another example of coordinated translational control.
2 Present investigations

2.1 Aims of the thesis

• To extend the methodological support allowing to study transcriptome-wide differences in translational efficiency:
  
  – Expand the polysome-profiling experimental method to make it available for large sets of small samples while maintaining the possibility to quantitatively measure translational control
  
  – Expand the anota computational method for analysis of changes in translational efficiencies to generalize it to polysome- and ribosome- profiling data quantified using DNA-microarrays or RNA sequencing and to adapt it to classical characteristics of RNA sequencing data

• To unravel a novel role for estrogen receptor alpha in cancer: namely to investigate the mechanisms by which this transcription factor also affects gene expression’s translational output
2.2 Results and discussion

2.2.1 Assessing differential translation and buffering using anota2seq

For translatomes quantified by DNA-microarrays, the anota algorithm had been developed to correct changes in polysome-associated mRNA for changes in cytoplasmic mRNA levels by using a linear regression method (in Figure 7A, the anota models are plotted for the treatment (solid line) and control conditions (dotted line); Larsson et al. 2010). This approach was shown to outperform the translational efficiency (TE) score approach (also called log-ratio method) which compares the log(translated mRNA data/cytoplasmic mRNA data) between conditions (Larsson et al. 2010). The first objective of Paper I was to develop anota2seq to allow similar modelling for RNA-sequencing data (accounting for the count nature of such data with a mean variance dependency which cannot be corrected by log-transformation alone) and to compare anota2seq to other methods for transcriptome-wide analysis of differential translation.

In the early days of next-generation sequencing, several studies used classical DNA-microarrays methods on log(RNAseq counts/total library size) (Perkins et al. 2009; Cloonan et al. 2008). It was shown later that RNAseq data should be analyzed using specific methods for count data. However, another strategy was taken by Law et al. (2014) when they established a new data transformation technique which, when applied to RNAseq data, "gives" properties of DNA-microarrays data. It permits that methods optimized for DNA-microarrays can be used on transformed RNAseq data. Using a similar strategy, anota2seq is a method which was adapted from the "DNA-microarrays anota method" (Larsson et al. 2010; Larsson et al. 2011). Indeed, anota2seq allows for transformation or normalization of RNAseq data prior to applying similar models as in anota.

As explained earlier, from RNA quantification of polysome- or ribosome-profiling, one can detect changes in translational efficiency leading to changes in protein levels (a change in the amount of translated mRNA that is not explained by a change in cytoplasmic mRNA) or buffering which opposes changes in protein levels despite alterations in corresponding cytoplasmic mRNA. A third mode of regulation of gene expression corresponds to having similar changes in cytoplasmic mRNA levels and levels of translated mRNA i.e. change in mRNA abundance. Paper II (regulation of gene expression upon depletion of ERα in a prostate cancer cell line) and Paper III (differences between serum starved HCT-116 cells (a colon-cancer cell line) with or without tumor protein p53 (p53)) are two examples where a subset of transcripts show increased or decreased abundance that were buffered at the level of translation. To allow for characterization of this mode of regulation, anota2seq includes the option to perform analysis of translational buffering and categorizes regulated genes in one
of three modes of regulation: changes in translational efficiencies leading to altered protein levels (in short, "translation"), changes in translational efficiencies leading to buffering ("buffering") or changes in mRNA abundance.

Similarly as in anota, the model fitted in anota2seq for analysis of changes in translational efficiency leading to altered protein levels consists of a linear regression with translated mRNA as dependent variable and cytoplasmic mRNA and the sample class variable as independent variables. A common slope for all sample categories is considered and the translational effect is defined as a difference in intercepts between conditions (Larsson et al. 2010). On the other hand, translational buffering is defined as changes in cytoplasmic mRNA level that are not paralleled by changes in levels of translated mRNA (Figure 10). As such, performing analysis of buffering considers cytoplasmic mRNA as dependent variable and translated mRNA as independent variable (together with the sample class). The "mRNA abundance" mode of regulation, corresponds to significant changes in both cytoplasmic and translated mRNA with changes in the same direction.

In order to compare the ability of different methods to identify genes regulated at the level of mRNA translation, we designed a simulation study. RNAseq counts were simulated for cytoplasmic and translated mRNA under two conditions for 4 categories of genes:

- truly unregulated genes (simulated from the same theoretical distributions for both conditions)
- genes regulated by differential translation (cytoplasmic mRNA counts were simulated from the same distribution but a positive or negative fold change (FC) was applied between the 2 conditions for the translated mRNA simulation)
- genes regulated by differential buffering (translated mRNA counts were simulated from the same distribution but a positive or negative FC was applied between the 2 conditions for the cytoplasmic mRNA simulation)
- genes regulated by differences in mRNA abundance (The same FC was applied
between conditions for both cytoplasmic and translated mRNA theoretical distributions)

We concluded from this study that anota2seq outperforms current methods notably by allowing distinction between changes in translational efficiency affecting protein levels and buffering. Indeed, anota2seq showed higher area under the receiver operating characteristic curve (AUC) and precision than other methods for identification of genes in the "differential translation" category (i.e. true differential translation). All methods except anota2seq, are based on the principle that a gene will be identified as differentially translated if the translated mRNA change between conditions is significantly different from the change in cytoplasmic mRNA levels, regardless of whether the difference is higher in translated mRNA or cytoplasmic mRNA. The inability of most methods to distinguish between differential translation leading to altered protein levels and buffering has led in the past to incorrect biological conclusions as explained in Larsson et al. (2010).

As mentioned earlier, regulation by translational buffering has been observed in other contexts (McManus et al. 2014; Artieri and Fraser 2014; Cenik et al. 2015; Lalanne et al. 2018). When discussing these different studies, we realized that translational buffering could be categorized in different contexts. The first context would be when translational buffering occurs in the context of transcript-dosage compensation (McManus et al. 2014; Artieri and Fraser 2014; Lalanne et al. 2018) or between different individuals (Cenik et al. 2015; Dassi et al. 2015) where differences between conditions are static. We envisaged that mechanisms at play in such cases are likely to differ from mechanisms mediating translational buffering upon adaptive responses which can be reverted (as when observing translational buffering upon depletion of \( \text{ER}\alpha \) which can be a therapy target). In this second context (exemplified in Paper II), we would use the term "translational offsetting". Finally, response to acute stress (Tebaldi et al. 2012) or delayed synthesis between mRNA and protein can cause transient translational buffering, called equilibration at the translational level.

As any model, our simulation study has limitations and its applicability scope is unclear. However, the parameters used for the control condition were estimated from a real polysome-profiling dataset (Guan et al. 2017) to maximize the confidence in inferred conclusions from this study. Furthermore, using such a simulation study allowed us to test additional parameters: for instance, how robust the different algorithms for translatome analyses are against increased variance and reduced sequencing depth. All algorithms perform well under increased variance and reduced sequencing depths as long as the number of mapped reads to mRNAs was higher than 5 million. As our simulation data are public, such explorations could easily be ex-
tended to verify for instance the impact of batch effects with increasing sizes as did Chothani et al. (2019). It would also be interesting to evaluate the impact of strong down-regulation of global translation. Indeed, as discussed in (Gandin et al. 2016), assessment of gene-level relative differences in translation between conditions can be influenced by reductions in translation of most genes quantified in parallel (as would be observed for instance upon mTOR inhibition).

Finally, even if the differences between the methods were smaller, anota2seq was shown to outperform current methods for statistical analysis of translational efficiency even in the absence of translational buffering. When further exploring the reason why some methods seemed to underperform, we observed that Xtail (Xiao et al. 2016) typically detects high amounts of non-differentially translated genes when tested under a NULL model (no true differences in gene expression) and that Babel poorly controls type I error (this was also noticed by Xiao et al. (2016) in their method comparison).

Thus, anota2seq allows efficient analysis of translatomes quantified using DNA-microarrays or RNAseq which is essential to further our knowledge of the role of translational control in cancer and other diseases.

2.2.2 Transcriptome-wide analysis of mRNA translation in tissue samples

Dysregulation of translation contributes to both initiation and progression of cancer by inducing global changes in protein synthesis rates and changes in the translational activity of specific mRNAs encoding cancer related proteins (Truitt and Ruggero 2016). Dysregulated mRNA translation can also mediate resistance towards targeted therapies (Boussemart et al. 2014). This occurs notably via activation of eIF4E which is hyper-active in most cancers, including those of the breast (De Benedetti and Graff 2004). Furthermore, preliminary analyzes performed in our research group on a small subsets of breast cancers showed that regulation at the translational level could unravel molecular subgroups of tumors which could not have been distinguished when only looking at the transcriptomic level (unpublished).

Breast cancer is a highly heterogeneous disease at the molecular level. A first indication for this heterogeneity is the wide range of ER expression in population-based cohorts (Osborne 1998; Wenger et al. 1993). Furthermore, despite ER expression correlating with proliferation and tumor aggressiveness (Wenger et al. 1993), overall ER-tumors usually show poorer prognosis than ER+ ones (Knight et al. 1977). This constituted an initial proof that breast tumors can be driven by many different molecular mechanisms. Intra-tumor heterogeneity is also common in this disease (Teixeira et al. 1995). Thus, studying breast cancer biology nowadays requires using patient material
from large cohorts that are representative of the variability of this disease. A cohort of 161 biobanked breast cancer tissue samples was identified with the aim to perform translome analysis.

Analyzing gene expression from tumor material implies the use of optimized experimental methods for low input of RNA. Thus, the polysome-profiling technique first needed to be adapted for large sample sizes with low RNA amounts. Indeed, in the classical linear gradient approach, mRNAs are separated by sedimentation and the entire volume is fractionated along the gradient in 26 fractions (Figure 11A; red numbers). Collecting the polysome-associated mRNA entails pooling a large volume (>3 mL) across many fractions (16 to 25 on Figure 11A) which is labor-intensive and can cause sample loss. Our optimization allowed to concentrate the polysome-associated mRNA in one or two fractions instead (fractions 16-17 on Figure 11B).

Using two conditions of a cell-line model (HCT116 p53-/- and HCT116 p53+/+), we performed polysome profiling using both methods in parallel and compare the results in terms of RNA quantity, quality and output of the analysis of differential translation using an-

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**Figure 11**: Polysome-profiling using the linear and the optimized non-linear gradient methods. Example of a polysome profile when using the classical method (A) or the optimized non-linear gradient (B). Fractions separations are shown as dotted lines. Fractions numbers are provided at the bottom of each plot.

(A) Modified from Gandin, V et al. (2016). "nanoCAGE reveals 5’ UTR features that define specific modes of translation of functionally related MTOR-sensitive mRNAs." In: Genome research 26(5), pp. 636-648. doi: https://doi.org/10.1101/gr.197566.115. ©2016 Gandin et al.; Published by Cold Spring Harbor Laboratory Press. This article, published in Genome Research, is available under a Creative Commons License (Attribution 4.0 International), as described at http://creativecommons.org/licenses/by/4.0/. (B) Modified from Liang C et al. "Polysome-profiling in small tissue samples." Nucleic Acids Res. 2018 Jan 9;46(1):e3. doi: https://doi.org/10.1093/nar/gkx940
ota2seq. High RNA quality and similar amounts were obtained from both methods. Next, we performed a differential expression analysis on polysome-associated mRNA obtained from both methods to assess gene-level differences between HCT116 p53+/+ and p53-/-.

When comparing FCs, a high Spearman correlation (0.74) was observed between those of the linear gradient polysome-associated mRNA data and those from the optimized non-linear gradient data. After adjustment for multiple testing using Benjamini-Hochberg false discovery rates (FDRs), p-values obtained from the optimized non-linear method appeared globally lower. Taken together, these results are consistent with both methods resulting in similar effect sizes and different variability (the optimized method providing data with lower variance among biological replicates).

Finally, we also validated the feasibility for the optimized non-linear gradient method to isolate efficiently translated mRNA from biobanked tissues. For this purpose, we selected 5 breast cancer tissue samples from the large cohort mentioned above (with 161 patient samples) and applied our new polysome-profiling method. RNAseq libraries were prepared for polysome-associated and cytoplasmic mRNA using Smartseq2 which has been developed for low input samples such as single-cell RNAseq (Picelli et al. 2014). High coverage of the translatome could be reached from breast cancer tissue samples and the sequencing depth was above 5 million reads mapped to protein coding mRNA for 3 samples and just below this threshold (between 4.5 and 5 million) for the remaining 2. In Paper I, we showed that statistical analysis of differential translation is not influenced by sequencing depth if it is above 5 million reads mapped to protein coding mRNAs. Anota2seq results were also quite stable in cases where 25% of samples have a sequencing depth of 2.5 million reads mapped to mRNAs and 75% have at least 5 million reads.

Thus studying novel mechanisms of regulation of mRNA translation in large collection of tissue samples is feasible using the optimized non-linear polysome-profiling method and anota2seq.

2.2.3 Investigating the mechanisms by which the transcription factor estrogen receptor alpha affects gene expression at the translational level

As described in section 1.2.1.3, the aim of Paper II was to study the role of ERα in coordinating gene expression at multiple levels. Indeed, in a previous publication (Takahizawa et al. 2015), potential non-nuclear functions of ERα were illustrated by showing that its depletion partially inhibited members of the mTOR and MAPK pathways. Therefore, we used BM67 cells (a prostate cancer cell-line derived from phosphatase and tensin homologue (PTEN)-deficient mice) and performed polysome-profiling fol-
ollowed by quantification of the translatome upon depletion of ERα. As expected from perturbation of a transcription factor, major differences were observed on the cytoplasmic mRNA level of many transcripts. However, at the translational level, observed regulations did not seem consistent with reduction in activity of important translation pathways; as seen by the limited amount of genes classified in anota2seq’s translation mode (i.e. changes in translational efficiency leading to altered protein levels) and by the absence of significant translationally suppressed cellular functions upon ERα depletion. Instead, alterations in mRNA levels were largely translationally offset. This unexpected result was carefully validated using independent technologies: the initial quantification was performed using DNA-microarrays; RNAseq resulted in very similar measurements of cytoplasmic and polysome-associated mRNA in both conditions; a subset of targets (selected in all different modes of regulation of gene expression) were validated using Nanostring; at the protein level, Western blotting (WB) confirmed regulation observed by polysome association for all tested targets. Thus, ERα up- and downregulates the mRNA abundance of many targets but a majority of these alterations are offset at the level of translation. We validated that, at least for a subset of targets, such translational offsetting leads to maintained protein levels despite mRNA modulations.

Next, we sought to pinpoint mRNA characteristics which could mediate translational offsetting upon ERα depletion. For this purpose, we explored features in different parts of the mRNA. Firstly, we performed nanoCAGE on shERα which allowed to precisely determine the position of transcription start sites (TSSs) and thereby analyzing characteristics of the 5'UTRs. Secondly, we explored presence of miRNA target sites on offset vs. non-offset mRNAs. We sequenced small RNAs in order to investigate whether ERα-dependent miRNA regulation could reconcile translational offsetting alterations. Lastly, we observed major differences in codon usage along mRNAs which are induced but translationally offset upon ERα depletion as compared to non-offset mRNAs.

mRNAs which are downregulated but translationally offset appeared to have shorter and less structured 5'UTRs than non-offset transcripts. Their median mRNA length is close to what has been hypothesized to be the "optimal" length for efficient translation (Kozak 1987) and they lack complex structures which could limit their translational efficiency.

Regarding analysis of miRNAs, we initially observed an enrichment of target sites (from the targetScan database (Lewis et al. 2005)) among mRNAs whose upregulation is translationally offset. Therefore, we hypothesized that ERα may upregulate specific miRNAs which would target these offset mRNAs thus mediating translational offset-
ting. Interestingly, our small RNA sequencing revealed that ERα depletion leads to more downregulation than upregulation of miRNAs and that no sign of upregulation was seen for miRNAs targeting mRNAs whose levels are induced but translationally offset. Accordingly, we concluded that ERα-regulated miRNAs do not seem to mediate offsetting at the level of translation. Yet, we detected that mRNAs whose levels are reduced upon ERα depletion but translationally offset (i.e. those lacking complex 5’UTR structures) generally lack miRNA target sites which could also limit their translational efficiency.

Intriguingly, when analyzing characteristics of transcripts whose levels are induced by ERα depletion but translationally offset, a major distinction with non-offset mRNAs is their codon usage. Specifically, ERα-induced but translationally offset transcripts are enriched in codons depending, for their translation, on tRNAs modified at their U34 position.

Initially, we noticed a substantial overlap between codons enriched in upregulated but translationally offset mRNAs and codons enriched in proliferation-related mRNAs in the study by Gingold et al. (see section 1.2.1.2 and Figure 6). Among these codons, the strongest over-representation was seen for AAA and GAA which require U34-modified tRNAs for efficient translation (Figure 8). Consistently, we showed that ERα depletion downregulates protein levels of ELP3 which is one of the enzymes catalyzing the modification. Moreover, translational offsetting of DEK Proto-Oncogene (DEK) (an offset target which we extensively validated) could be reversed by re-expression of ELP3 in BM67 cell-lines where ELP3 had previously been knocked out. Finally, a strong reduction of these mcm5s^2U modifications was quantified by mass spectrometry upon ERα inhibition in a breast cancer cell line. In conclusion, ERα regulates expression of U34-modifications and selectively offsets mRNA abundance at the translational level.

Studying mechanisms of translational offsetting allowed to formulate new perspectives in the role that ERα plays in cancer. Functionally, this implies that upon ERα depletion, transcripts enriched in "proliferation signature codons" will not be synthesized despite high levels of mRNA molecules. Furthermore, we showed that the anti-proliferative effect of ERα-inhibitors was strongly reduced in cells where the U34-modifying pathway had been inactivated. In melanoma, perturbation of the U34-modifying pathway was associated with resistance to B-Raf proto-oncogene, serine/threonine kinase (BRAF) inhibitors (Rapino et al. 2018). In breast cancer, resistance to endocrine therapies is a major concern which limits new survival benefits (Clarke et al. 2015). Studying the U34-modifying pathway as a potential target to overcome treatment resistance in ERα-dependent cancers may be a promising strategy.
Mechanistically, an intriguing remaining question is whether and to which extent reduced abundance of U34-modified tRNAs would influence translation elongation vs. initiation rates. However, this might be challenging to decipher as reduced elongation rates can themselves limit initiation by accumulation of ribosomes towards the 5' end of the coding sequence. In our study, we observed translational offsetting of a large amount of mRNAs from polysome-profiling data which implies that for such transcripts, the same amount of mRNA is associated with >3 ribosomes (we used this threshold to isolate efficiently translated mRNA) following ERα depletion whereas the mRNA level is increased. We showed that this is mediated by reduced availability of U34-modified tRNAs. Ranjan and Rodnina (2017) proposed that U34 enzymes causes ribosome pausing during translation elongation and suggested that lack of thiolation of tRNA\textsuperscript{UUU}\textsubscript{Lys} increases the residence time of ribosomes on AAA codons by about 40% in a prokaryotic translation system. We investigated further whether such an increase in translocation time at codons requiring U34-modified tRNAs could explain the translational offsetting observed in Paper II. For this purpose, we selected one target which was induced but translationally offset upon ERα depletion: DEK. We chose this target because it has an extreme frequency of AAA, CAA and GAA codons compared to other mouse transcripts (Figure 12A). We validated its FC on cytoplasmic mRNA using qPCR (Figure 12B). We used ribosome flow model (RFM)s (Reuveni et al. 2011) to estimate translational output of the DEK mRNA upon several hypothetical reduction of translation elongation rate at codons decoded by U34-modified tRNAs. As input of the RFM, we needed to give the coding sequence of DEK, a range of initiation rates and codon-specific residence times. We sought to assess if the combination of increased residence time coupled with increased mRNA level could lead to maintained protein synthesis as compared to the control condition (Figure 12C). The RFMs indicated that ribosome residence time on codons requiring U34-modified tRNAs cannot alone explain observed translational offsetting under any initiation rate (Figure 12D). Therefore, ribosome drop-off could be a potential mechanism restricting translation under conditions where tRNAs are hypomodified at the U34 position but additional experiments are required to explore this further and eventually to understand how translational control can be used to maintain homeostasis.

2.2.3.1 Reflections on ethical considerations in bioinformatics projects My doctoral studies have focused on statistical method development and bioinformatics analyzes of translatomes from experiments mainly performed on cell lines. Reflections on ethical considerations instinctively arise in research projects for instance when experiments need to be performed on animals and these reflections are then of utmost importance. However, in other contexts such as data analysis, ethical discussions are often overlooked. Paper II provided an example of interesting ethical considerations in data analysis which will be described here.
Figure 12: Increased ribosome residence time at AAA, CAA and GAA codons cannot alone explain DEK translational offsetting. We use the RFM to assess whether an increase in ribosome residence time on codons requiring U34 tRNA modifications can alone explain such a translational offsetting. We fit 4 RFMs: one with default residence codon times (calculated based on mouse tRNA copy numbers) and three other models with residence time on AAA, CAA and GAA codons increased by 20%, 50%, 100%. (A) Distribution of frequencies of AAA, CAA and GAA in coding sequences of the mouse transcriptome (from consensus coding sequence (CCDS) database). (B) DEK mRNA relative expression from shERα and control cells measured by qPCR. (C) Residence time (used as input in the RFM) of the ribosome at each position along the DEK coding sequence for baseline condition (gray) and when the residence time on AAA, CAA, GAA codons is increased by 20% (yellow), 50% (orange), 100% (red). Residence time is averaged across chunks of 12 codons. (D) Estimated protein abundance predictor as defined in Reuveni et al. (2011) showing the impact of increased residence times in case of unchanged abundance (plain lines) and when DEK mRNA levels is increased by 1.95 fold (dotted lines).
One classical example which raises questions related to scientific reasoning and ethics comes when the initial hypothesis of a study cannot be confirmed by the analysis. More specifically, a study has been designed based on a scientific hypothesis; it was conducted according to this design and then led to a negative result. Let’s set apart the discussion about what should be considered as a negative result for now and focus on the following question: should one publish what would be considered as a negative result or should other options be considered? A common "excuse" for not trying to publish negative results is that it would be refused by journal editors themselves. Publication bias, also called positive-outcome bias, is an unfortunate reality in medical research. In clinical trial research for instance, studies with statistically significant results are definitely more likely to be published and more likely to get published early. This kind of bias can have deleterious consequences, notably because meta-analyses can only be based on published results. This issue has recently begun to be addressed by some journals which are dedicated to publication of negative results. Another question is: would it be unethical to publish these results as the "positive" results of another post-hoc hypothesis? Should this alternative be considered as a misleading way to make one's research more interesting than what is it in reality? Or should it be considered as the starting point of a potentially promising new hypothesis? Firstly, the answer to these questions depends on whether the research question is to be considered as having a "hypothesis confirmation" or "hypothesis generation" goal. The former would imply that a lot of research has previously been performed to lead to the hypothesis. In this case, publishing the negative result should probably be the reasonable option. An example of this would be phase III clinical trials, which are confirmatory trials and may lead to approval of new drugs or indications. In this case, a statistical analysis plan should be pre-defined and any additional analyzes should be interpreted as having lower levels of evidence. There should however be more freedom in research that is further away to applications for patients and research that is performed to generate new hypotheses. In this case, it should be acceptable to consider a post-hoc analysis as promising even if its level of evidence is not as high as the initial hypothesis.

In **Paper II**, the initial hypothesis was that ERα would not only regulate transcription of specific targets but also translational output. Indeed, ERα has been shown to interact with classical pathways which impinge on translational control such as the mTOR pathway. We therefore wanted to test whether ERα’s transcriptional output would also be regulated at the translational level notably via this pathway. As described above, we could however not confirm from our transcriptome-wide studies of changes in translation that ERα depletion impacted the mTOR pathway. However, the analysis of data from these experiments revealed other interesting mechanisms upon
ERα depletion (see Paper II). We published our results in the context of the second hypothesis, stating that this was not the initial hypothesis. Furthermore, additional validation experiments have been performed in order to give more strength to this post-hoc hypothesis. Several reflections were raised before writing the manuscript to decide in which order the "story" should be told and whether to omit the initial negative result. In the final version of this paper, the initial hypothesis is discussed in the introduction. Of course, ethics was not the only consideration when designing the manuscript but even in projects where the context of the research offers freedom in the way the analysis can be done, it is interesting to reflect upon the integrity and validity of scientific results from post-hoc hypotheses. Having stated that more freedom should be allowed in the data analysis of basic research, this paragraph should not be concluded before adding that data dredging is of course not to be advised.
3 Conclusion

Because of its tight connection with uncontrolled proliferation (Faller et al. 2015), immune response (Piccirillo et al. 2014), altered metabolism (Cunningham et al. 2014), invasion and metastasis (Robichaud et al. 2015); translational dysregulation is now itself considered as a hallmark of cancer (Vaklavas et al. 2017). This thesis provides methodological advances allowing to study further mechanisms by which translation influences cancer cell phenotypes. Indeed, our optimized gradient polysome-profiling method allows to isolate translatomes from tissue samples. Furthermore, we demonstrated that the anota2seq algorithm outperforms current methods for statistical identification of differences in translational efficiencies.

Furthermore, this thesis shines a light on an underappreciated mode of gene expression regulation whereby translation acts as an offsetting process which opposes protein levels despite fluctuations in corresponding mRNA abundance. Common mechanisms regulating translational offsetting are yet to be discovered but upon depletion of ERα in prostate cancer, the most influential regulatory pathway impinged on modification of specific tRNAs. This result could have implications in our understanding of how the proteome of hormone-dependent diseases is controlled as well as in strategies to tackle resistance to ERα-targeted therapies.
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On an even more personal note, I should maybe say that I would not claim that the PhD path has always been fun for me (yes, it mostly should!) and I want to take this as an opportunity to thank people whose presence along the way has been so essential: [in Frenglish in the text] les sis’, les couz’ (& assimilés), les cops’ (from around here, d’ailleurs, de Génération.s Stockholm), les parents (& beau-) and most particularly Robin!
References


Li, JJ, Chew, GL and Biggin, MD (2017a). “Quantitating translational control: mRNA abundance-dependent and independent contributions and the mRNA sequences that specify them”. In: Nucleic Acids Research 45(20), pp. 11821–11836.


more than 127,000 breast cancer patients”. In: *Breast Cancer Research and Treatment* 28(1), pp. 9–20.


