Quantification of long non-coding RNAs using qRT-PCR: comparison of different cDNA synthesis methods and RNA stability

Tomasz Kolenda¹,²,³, Marcel Ryś¹,², Kacper Guglas¹,²,³, Anna Teresiak¹, Renata Bliźniak², Jacek Mackiewicz⁴,⁵,⁶, Katarzyna Lamperska²

¹Chair of Medical Biotechnology, Poznan University of Medical Sciences, Poznan, Poland
²Laboratory of Cancer Genetics, Greater Poland Cancer Centre, Poznan, Poland
³Postgraduate School of Molecular Medicine, Medical University of Warsaw, Warsaw, Poland
⁴Department of Medical and Experimental Oncology, Heliodor Swiecicki Clinical Hospital, Poznan University of Medical Sciences, Poznan, Poland
⁵Department of Biology and Environmental Sciences, Poznan University of Medical Sciences, Poznan, Poland
⁶Department of Diagnostics and Cancer Immunology, Greater Poland Cancer Centre, Poznan, Poland

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Abstract

Introduction: Long non-coding RNAs (lncRNAs), a class of regulatory RNA molecules, are over 200 nucleotides long and could be used as a new potential biomarker, but their detection methods such as qRT-PCR are still not validated, and the influence of RNA degradation on lncRNA quantification is not clear. In this study, commercially available cDNA synthesis kits were tested and the influence of RNA degradation was compared.

Material and methods: Total RNA from FaDu cells was isolated and high quality RNA and highly degraded RNA samples were used. Reverse transcription was performed using three different commercially available kits and quantifications were performed using lncRNA Primer Plate and SYBR Green I Master by LightCycler 96. qRT-PCR was performed using three different cDNA samples and results are presented as the mean Ct values. A p-value < 0.05 was considered to be significant.

Results: Lower lncRNA Ct values (61/90; 67.78%) after qRT-PCR quantification were observed for cDNA synthesized using random hexamer primers preceded by polyA-tailing and adaptor-anchoring steps. It was observed that 9/90 (10.00%) lncRNAs were not detectable using different cDNA synthesis methods. For 75/90 (83%) lncRNAs, RNA degradation weakly influenced lncRNA Ct values and no differences were observed between high quality RNA and degraded samples. Seventy percent of examined lncRNAs showed significantly different Ct values depending on RNA degradation.

Conclusions: cDNA synthesis kits with random hexamer primers preceded by polyA-tailing and adaptor-anchoring steps allows enhancement of lncRNA quantification specificity and sensitivity. In most cases degradation of RNA samples does not affect lncRNA quantification because these molecules have good stability.

Key words: lncRNA, cDNA synthesis, qRT-PCR, RNA stability and degradation.
Introduction

The idea of using RNA as a biomarker is not new [1, 2]. Long non-coding RNAs (lncRNAs) are molecules longer than 200 nucleotides. They are actively transcribed and do not encode any type of proteins. Knowledge about lncRNAs is still limited and thus this is an important area for further research. lncRNA molecules possess many functional domains such as RNA or DNA binding sites and protein binding sites. They may also undergo a conformational switch. Because of these domains, lncRNAs may have crucial physiological functions, such as controlling transcription, post-transcription processes, or translation, or modeling epigenetic modifications. lncRNAs participate in cellular processes such as proliferation, apoptosis, response to stress, and regulation of cell metabolism or phenotype. Disturbances in lncRNA expression are associated with cancer processes, and a growing number of studies have focused on lncRNAs as new biomarkers [3].

Detection of expression of lncRNA originating from tissue, urine, peripheral blood, serum, saliva, or urine samples can be easily performed using molecular biology methods [4–9]. It was also reported that lncRNAs can be extracted from exosomes, which are thought to be mediators in cell-cell communication [10]. However, not all lncRNAs are present in each type of biological material. For example, Tang et al. observed that HOTAIR, HULC, MALAT1, MEG-3, NEAT-1, and UCA1 were present in cancer and adjacent noncancerous samples from OSCC (oral squamous cell carcinoma) patients, but only HOTAIR and MALAT1 from this group of lncRNAs were detected in some saliva samples [8].

There are many available methods to study lncRNAs: i) lncRNA immunoprecipitation; ii) lncRNA in situ hybridization; iii) Au-NP assay (gold nanoparticle-based); iv) lncRNA northern blot analysis; v) estimation of methylation status using HRM analysis (high resolution melting); vi) microarray or RNA sequencing; and vii) qRT-PCR or newly developed ddPCR (droplet digital PCR) [5, 6, 11–13]. The most frequently used method in lncRNA studies is a hybridization assay, especially the qRT-PCR method based on SYBR-green dye and TaqMan probes [14]. The qRT-PCR experiments can be laboratory-specific. Various research groups use different primers, reference genes (or set of genes) and amplification strategies, which sometimes makes it impossible to compare the results [15, 16]. It seems that only commercial qRT-PCR platforms could help to solve this problem. The commercially available qRT-PCR lncRNA platforms, such as the LncProfiler qPCR Array Kit (SBI), provide the possibility of simple and quick quantification of 90 lncRNAs in a single run, based on Ct (threshold cycle) analysis. In contrast to qRT-PCR, the microarray method is well-validated technology and is more comparable among experiments, but it is more expensive [17]. Moreover, using a commercial lncRNA microarray platform, expression of over 30,000 lncRNAs can be evaluated without using the sophisticated bioinformatics methods required for NGS (next generation sequencing) data extraction [18, 19]. However, in qRT-PCR and microarray methods, detection is limited to only known lncRNA transcripts.

As mentioned above, one of the most popular methods used in lncRNA research is qRT-PCR, but this method is not standardized and different approaches to prepare cDNA are used. Moreover, the influence of RNA degradation on quantification of lncRNA is not clear. These problems should be solved before using lncRNAs as a new class of biomarker in clinical practice.

In this study, different cDNA synthesis kits were evaluated and the quality of RNA samples was compared. Kits were based on the following: i) random hexamer primers preceded by polyA-tailing and adaptor-anchoring steps; ii) simple reaction using a blend of random hexamer primers and oligo(dT); iii) only random hexamer primers; and iv) only oligo(dT) (Figures 1 A–D).

Material and methods

The FaDu cell line, which is a model of hypopharynx squamous cell carcinoma, was used. The FaDu cell line was maintained in DMEM high glucose (4.5 g/l) medium (Sigma-Aldrich) supplemented with 8.85% (v/v) fetal bovine serum (Sigma-Aldrich), 1.77 mM L-glutamine (PAA), 0.885% (v/v) MEM non-essential amino acid solution (PAA), 0.885% (v/v) penicillin-streptomycin (PAA), and 8.85 mM HEPES (Sigma-Aldrich). Mycoplasma detection tests were performed routinely using the VenorGeM Mycoplasma PCR Detection Kit (Minerva Biolabs).

RNA isolation, quantification, and degradation protocol

Total RNA from FaDu cells was isolated using a High Pure miRNA isolation kit (Roche), according to the isolation protocol for total RNA (including the lncRNA fraction) from tissue and cell line samples. Quality and quantity of RNA samples were assessed using a NanoDrop spectrophotometer (Thermo Scientific), followed by 28S and 18S RNA band estimation (native 1% agarose gel electrophoresis in TAE buffer).

One high quality RNA sample was aliquoted (1 μg of total RNA) and used for the reverse transcription reaction using three different cDNA synthesis kits. The best cDNA kit was then chosen for further experiments.
To test the influence of RNA integrity on lncRNA amplification, an RNA degradation protocol was developed. Aliquoted RNA (1 µg/tube) was incubated for 0, 3, 6, 8, and 10 days at room temperature. At all time points, the RNA quality was tested using 28S and 18S rRNA band estimation (native 1% agarose gel electrophoresis in TAE buffer). Additional RNA samples on day 0 and day 10 were tested using a NanoDrop spectrophotometer. High quality (visible 28S and 18S rRNA bands), degraded (lack of 28S rRNA band and visible 18S rRNA band), and highly degraded RNA samples (lack of a 28S or 18S rRNA band, visible smear of RNA) were used for cDNA synthesis and the qRT-PCR reaction was performed.

**cDNA synthesis**

Reverse transcription was performed using three different commercially available kits: i) LncProfiler qPCR Array Kit (SBI); ii) iScript cDNA Synthesis Kit (Bio-Rad); and iii) First Strand cDNA Synthesis Kit (Fermentas). For all reactions, the same amount of total RNA (1 µg/reaction) from the same isolation was used. All experiments (RNA isolation, cDNA synthesis and qRT-PCR) were performed in triplicate.

**LncProfiler qPCR array kit (SBI)**

Reverse transcription is based on 3 steps: i) poly-A tailing; ii) annealing anchor dT adaptor; and

![Diagram of cDNA synthesis using different kits](image)

**Figure 1.** Comparison of different reverse transcription kits used to quantify lncRNA employing qRT-PCR methods. A – cDNA synthesis using random hexamer primers preceded by polyA-tailing and adaptor-anchoring steps, B – simple reaction using blend of random hexamer primers and oligo(dT), C – reaction based on random hexamer primers, D – based only on oligo(dT)
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Collecting drops by brief centrifugation. Next, 4 µl of 5× reaction buffer, 1 µl of RiboLock Ribonuclease Inhibitor, 2 µl of 10 mM dNTP mix, and 2 µl of M-MuLV reverse transcriptase were added. Reactions were incubated at 37°C for 60 min followed by 10 min at 70°C (for reactions with hexamer primers, the incubation procedure was followed by a sequential incubation step of 10 min at 25°C).

qRT-PCR reaction

cDNA was mixed with 1.750 ml of 2× LightCycler 480 SYBR Green I Master buffer (Roche) and 1.480 ml of nuclease free water, and 26 µl of the mixture was dropped into wells on a 96-well qRT-PCR plate. Next, 4 µl of IncRNA primers from Primer Plate (component of the LncProfiler qPCR Array Kit) was loaded onto the plate and the qPCR reaction was performed using the following protocol: preincubations (50°C for 2 min and 95°C for 10 min), 60 cycles of 2-step amplification (95°C for 15 s and 60°C for 1 min), and a melting step. Reactions were performed using a LightCycler 96 (Roche).

Statistical analysis

To compare Ct values, the statistical analysis was performed using GraphPad Prism 5 software with an unpaired t-test or a one-way analysis of variance (ANOVA) followed by Tukey’s post-hoc test. All qRT-PCR experiments were performed three times using three different cDNA samples and are presented as the mean Ct values. Error bars represent the standard deviation (SD) and a p-value < 0.05 was considered to be significant.

Results

In this study, different cDNA synthesis kits based on: i) random hexamer primers preceded by polyA-tailing and adaptor-anchoring steps; ii) simple reaction using a blend of random hexamer primers and oligo(dT); iii) using only random hexamer primers; and iv) only oligo(dT) were tested. We assumed that the lower the Ct value was, the more sensitive and precise was the method of cDNA synthesis. Comparison of mean qRT-PCR Ct values is presented on a heat map (Figure 2 A). Lower Ct values were observed for 61/90 (67.78%) of examined IncRNAs where cDNA was synthesized using random hexamer primers preceded by polyA-tailing and adaptor-anchoring steps. However, in the case of 15/90 (16.67%) IncRNAs, higher Ct values were observed for other methods (Figures 2 A and B).

Heat map and clustering showed that qRT-PCR Ct values for cDNA obtained using random hexamer primers preceded by polyA-tailing and adaptor-anchoring steps are distinct from other examined cDNA synthesis kits, which are similar to each other and clustered together (Figure 2 A). Moreover, 9/90 (10%) of the examined IncRNAs were detected only in the cDNA samples obtained by random hexamer primers preceded by polyA-tailing and adaptor-anchoring steps. However, for 5/90 (5.56%) IncRNAs, no IncRNA amplification was observed, and these IncRNAs were detected in cDNA samples synthesized using a combination of random hexamer primers and oligo(dT), and with only random hexamer primers or only oligo(dT) (Figures 2 A and B).

רו For the most frequently described IncRNAs in cancers and in head and neck squamous cell carcinomas in particular [3], significant changes were observed among qRT-PCR Ct values obtained using different cDNA synthesis kits. For H19 (p < 0.0001), HOTTIP (p = 0.0011), the GAS5 family (p < 0.0001), the MEG3 family (p = 0.0001), Air (p < 0.0001), and HOTAIRM1 (p < 0.0001), lower Ct values were observed (32.55 ± 3.12 vs. 42.49 ± 3.52), while for ANRIL (p = 0.0045) and HOTAIR (p = 0.0053), higher Ct values (34.18 ± 0.33 vs. 31.68 ± 1.46) were observed for cDNA obtained with random hexamer primers preceded by polyA-tailing and adaptor-anchoring steps compared to the rest of the tested kits (Figure 2 C).

The influence of total RNA degradation on qRT-PCR quantification was then investigated. Freshly isolated RNA was aliquoted (V = 5 µl/sample, C = 1 µg/sample) and incubated at room temperature for 3, 6, 8, and 10 days. Degradation of total RNA was observed (Figure 3 A). After 3 days of incubation, no 285 band but a smear of the 18S rRNA band was observed until day 10, when RNA was highly degraded (no 285 or 185 rRNA bands; Figure 3 A). The degradation process caused changes in RNA sample absorbance after 10 days of incubation that was measured using a NanoDrop spectrophotometer. The UV absorbance increased and is observed as an effect of apparent concentration change from 206.3 ng/µl measured immediately after isolation to 235.27 ng/µl measured after 10 days of incubation (p = 0.0227; Figure 3 B).

cDNA synthesis using random hexamer primers preceded by polyA-tailing and adapter-anchoring steps and qRT-PCR using the LncProfiler qPCR Array kit were performed for RNA samples from the day of isolation as well as after incubation for 3 and 10 days. For 83% (75/90) of IncRNAs, there were no significant differences (p > 0.05) in Ct values between high quality RNA and degraded RNA (day 3 or 10). For IncRNAs: Jpx, PSF inhibiting RNA, PTENP1, SAf, SNHG5, and the mRNA GAPDH there were no differences between Ct values of cDNA synthesized using RNA samples after isolation or after incubation for 3 days, but significantly higher Ct values were observed for the samples when the qRT-PCR was performed based on RNA incu-
Figure 2. Comparison of mean Ct values of qRT-PCR reactions based on cDNA synthetized using different commercially available cDNA synthesis kits. 
A – Heat map and clustering of mean Ct values of qRT-PCR reactions. Blue color indicates lower Ct values, red means higher, and grey means lack of amplification. B – Comparison of cDNA synthesis using random hexamer primers preceded by polyA-tailing and adaptor-anchoring steps with the rest of tested cDNA synthesis kits. C – Examples of the most changed Ct values

Data are presented as the mean Ct values; error bars represent SD; *p < 0.05, **p < 0.005, ***p < 0.0001.
bated for 10 days (34.92 ±0.28 vs. 37.71 ±1.17, p = 0.0225; 36.51 ±0.05 vs. 38.78 ±0.40, p = 0.0035; 37.72 ±0.72 vs. 41.28 ±1.75, p = 0.0221; 32.90 ±0.21 vs. 35.01 ±0.87, p = 0.0291; 29.55 ±0.58 vs. 33.32 ±0.58, p = 0.0018, and 25.10 ±1.10 vs. 27.76 ±0.81, p = 0.0150, respectively).

Moreover, these differences were also observed between samples incubated for 3 and 10 days (35.25 ±0.62 vs. 37.71 ±1.17; 36.98 ±0.54 vs. 38.78 ±0.40; 37.48 ±0.50 vs. 41.28 ±1.75; 32.82 ±0.53 vs. 35.01 ±0.87; 30.15 ±0.67 vs. 33.32 ±0.58, and 25.05 ±0.53 vs. 27.76 ±0.81, respectively). For the lncRNAs Tmevpg1 and Xist higher Ct values were noted only between days 3 and 10 (32.52 ±0.52 vs. 33.77 ±0.41, p = 0.0386, and 35.34 ±0.35 vs. 36.97 ±0.82, p = 0.0306, respectively). qRT-PCR performed using RNA incubated for 10 days provided better results (lower Ct values) compared to cDNA based on RNA after isolation in the case of NEAT1 (family), Nespas, NRON and ST7OT (27.45 ±0.50 vs. 28.91 ±0.02, p = 0.0490; 30.97 ±1.18 vs. 34.36 ±0.66, p = 0.0217; 32.87 ±0.36 vs. 34.01 ±0.13, p = 0.0263, and 28.04 ±0.41 vs. 29.74 ±0.36, p = 0.0214, respectively), and for lncRNA-RoR compared to cDNA based on RNA incubated for 3 days (34.27 ±0.31 vs. 35.36 ±0.23, p = 0.0238). For the lncRNAs MER11C and NTT lower Ct values were observed for RNA incubated for 10 days as well as for 10 days compared to RNA after isolation (30.51 ±0.48 and 30.31 ±0.60 vs. 32.15 ±0.01, p = 0.0257, and 32.90 ±0.49 and 32.72 ±0.45 vs. 34.97 ±0.23, p = 0.0047, respectively; Figure 4).

Discussion

Similarly to most researchers, we also chose the qRT-PCR method to study lncRNA expression because of its simplicity and quick sample analysis [20]. However, the main issue for qRT-PCR is selection of the proper methodology in the experimental workflow, but there are no experimental data comparing various approaches to this issue. In this study, we compared three different commercially available cDNA synthesis kits. Based on total RNA (including the lncRNA fraction) we obtained the cDNA, and this could be used to quantify expression of lncRNA using the commercially available LncProfiler qPCR Array Kit (SBI). We assumed that the lower the Ct value was, the more sensitive and precise was the method. Comparing three different approaches, we observed the best results (low Ct values) for the method using cDNA synthesis by random hexamer primers preceded by polyA-tailing and adaptor-anchoring steps. Generally, results of simple reactions using a blend of random hexamer primers and oligo(dT) or just random hexamer primers or oligo(dT) alone could incorrectly suggest that there was no expression of some lncRNAs or that the expression was at a very low level. It should be noted that some of the lncRNAs possess endogenous polyA tails but others do not. Moreover, most lncRNAs are present in samples at low copy numbers, which causes difficulties for their quantification [21, 22]. These two lncRNA features require the use of cDNA kits with an additional step of adding polyA tails and annealing anchor (dT) adapters. This approach allows enhancement of the specificity and sensitivity of lncRNA quantification, which we observed. Unfortunately, in most of the studies, cDNA is prepared using kits containing a mixture of oligo(dT) and/or random hexamer primers [14, 23, 24], where oligo(dT) is not functional for lncRNA molecules without a polyA tail. It is well known that specific primers decrease background priming, but random and oligo(dT) primers maximize the number of mRNA molecules that can be analyzed from a small sample of RNA [25]. We observed that artificially adding polyA tails and then

![Figure 3](image-url)
Figure 4. Comparison of mean $Ct$ values of changed lncRNAs and GAPDH mRNA amplified with cDNA synthesis using random hexamer primers preceded by polyA-tailing and adaptor-anchoring steps and RNA samples just after isolation and after 3 and 10 days incubation at room temperature.

Data are presented as the mean $Ct$ values; error bars represent SD; *$p < 0.05$, **$p < 0.005$. 
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Figure 4. Cont.
using annealing anchor (dT) adapters (analog of oligo(dT) primers) can help to quantify lncRNAs more precisely.

One possible explanation of the difficulties in quantification of some lncRNAs observed by us is an influence of molecular structure features such as length, GC content and molecule folding. It is well known that short RNA molecules such as miRNAs are difficult to quantify by qRT-PCR and require specific cDNA synthesis kits with adapter or polyA tail addition. Longer molecules seem to be easier to quantify than short RNAs such as miRNAs. On the other hand, a lot of secondary structures such as hairpin loops are problematic for cDNA synthesis and require a higher temperature for relaxing RNA molecules, making them available for reverse transcriptase [26, 27]. Moreover, one of the kits used by us possesses the DTT (dithiothreitol) enhancer, which helps to amplify templates rich in GC sequences, and it contributed to the better qRT-PCR results [28]. Because of the structural characteristics, some lncRNAs seem to need addition of a polyA tail or a higher temperature for a longer time during cDNA synthesis. We suggest that some of the results could be overlooked because an improper cDNA synthesis approach was used and specific lncRNA was not quantified.

The next problem is the stability of lncRNAs and the influence of RNA integrity on qRT-PCR quantification. Because of the length of lncRNAs (more than 200 nt), they are considered to be less stable and easier to degrade compared with short RNA molecules such as miRNAs. We analyzed the influence of RNA stability on quantification of lncRNAs using a cDNA kit (polyA tailing with anchor (dT) adapter annealing and cDNA synthesis using random primer mix), and we compared samples of high quality total RNA (visible 28S and 18S rRNA bands), degraded (no 28S band and visible smear of 18S rRNA band) and highly degraded RNA (no 28S or 18S rRNA bands). Degradation causes changes in RNA structure increasing the amount of short RNA molecules, which strongly affects RNA detection [29, 30]. Our results indicated that most of the examined lncRNAs are stable, even when the total RNA is highly degraded. Our observations are supported by the results obtained by Kraus et al. They reported that some lncRNAs are more stable than miRNAs that are about 22 nucleotides long [31, 32].

However, some long transcripts, such as mRNAs, are sensitive to degradation, and in this case, RNA integrity is an important factor affecting qRT-PCR quantification. It is postulated that the lncRNA half-life depends on its coding place in the genome, on posttranscriptional modifications, and on subcellular localization and its function [33]. Some authors have noted that localization of lncRNAs in the genome may influence their transcript stability, especially intragenic and cis-antisense lncRNAs compared with those derived from introns (half-life more than 16 h) [8, 33]. We noted that some lncRNAs are stable for much longer. After 3 days, we did not observe changes in Ct values in most of the examined lncRNAs, while after 10 days, changes in Ct values were observed only in some lncRNAs. Detection of some lncRNAs in saliva confirms the high stability of these molecules [8]. Moreover, lncRNAs isolated from the plasma are resistant to RNase A digestion and overnight incubation at room temperature [9]. Generally, low RNA stability may cause difficulties in analyzing some long coding or non-coding RNA transcripts obtained from archived formalin-fixed paraffin-embedded (FFPE) blocks [34]. This problem is solved by applying the simple modification of measuring the lncRNA expression level by reaction with three different pairs of non-overlapping primers [35]. However, RNA is both degraded and modified under the archiving process in FFPE blocks, which makes this RNA more difficult to analyze [36].

It is well known that the main features of an ideal biomarker molecule are simplicity of acquisition, diversity of sources where it occurs, and ease of measurement methodology [20, 37]. We postulate that under appropriate processing conditions, lncRNA seems to meet the characteristics of biomarker molecules. However, standardization procedures should be applied before clinical use of lncRNA biomarkers.

In conclusion, we recommend the following: i) use of cDNA kits designed for lncRNA or cDNA synthesis preceded by polyA tailing with anchor (dT) adapter annealing; ii) use of highly thermostable reverse transcriptase designed for long reactions at higher temperatures; and iii) use of high RNA quality is generally recommended, but most lncRNAs seem to be stable and can be quantified in degraded RNA samples.

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Conflict of interest

The authors declare no conflict of interest.
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