mRNA degradation controls differentiation state-dependent differences in transcript and splice variant abundance

Peter A. C. ’t Hoen1*, Michael Hirsch2, Emile J. de Meijer1, Renée X. de Menezes1,3, Gertjan B. van Ommen1 and Johan T. den Dunnen1

1Center for Human and Clinical Genetics and Leiden Genome Technology Center, Leiden University Medical Center, 2300 RC Leiden, The Netherlands, 2School of Information Systems, Computing and Mathematics, Brunel University, Uxbridge, Middlesex, UB8 3PH, UK and 3Department Epidemiology and Biostatistics, VU Medisch Centrum, Amsterdam, The Netherlands

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ABSTRACT

Expression profiling experiments usually provide a static snapshot of messenger RNA (mRNA) levels. Improved understanding of the dynamics of mRNA synthesis and degradation will aid the development of sound bioinformatic models for control of gene expression. We studied mRNA stability in proliferating and differentiated myogenic cells using whole-genome exon arrays and reported the decay rates (half life) for ~7000 mRNAs. We showed that the stability of many mRNAs strongly depends on the differentiation status and contributes to differences in abundance of these mRNAs. In addition, alternative splicing turns out to be coupled to mRNA degradation. Although different splice forms may be produced at comparable levels, their relative abundance is partly determined by their different stabilities in proliferating and differentiated cells. Where the 3'-untranslated region (3'-UTR) was previously thought to contain most RNA stabilizing and destabilizing elements, we showed that this also holds for transcript isoforms sharing the same 3'-UTR. There are two splice variants in Itga7, of which the isoform with an extra internal exon is highly stable in differentiated cells but preferentially degraded in the cytoplasm of proliferating cells. In conclusion, control of stability and degradation emerge as important determinants for differential expression of mRNA transcripts and splice variants.

INTRODUCTION

Although the abundance of messenger RNAs (mRNAs) in the cell is the net result of mRNA synthesis and degradation, expression profiling experiments usually take a snapshot of the mRNA levels in the cell and do not capture the dynamics of mRNA synthesis and breakdown. This lack of information on the dynamic component of the regulation of mRNA levels limits biological investigations in general and proper modeling of transcriptional networks in systems biology approaches in particular. When mRNA synthesis is shut down completely, the effect may be immediate for very short-lived mRNAs but only manifest after hours or days for mRNAs with long half lives. The current study fills in part of this gap and surveys mRNA degradation rates during muscle differentiation on a genome-wide scale. With the availability of these data, the genetic network orchestrating muscle differentiation can be revised and improved.

From the results of previous genome-wide analyses of mRNA degradation rates, we know that there are large differences between the stabilities of individual mRNAs, and that mRNA stability is strongly related to the molecular function of the encoded protein (1–8). However, the mechanisms underlying differential stability of transcripts remain poorly understood. MicroRNA (miRNA)-mediated mRNA degradation emerged as an important controlling factor for mRNA half life. Earlier research also characterized a plethora of RNA binding proteins that may affect mRNA stability (9). Recent studies in differentiating embryonic stem cells (7) and T-cells (6) demonstrated that mRNA stability may be affected by the cell’s biological state. Furthermore, an important subgroup of transcripts use alternative polyadenylation sites depending on the differentiation status (10). This results in 3'-untranslated regions (3'-UTRs) with variable lengths and probably contributes to the differential stability of these transcripts (10). The deciding factors determining the choice for shorter or longer 3'-UTRs are currently unknown.

*To whom correspondence should be addressed. Tel: +31 71 5269421; Fax: +31 71 5268285; Email: p.a.c.hoen@lumc.nl

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We were interested in the control of decay rates during myogenic differentiation and studied proliferating and differentiated myogenic C2C12 cells, a well-established system to study differentiation of muscle cells. Cells were treated with actinomycin D to arrest de novo RNA synthesis, with sampling at short time intervals. We used Affymetrix whole-genome exon arrays covering all known and predicted exons. Where previous genome-wide studies determined expression levels and stability of the 3' ends of transcripts, these arrays allowed us to assess expression level per exon and thereby, for the first time, the effect of alternative splicing on mRNA decay rates. Our results demonstrated that changes in mRNA abundance during differentiation of muscle cells are correlated with changes in decay rates and that the ratios of specific splice variants are controlled at the level of mRNA stability.

**MATERIALS AND METHODS**

**Cell culture, RNA isolation, microarray hybridization**

C2C12 cells were grown in proliferation medium [Dulbecco's modified Eagle's medium (DMEM) with 10% fetal bovine serum (FBS), 2% Glutamax, 1% glucose and penicillin/streptomycin]. Myogenic differentiation was induced by shifting to differentiation medium (DMEM with 2% FBS, 2% Glutamax, 1% glucose and penicillin/streptomycin). C2C12 cells were maintained in differentiation medium for 8 days, when long and multinucleated myotubes had formed. Cultures of proliferating and differentiated C2C12 muscle cells in 6-well dishes containing 3-ml medium were treated with 5 μg/ml actinomycin D to arrest the transcription machinery. At seven different time points after addition of actinomycin D (0, 10, 20, 30, 60, 150, 480 min), cells were lysed and RNA was isolated with the Machery Nagel Nucleospin RNA II Kit. There was no notable decrease in cell viability within this time period. The experiment was performed in duplicate. After depletion of ribosomal RNAs, mRNA was reverse transcribed with random primers containing a T7 tag, and labeled with biotin during the T7 RNA amplification (which retains strand specificity), according to standard Affymetrix protocols. Equal amounts of amplified RNA were hybridized to 28 Affymetrix Mouse Exon v1 arrays [7 time points × 2 replicates × 2 conditions (proliferating/differentiated)].

**Affymetrix exon arrays**

Affymetrix Mouse Exon v1 arrays contain 1236087 non-control probe sets. Most probe sets consist of four probes and the intensity value of each probe set is the robust mean of the individual probe sets. Most exons are covered by at least one probe set. There is a subset of exons, called core exons, which represent all the exons for which there is substantial experimental proof for their existence. Next to the core set, the array covers a huge set of computationally predicted exons. The intensities of probe sets in most of these latter exons (in our experiment 91 versus 51% for core exons) are not above background, and these have not been considered in the analysis described here. Data from these arrays can be analyzed at the exon or the gene level. The presented gene level summarization, giving only one expression value per gene, considers the intensities of probes in core exons only.

**Data analysis—normalization**

In our experimental set-up, normalization is not straightforward, since an important assumption underlying standard normalization procedures, equality of the total amount of mRNA under all conditions studied, is violated. Despite the decrease in the total amount of mRNA in the cell at later time points, equal amounts of RNA have been hybridized to the chips. This means that the samples from the later time points will be enriched for stable transcripts and depleted for fast decaying transcripts. We tried to use spike-in RNAs to correct for the differences in input RNA but this was not successful due to the low number of spike-in RNAs used. Thus the only alternative was to use a standard normalization method, with the intensities of more stable transcripts increasing at later time points.

The median polish summarization method [as implemented in RMA (11)] was applied to summarize the intensities of probes in a probe set and probes in a gene. VSN [variance stabilization and normalization (12)] was used for normalization. VSN and RMA normalization generated similar values in the high intensity range. In the lower intensity range, the variance stabilizing properties of the VSN algorithm corrects for noise-induced artificially inflated differences between arrays in near-background reads. As an additional advantage, the VSN data are on a natural logarithm scale compatible with our exponential decay model, whereas the RMA data are on a log₂ scale. Since the arrays contain many negative and positive control spots, the effect of removing those before the normalization was investigated. This had no effect on the normalization of the non-control spots. As a threshold for background intensity levels, we used the mean + 2 SD of around 4000 negative control probes on the array.

All normalization and subsequent data analysis was performed in the statistical programming language R (13). Raw and normalized data have been submitted to GEO and are available under series GSE14387.

**Data analysis—model fit**

To model decay rates, a standard linear model (1) was fitted with normalized intensity Y of each probe set j as dependent variable and time as independent, continuous variable.

\[ Y_j = \alpha_j + \beta_j \times \text{time} + \epsilon_j \]  

where time is a vector (0, 10, 20, 30, 60, 150, 480); \( \alpha_j \) represents the intercept (estimated expression level at \( t = 0 \)); \( \beta_j \) represents the slope (negative of the decay constant).

The appropriateness of this model is discussed in the Supplementary Methods and in the caption of Supplementary Figure S1. Replicate time points were independent data points in the model. Two separate models were fit for the proliferating and differentiated cells. Decay
constants were said to be significantly different between proliferating and differentiated cells when there was no overlap between their 95% confidence intervals (as determined from the standard error on β, and given the t-distribution and 12 degrees of freedom). Decay constants for entire transcripts were calculated using the exact same model, but after summarizing core probe sets in a transcript into a single expression value by RMA. The half life is given by −ln(2)/β.

For identification of probe sets with a behavior different from the other probe sets in the gene (possible alternative splicing events), we considered the mixed effect model (2) (nlme package; lme function in R), where time is a continuous variable and treated as a fixed effect, where the probe sets were considered as random effects and where we included an interaction term between the probe sets and the differentiation status. In formula form (14):

\[ Y_{ijk} = \beta \times \text{time} + b_j + b_{jk} + \varepsilon_{ijk} \]  

where \( b_j \sim N(0, \sigma_j^2) \), \( b_{jk} \sim N(0, \sigma_k^2) \), \( \varepsilon_{ijk} \sim N(0, \sigma_i^2) \) and \( Y_{ijk} \) represents normalized intensity of probe set \( j \) (\( i = 3, \ldots, n \) probe sets in the gene) with differentiation status \( k \) (\( k = 1, 2 \)) in gene \( i \); \( \beta \) represents time vector (0, 10, 20, 30, 60, 150, 480); \( \beta \) represents fixed effect; \( b_j \) represents random intercept; \( b_{jk} \) represents the interaction between the probe set and differentiation status and \( \varepsilon_{ijk} \) represents error term.

Data analysis—annotation and visualization

The R package Xmap (15), interfacing to a local Ensembl install (mouse release 47; NCBI genome build 37) was used for annotation and visualization of the exon-level data.

Quantitative PCR analysis

Quantitative polymerase chain reaction (qPCR) analysis with SYBR-Green detection was performed as described earlier (16). Bgn was used in the RNA decay studies as a reference gene to correct for differences in input cDNA. Bgn was found to be extremely stable and did not change significantly in Ct value over the time period analyzed. Gapdh was used as a reference gene for the nuclear versus cytoplasmic comparison.

3′-RACE

To determine the polyadenylation sites in the Itga7 transcript, we performed 3′-RACE (rapid amplification of cDNA ends). We performed a reverse transcription (RevertAid RNAseH-M-MuLV, MBI-Fermentas) on 500 ng of total RNA starting at the polyA site with an anchored primer oligo-dT primer (sequence: GCTGCAGACCGGTTTAAACGGCaCCGCACTTTTTTTTTTTTTTTTT). Subsequently, we performed two rounds of PCR [30 cycles, Faststart Taq (Roche)]. The forward primer was designed in exon 22 (sequence: CTGTTAGTCCTGGCTCTGCT); the sequence of the reverse primer in the first round was GCTGCAGACCGGTTTAAAC, whereas the sequence of the reverse primer in the second round of nested PCR was GCGTATAACACCGCAGGC.

GT. Bands were excised from agarose gel and sequenced on an ABI-3700 sequencer.

Isolation of nuclei

Cells were scraped in phosphate buffered saline and pelleted by centrifugation for 5 min at 1500g. The cells were resuspended in cold lysis buffer (10 mM Tris-HCl pH 7.4, 3 mM MgCl2, 10 mM NaCl, 1 mM EDTA) and pelleted by centrifugation for 5 min at 1500g. After resuspending in lysis buffer including 0.5% (v/v) Triton X-100, cells were disrupted in a Dounce homogenizer with 4 × 10 strokes of the pestle. Nuclei were pelleted by centrifugation for 2 × 5 min at 1500g and washed with lysis buffer (without Triton X-100). Nuclei were purified by sucrose gradient centrifugation. To this end, a centrifuge tube was filled with 4 ml of high sucrose buffer (2 M sucrose, 5 mM MgAc, 0.1 mM EDTA, 10 mM Tris–HCl pH 8.0, 1 mM DTT) and nuclei were added after mixing with 4.4 ml of low sucrose buffer [0.32 M sucrose, 3 mM CaCl2, 2 mM MgAc, 0.1 mM EDTA, 10 mM Tris–HCl pH 8.0, 1 mM DTT, 0.5% (v/v) Triton X-100], with subsequent centrifugation for 30 min at 30 000g in an SW41 rotor. Pellet was resuspended in 300 µl of storage buffer [50 mM Tris–HCl pH 8.3, 40% (v/v) glycerol, 2 mM MgCl2, 0.1 mM EDTA], aliquotted and snap frozen in liquid nitrogen.

RESULTS

Experimental setting

Cultures of proliferating C2C12 muscle precursor cells and cultures of C2C12 cells differentiated to elongated and fused myotubes were treated with actinomycin D to arrest the transcription machinery. At seven different time points after addition of actinomycin D (0, 10, 20, 30, 60, 150, 480 min), RNA was isolated and hybridized to Affymetrix Mouse Exon v1 arrays (in duplicate, 28 arrays in total). With the selected time intervals, it is possible to accurately estimate the degradation rates of relatively short-lived transcripts. For very stable transcripts (\( t_{1/2} > 480 \) min) the estimate becomes inaccurate, but longer incubation times resulted in a significant decrease in cell viability severely complicating the analysis.

To model mRNA degradation, we consider an exponential decay model. This model becomes linear on the logarithmic scale used to report the normalized exon-array signals (model 1; see Supplementary Methods and Supplementary Figure S1 for details on the model and the goodness-of-fit). The slope in this linear model is the (negative of the) decay constant, whereas the intercept reflects the expression level before addition of actinomycin D.

Estimation of decay rates

All mRNAs decay over time. The normalization method used (see ‘Materials and Methods’ section), centers the average slopes around zero (Figure 1A). Thus, stable transcripts appear to have positive slopes relative to the average, whereas transcripts with fast degradation will have negative slopes. The relative differences in decay rates (slow decaying versus fast decaying) can be well
estimated using this model (Figure 1B, Supplementary Figure S1 and Supplementary Methods). The maximum slope for expressed genes is 0.0028 intensity units/minute, the mean is \(0.00042\) (median \(0.00020\)), and the minimum \(0.0073\). If we assume that the slowest decaying transcript is stable over the period analyzed, all slopes would need to be corrected with \(0.0028\). The half life is calculated as \(\ln(2)/\text{slope}\) and the shortest half life we observed is then 69 min, the median half life of all expressed transcripts 231 min. This is in a similar range as reported before for proliferating C2C12 cells (8). The decay rates determined on the microarrays correlate well with those measured by quantitative PCR [Figure 2, Pearson’s correlation coefficient: 0.94 (\(P = 3.2 \times 10^{-6}\))], in spite of the array data appearing to overestimate the half life of short-lived transcripts and to underestimate those of stable transcripts. Relative decay rates for all genes with detectable expression (6852 in total) are reported in Supplementary Table S1.

Factors determining mRNA half life

It has been noted that mRNA stability appears to correlate with the cellular function of the encoded proteins (1–8). To study this correlation in our data we functionally classified the most stable and unstable mRNA transcripts in proliferating and differentiated myoblasts measured with qPCR with primers designed in the 3′-UTR are plotted on the \(x\)-axis, and the slopes measured with microarray probe sets in the same regions are plotted on the \(y\)-axis. The correlation coefficient is the Pearson correlation coefficient.

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**Figure 1.** Estimated relative decay rates measured on microarrays. Distribution of the slopes (negative of degradation rate constants) (A) and \(R^2\) values of the linear model (B) for all probe sets (dashed line) and for probe sets in genes expressed above background (solid line). The majority of probe sets do not give detectable signals and have very flat (slope near zero) and noisy (low \(R^2\) values) decay curves. In contrast, expressed genes frequently demonstrate more positive or negative slopes, and their decay rate constants can be reliably measured (high \(R^2\) values). This is expected since background signals remain constant over time, thus generating a slope close to zero, whereas true signals tend to increase or decrease, generating a slope away from zero.

**Figure 2.** Correlation of microarray- and quantitative PCR-based degradation rates. For both microarray and qPCR assays, the natural logarithm of the expression level was plotted against the time after start of actinomycin D treatment and a linear curve was fitted. The slope of this curve is the negative of the decay constant. The slopes for the Bmp4, Ccng2, Cebp, Pde4dip-short, Pde4dip-long and Itga7-exon27 genes in proliferating and differentiated myoblasts measured with qPCR with primers designed in the 3′-UTR are plotted on the \(x\)-axis, and the slopes measured with microarray probe sets in the same regions are plotted on the \(y\)-axis. The correlation coefficient is the Pearson correlation coefficient.
expression, including histones. RNAs coding for structural and contractile muscle proteins, which are only expressed in differentiated cells, appear to be very stable. Transcripts in the category ‘muscle development’ are either very short- or very long-lived, because this category contains both regulatory and structural proteins.

Short mRNAs tend to be less stable than average (Supplementary Figure S2). The median half lives of mRNAs consisting of one, two, or three exons in proliferating cells are 172, 189 and 199 min, respectively. This is significantly shorter ($P \leq 1 \times 10^{-5}$ Kolmogorov–Smirnov test) than the average half life in proliferating cells (237 min). Similar results were obtained for differentiated cells. Figure 3 visualizes the statistically significant shift to shorter half lives in mRNAs with three, two or one exons.

There is no correlation between 3′-UTR length and transcript stability (Supplementary Figure S3).

### mRNA stability is dependent on the cellular state

We calculated the differences in decay rates between differentiated and proliferating cells. There were 170 mRNAs displaying significant differences in stability between the two states; 99 of those were more stable in proliferating cells and 71 were more stable in differentiated cells (Supplementary Table S1). Interestingly, many of the mRNAs with differences in stability were also differentially expressed between the two conditions. Table 2 summarizes some of the results for transcripts playing an established role in muscle development. Slower degradation in differentiated than in proliferating cells goes with higher expression levels in differentiated cells, and vice versa. Myog and Cdkn1a (p21), two important regulators of differentiation previously shown to have increased expression and stability in differentiated cells (17–19) are amongst the genes with clearly different mRNA degradation rates: The half life of the Myog mRNA increased from 165 min in proliferating cells to 268 min in differentiated cells; the half life of the Cdkn1a mRNA increased from 138 to 371 min. The generality of these observations is illustrated by the strong positive correlation obtained after plotting the differences in gene expression against the differences in decay constants (Figure 4, Pearson’s correlation coefficient: 0.57; $P < 2.2 \times 10^{-16}$). Thus, control of mRNA stability and degradation emerges as an important determinant for the regulation of steady-state mRNA levels.

### Relative abundance of splice isoforms is controlled by mRNA stability

The observation that the cellular state determines the overall decay rates of transcripts triggered us to investigate the role of mRNA degradation in controlling the ratio between different splice isoforms. To this end, we used the mixed effect model (2) to identify genes where the decay constant of a particular probe set differed more than 2 SDs from the decay constant of all other probe sets in the gene. Sixty-nine probe sets met these criteria (Supplementary Table S2). After further filtering for

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Table 1. Functional classification of fast and slow decaying transcripts in proliferating and differentiated cells

<table>
<thead>
<tr>
<th>Category</th>
<th>Fast P-value</th>
<th>Slow P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proliferating</td>
<td>3.7 $\times 10^{-8}$</td>
<td>3.6 $\times 10^{-4}$</td>
</tr>
<tr>
<td>Establishment and/or maintenance of chromatin architecture</td>
<td>7.1 $\times 10^{-4}$</td>
<td></td>
</tr>
<tr>
<td>Differentiated</td>
<td>4.6 $\times 10^{-10}$</td>
<td>1.1 $\times 10^{-4}$</td>
</tr>
<tr>
<td>Regulation of cellular processes</td>
<td>8.3 $\times 10^{-5}$</td>
<td></td>
</tr>
<tr>
<td>Regulation of transcription</td>
<td>3.1 $\times 10^{-7}$</td>
<td></td>
</tr>
<tr>
<td>Negative regulation of protein kinase activity</td>
<td>5.0 $\times 10^{-5}$</td>
<td></td>
</tr>
<tr>
<td>Cell cycle</td>
<td>2.1 $\times 10^{-7}$</td>
<td></td>
</tr>
<tr>
<td>Development</td>
<td>1.3 $\times 10^{-4}$</td>
<td></td>
</tr>
<tr>
<td>Muscle development</td>
<td>1.1 $\times 10^{-4}$</td>
<td></td>
</tr>
<tr>
<td>Muscle contraction</td>
<td>7.6 $\times 10^{-28}$</td>
<td></td>
</tr>
</tbody>
</table>

Gene annotation was derived from Gene Ontology. The biological processes significantly overrepresented in the 100 fastest and slowest decaying transcripts in proliferating and differentiated cells are displayed, together with the $P$-value from the hypergeometric test used for testing overrepresentation. Given multiple testing issues, $P < 1 \times 10^{-4}$ was considered to be significant.

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![Figure 3](https://academic.oup.com/nar/article-abstract/39/2/556/2409251/1)
Table 2. Half lives of myogenesis-related genes change during differentiation

<table>
<thead>
<tr>
<th>Gene</th>
<th>Fold-change differentiated versus proliferating cells</th>
<th>Half life proliferating cells (min)</th>
<th>Half life differentiated cells (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tnc</td>
<td>-3.86</td>
<td>444</td>
<td>229</td>
</tr>
<tr>
<td>Mbnl3</td>
<td>-2.72</td>
<td>342</td>
<td>264</td>
</tr>
<tr>
<td>Musk</td>
<td>2.82</td>
<td>246</td>
<td>438</td>
</tr>
<tr>
<td>Lrp4</td>
<td>3.29</td>
<td>236</td>
<td>416</td>
</tr>
<tr>
<td>Cdkn1a</td>
<td>3.73</td>
<td>138</td>
<td>371</td>
</tr>
<tr>
<td>Tnnt2</td>
<td>4.75</td>
<td>242</td>
<td>500</td>
</tr>
<tr>
<td>Tmod1</td>
<td>7.23</td>
<td>201</td>
<td>455</td>
</tr>
<tr>
<td>Acta1</td>
<td>7.75</td>
<td>322</td>
<td>555</td>
</tr>
<tr>
<td>Myog</td>
<td>9.09</td>
<td>165</td>
<td>268</td>
</tr>
<tr>
<td>Cacna1s</td>
<td>17.01</td>
<td>229</td>
<td>798</td>
</tr>
</tbody>
</table>

Only genes annotated with ‘muscle development’ in Gene Ontology and a minimum absolute fold change of two are shown.

Table 3. Location of probe sets with aberrant slopes as compared with other probe sets in the gene

<table>
<thead>
<tr>
<th>Number of probe sets</th>
<th>Probe set location</th>
<th>Genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>14</td>
<td>5'-UTR, alternative 5'-UTR, or coding part of first exon</td>
<td>Thrg4, Foxm1, Casq1, Trim37, Ly75, Casq2, Ezh2, Ap2a1, Ncapd3, Serpine1, Oshp2, Mamdc2, Bub1b, Mybph</td>
</tr>
<tr>
<td>10</td>
<td>3'-UTR or alternative 3'-UTR</td>
<td>Klf20a, Ankrd2, Myom2, Foxo6, Cp, Men5, Igb1, Cacna1s, Pde4dip, Xirp1</td>
</tr>
<tr>
<td>9</td>
<td>Internal exon subject to alternative splicing</td>
<td>Ads11, Myom1*, Igb7*, Pygm, Rbn9, Sec24c, Tnnt3</td>
</tr>
<tr>
<td>6</td>
<td>Internal exon not known to be subject to alternative splicing</td>
<td>Tcfap4, Atp1a2*, Cabc1, Myom2, Mmp15</td>
</tr>
</tbody>
</table>

*Two probe sets in two different exons affected.

*According to Ensembl annotation.
Figure 5. Association between RNA degradation and alternative splicing in Pde4dip. (A) Exon structure of the Pde4dip-short and the Pde4dip-long mRNA and the natural logarithm of the ratio (y-axis) of the expression levels of the different exons (x-axis) between differentiated and proliferating cells, as assayed before actinomycin D treatment. Note: using CAGE-seq (cap analysis gene of expression analyzed by high-throughput sequencing, investigating transcription start sites) data, we found that the annotation of the first exon from ENSMUST00000090750 in ENSEMBL is incorrect (20), explaining its aberrant behavior. (B) Plot of the decay rates in differentiated (closed symbols) and proliferating (open symbols) cells for the different probe sets in the Pde4dip gene, ordered from 5' to the 3' end of the gene. Less stable parts of the RNA have more negative slopes. (C) Relative abundance (arbitrary units; a.b.u) of the short and long transcripts of the Pde4dip gene, as assayed by qPCR with primers in the two different 3'-UTRs. (D) Estimated half lives of the short and long transcripts of the Pde4dip gene, as assayed by qPCR with primers in the two different 3'-UTRs. Error bars reflect standard error of the mean.
**Figure 6.** Association between RNA degradation and alternative splicing in *Itga7*. (A) Agarose gel displaying the products of 3′-RACE for *Itga7* with forward primer in exon 22 in proliferating (P) and differentiated (D) cells and a 200-bp size marker (M). The longer fragment (more abundant in differentiated cells) contains exon 23, whereas the shorter fragment lacks exon 23. The size difference equals the size of exon 23 (113 bp). Subsequent sequencing confirmed this and mapped the polyadenylation site for both fragments at the previously identified and annotated polyA
stable in differentiating than in proliferating cells. qPCR experiments (Figure 5D) validate the longer half life of the long isoform in differentiated cells. Thus, the ratio between the long and the short isoform is regulated, at least in part, at the level of mRNA degradation, and depends on the differentiation status of the cells. 

Iga7. Iga7 is an example of a gene that codes for different transcript isoforms that share the same 3'-UTR but differ in stability. Iga7 codes for integrin α7, a transmembrane receptor important for maintaining the link between the extracellular matrix and the cytoskeleton of the muscle fiber (22,23). Iga7 is known to undergo alternative splicing, resulting in differences in both the extracellular (N-terminal) and cytoplasmic (C-terminal) domain. The splice isoform containing an additional exon (here indicated as 23, referred to in other publications as transcript α7A) is specific for skeletal muscle and expressed mainly during myogenic differentiation (24). Since this exon is out-of-frame, inclusion of this exon gives rise to a protein with an alternative C-terminal amino acid sequence, which may activate alternative signaling cascades. By 3’-RACE, with a primer in exon 22, we demonstrated that the transcripts with or without exon 23 indeed share the same 3’-UTR and use the same polyadenylation site (Figure 6A). As expected, transcripts containing exon 23 are almost exclusively present in differentiated cells (Figure 6B and D). Both exon-array (Figure 6C) and qPCR experiments (Figure 6E) with primers in exon 23 indicate that transcripts containing exon 23 are very unstable in proliferating cells. Moreover, analysis of the nuclear fraction of the mRNA shows that the splice variant that includes exon 23 is made in proliferating cells and is more abundant in the nucleus than in the cytoplasm (Figure 6F and G), and therefore preferentially degraded in the cytoplasm of the proliferating cells. This example shows that the stability of splice variants is actively controlled and dependent on the differentiation status.

DISCUSSION

We reported around 7000 mRNA half lives in proliferating and differentiated myoblast cells. Given the 480 min time interval analyzed, the estimates for transcripts with half lives shorter than 480 min are more precise than those for very stable transcripts. The reported half lives can guide the choice of evaluation time points in future studies where transcript levels are induced or repressed [with the obvious exception of small interfering RNA (siRNA) experiments influencing mRNA degradation itself]. Knowledge on mRNA half life can be used in the analysis of gene expression profiling studies to identify genes that are in fact completely shut down, whereas the data as such only indicate that expression is slightly reduced. In addition, the data can be used to flag transcripts or transcript isoforms that cannot be valued since their mRNA half lives exceed the time points analyzed, or demand special attention due to significantly altered half lives.

Furthermore, we show that transcripts, including specific splice variants, with a role in myogenesis and differential expression between proliferating and differentiated myoblasts frequently demonstrate different stabilities in the two states. Transcripts with higher expression in differentiated than in proliferating cells are more rapidly degraded in proliferating cells and vice versa. A cell is in some cases reminiscent of a Dutch auction. When the clock is down to zero and there appears to be no demand, the commodities are to be destroyed.

Despite the characterization of a multitude of proteins and pathways involved in mRNA degradation (9), we are only beginning to understand the mechanisms that control the stability and degradation of individual transcripts. The 3’-UTR is very important for transcript stability given the presence of miRNA target sites, AU-rich elements (AREs) and binding sites for other RNA-binding proteins. Our observation that decay constants of first or last probe sets in a gene frequently differ (are higher or lower) from the other probe sets is consistent with the reported extensive use of differential 3’-UTRs (and alternative promoters) depending on the tissue or cellular state (10,25–27). In differentiating T-cells (10) and in cancer cells (27), differential expression of transcripts has been related to alternative 3’-UTR (polyadenylation site) use, resulting in differences in mRNA stability.

It is not yet clear which RNA binding proteins are responsible for the control of mRNA stability during myogenic differentiation. The HuR and KSRP can transiently affect the stability of transcripts containing AREs...
during myogenic differentiation (17–19, 28). However, only a minority of the transcripts with variable decay rates contain AU-rich elements [ARE motifs, predicted by the ARED algorithm (29)] in their 3'-UTR. From our digital gene expression studies (20), two members of the tristetraprolin family of RNA binding protein emerge as other important candidate factors for differentiation state-dependent control of mRNA degradation: ZFP36L1, which has been shown to accelerate degradation of tumor necrosis factor-alpha (TNF-α), vascular endothelial growth factor (VEGF) and granulocyte macrophage-colony stimulating factor (GM-CSF) (30–32) and ZFP36L2. The former is 10-fold increased and 3-fold decreased in expression during myogenic differentiation. The muscle-specific RNA binding protein CUGBP has been shown to stabilize mRNAs by binding to GU-rich elements in the 3'-UTR (8, 33). However, CUGBP shows little variation in expression between proliferating and differentiated cells and is therefore less likely to be responsible for the differences in decay rates observed in proliferating and differentiated myoblasts. In addition to RNA stabilizing and destabilizing proteins, also miRNAs and components of the nonsense-mediated decay machinery have been shown to have tissue-dependent expression levels and activities and may contribute to the observed differences in mRNA stability (34–38).

One of the new aspects of the current study is the notion that also splice variants that share the same 3'-UTR but include alternative internal exons may differ in stability. It was previously thought that the splicing machinery determines the abundance of the different splice isoforms aided by tissue-specific splicing factors like FOX, CELF, PTB, MBNL, hnRNP A/B (26, 39, 40). From our study, it appears that the presence of stabilizing or destabilizing elements in the alternative exons or differences in structural characteristics of the synthesized transcripts may also contribute to their differential expression. We show this in detail for the Itga7 transcripts variants with or without exon 23. The transcript including exon 23 appears to be more stable in differentiated cells, whereas in proliferating cells it is subject to fast degradation. A comparison of the nuclear fraction of the RNA with total RNA (nuclear and cytoplasmic) shows that the degradation of the transcript containing exon 23 mainly occurs in the cytoplasm. This observation is in line with a recent publication showing major differences between transcript variants in their nucleocytoplasmic distribution, at least partly attributable to mechanistic differences between nuclear and cytoplasmic mRNA degradation (41). We investigated whether exon 23 contains any known destabilizing elements such as AREs or PUFs (7), but could not find any. It is still possible that exon 23 contains an mRNA binding site but prediction programs failed to predict a good candidate. Thus, the mechanism for its preferential cytoplasmic degradation in proliferating myoblast remains elusive.

We conclude that mRNA degradation is a key controlling factor in gene expression regulation, possibly as important as transcription factor-mediated induction of mRNA synthesis. The mechanisms of mRNA degradation are currently less well understood than the mechanisms behind transcription factor activity regulation. We have gained several mechanistic insights in the process of mRNA degradation, but many mechanistic aspects deserve more attention in future studies. Knowledge on mRNA degradation rates, as presented in this study, is important for the proper interpretation of many biological studies. Decay rates need to be taken into account when choosing a time point for evaluation of the perturbation of a biological system, aid the modeling of transcriptional networks, and should be incorporated as important parameters in systems biology approaches.

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**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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