METHOD

Evaluation of critical design parameters for RT-qPCR-based analysis of multiple dUTPase isoform genes in mice

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The coupling of nucleotide biosynthesis and genome integrity plays an important role in ensuring faithful maintenance and transmission of genetic information. The enzyme dUTPase is a prime example of such coupling, as it generates dUMP for thymidylate biosynthesis and removes dUTP for synthesis of uracil-free DNA. Despite its significant role, the expression patterns of dUTPase isoforms in animals have not yet been described. Here, we developed a detailed optimization procedure for RT-qPCR-based isoform-specific analysis of dUTPase expression levels in various organs of adult mice. Primer design, optimal annealing temperature, and primer concentrations were specified for both nuclear and mitochondrial dUTPase isoforms, as well as two commonly used reference genes, GAPDH and PPIA. The linear range of the RNA concentration for the reverse transcription reaction was determined. The PCR efficiencies were calculated using serial dilutions of cDNA. Our data indicate that organs involved in lymphocyte production, as well as reproductive organs, are characterized by high levels of expression of the nuclear dUTPase isoform. On the other hand, we observed that expression of the mitochondrial dUTPase isoform is considerably increased in heart, kidney, and ovary. Despite the differences in expression levels among the various organs, we also found that the mitochondrial dUTPase isoform shows a much more uniform expression pattern as compared to the reference genes GAPDH and PPIA.

The dUTPase enzyme family is essential in maintaining genome integrity [1]. It has been also found to be almost ubiquitous among free-living organisms from bacteria to man and is encoded in diverse evolutionary strains of herpes viruses and retroviruses as well [2]. The function of the enzyme is to remove dUTP from the cellular pool via its enzymatic action to catalyze pyrophosphorolysis of dUTP to result in dUMP and pyrophosphate products. This reaction has dual importance for cellular physiology, since on the one hand it sanitizes the nucleotide pool, while on the other hand it also produces dUMP, the key metabolite for de novo thymidylate biosynthesis. The significance of low dUTP level within the cells can be appreciated in light of the cellular response to increased concentration of uracil in DNA. Uracil can appear in DNA by two routes, either via cytosine deamination...
(that may occur spontaneously or through regulated cytosine deaminases) or thymine-replacing incorporation under high cellular dUTP level. Cytosine deamination is a mutagenic reaction that will lead to an exchange of a G:U base pair to an A:T base pair if not promptly repaired. Thymine-replacing uracils are not mutagenic per se; however they may perturb those DNA-protein interactions where the methyl group on the thymine base plays an important role. Both uracils from deaminated cytosines or thymine-replacing incorporation are targeted by uracil excision repair (base excision repair, BER). If dUTP levels are high in the cellular milieu, repair synthesis will reintroduce uracils, creating a hyperactive futile cycle leading to chromosome fragmentation and cell death (thymine-less cell death). Correct and efficient action of dUTPase is therefore of crucial importance to prevent thymine-less cell death [3,4].

It has been well established that dUTPase action is of most significance in cells with actively ongoing DNA synthesis. In eukaryotic organisms, dUTPase is usually present in two isoforms, one of these is nuclear and the other is mitochondrial/cytoplasmic [5–8]. In agreement with its physiological role, in in vitro cellular experiments the expression of the nuclear isoform of dUTPase was shown to be under cell cycle control. However, it is also known that in human cells the mitochondrial isoform is constitutively expressed. Although the expression pattern of dUTPase and its regulation in isolated eukaryotic cell lines has been covered in insightful details in numerous articles, much less is known about the expression pattern of dUTPase in living organisms. There is especially a lack of information regarding the expression pattern of the two distinct isoforms, since expression databases provide only a combined value. In plants, dUTPase level was associated with the mitotic status of meristem tissues [9]. In Drosophila melanogaster, the expression of dUTPase was shown to be under developmental control [5,10]. In cancer tissues, overexpression of the nuclear isoform of dUTPase was associated with poor prognosis [11–13].

While there is a lack of detailed focused investigations, the Expression Atlas database obtained by large-scale systems biology approaches provide a wealth of data on dUTPase expression levels [14–18]. The reliability of such data is limited, and conflicting results can be found. Moreover, none of the data provide any specific information for both of the isoforms of dUTPases.

In light of the physiological importance of dUTPase and lack of data on its isoform-specific expression level, our aim was to develop a highly reliable and well-controlled reverse transcription/quantitative real-time polymerase chain reaction (RT-qPCR) method that can be used for parallel determination of the expression levels of both dUTPase isoforms. We selected mouse as an animal model for these studies. In mice, the two isoforms are generated via alternative splicing, thus for isoform-specific detection, amplification of the differing regions of the two transcripts was necessary. Both of these regions have high GC content and therefore a complex secondary structure that makes the optimization process rather difficult [19]. Despite the significance of RT-qPCR, there has been a lack of consensus on how to perform and interpret experiments using this method. This issue was addressed by the MIQE guidelines in order to provide more reliable results [20]. Here, we present a detailed optimization procedure for the RT-qPCR conditions and report the performance parameters of the optimized method in compliance with the MIQE guideline. We also use this protocol to characterize the organ-specific expression patterns of dUTPase isoforms in adult mice.

**Results and Discussion**

**Optimization of RT-qPCR conditions**

**Primer selection and annealing temperature optimization**

The Dut gene encodes for both nuclear and mitochondrial isoform of the enzyme dUTPase in mouse. These

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Fig. 1. Selection of primers. (A) On the top, genomic sequences of both nuclear and mitochondrial isoforms are shown. Exons are indicated with Roman numerals in rectangles and shown with alternating blue and pink colors; introns are simplified as lines (for longer introns, lines are broken). In the middle, schematic illustration of primer candidates for amplification of both isoforms—after reverse transcription—of the Dut transcript is depicted. Exons are numbered and shown with alternating blue and pink colors. Common part of the sequence where the reverse primers are located is enlarged. All primers are indicated with arrows. On the bottom, PCR products amplified with the Rev1 reverse and the appropriate forward primers for both isoforms are shown. (B) Amplification curves for the mitochondrial isoform with the Rev1 reverse primer performed at a range of annealing temperatures from 53.1 to 69 °C using a gradient thermal block. RFU, relative fluorescence unit. (C) Melting curve analysis of the PCR products of the amplification introduced in panel B. (D) Agarose gel electrophoresis of the PCR products of the amplification introduced in panel B. The specific product is indicated with an arrow. (E) Quantification cycles (Cq) at a range of annealing temperature from 53.1 to 69 °C with all four reverse primer candidates for the mitochondrial isoform. The solid squares indicate specific products as determined with agarose gel electrophoresis and melting curve analysis. Open squares indicate aspecific products. (F) The same representation for the nuclear isoform as shown in panel E.
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A

DNA

Mitochondrial

Nuclear

mRNA

Mitochondrial

Nuclear

PCR product

Mitochondrial

Nuclear

mDut-Fw

78 bp PCR product

nDut-Fw

53 bp PCR product

mDut Fw-Rev1

B

Amplification

Mitochondrial

mDut-Fw

Rev1

C

Melt Peak

Mitochondrial

mDut-Fw

Rev1

Rev2

Rev3

Rev4

EF

E

mDut

Rev1

Rev2

Rev3

Rev4

F

nDut

Rev1

Rev2

Rev3

Rev4

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isoforms differ only in the first exon via alternative splicing (Fig. 1A). For the quantification of these transcripts, forward primers were designed in the first different exons. As opposed to the limited possibility for the forward primer design, several reverse primers could be applicable. Since the amplicon length can remarkably influence the efficiency of PCR, with short segments characterized with more efficient amplification, we deserved to amplify short regions of cDNA [21]. The design of primers and the prediction of the specificity were carried out with the Primer-BLAST tool [22]. Unfortunately, in silico prediction of the specificity of the products with each potential primer pairs did not result in a conclusive optimal selection of adequate primers in terms of specificity. Therefore, we decided to select three exonic (Rev1, Rev2, and Rev3) and one exon junction spanning (Rev4) common reverse primers for both isoforms to be experimentally investigated (Table 1).

We realized that the relatively high GC content of the first exon of the nuclear and the first two exons of the mitochondrial isoforms (70% for the nuclear and 75% for the mitochondrial isoform, respectively) may be associated with a complex secondary structure and accordingly a decrease in PCR efficiency [19]. Hence, the secondary structure of the transcripts was analyzed with the OligoAnalyzer tool of Integrated DNA Technologies. The secondary structure of the PCR product with the isoform-specific forward primers and Rev4 and Rev1 reverse primers is shown in Fig. S1. We selected the Rev4 and Rev1 reverse primers for this comparison since these will lead to the production of the longest and shortest PCR products. Using the Rev4 reverse primer, a longer PCR product is formed which results in a more complex secondary structure characterized with a higher melting temperature and greater change in Gibbs free energy as compared to the PCR product produced with Rev1 reverse primer (Fig. S1E).

For both nuclear and mitochondrial isoforms, all four candidate reverse primers were tested at a range of annealing temperature from 53.1 to 69 °C using a gradient thermal block. To test all primer pairs, the same amount of cDNA template was used in the PCR in order that the quantification cycle (Cq) values of different primers can be compared for each target. Figure 1B shows the amplification curves for the mitochondrial isoform with the Rev1 reverse primer. Amplification associated with higher annealing temperature results in lower Cq value, indicating more efficient amplification. Melting curve analysis (Fig. 1C) and agarose gel electrophoresis (Fig. 1D) were performed to assess the specificity of the amplification reaction. In case of the PCR product for the mitochondrial isoform with the Rev1 reverse primer, the 78-base pair (bp) length specific product is formed in the range of annealing temperature from 66 to 69 °C with a melting temperature of 89.5 °C; however, at lower temperature aspecific products appear (cf aspecific products with approximately 84 °C melting temperature on Fig. 1C). For the nuclear isoform with the Rev1 reverse primer, the image of agarose gel electrophoresis is shown in Fig. S2A. Between 55.4 and 69 °C, 53-bp length specific product is formed. Figure S2B presents the agarose gel electrophoresis of the PCR products for both isoforms with Rev2 reverse primer in the same range of annealing temperature. For the nuclear isoform, the 73-bp length specific product is formed from 58.6 to 69 °C, while for the mitochondrial isoform the 98-bp specific product is formed only at 68 °C.

For comparison of the candidate reverse primers, the Cq values were plotted on the same diagram in the investigated temperature range (Fig. 1E,F for the mitochondrial and nuclear isoforms, respectively) also marking the specificity of the products. In case of the mitochondrial isoform, despite the specificity of the product with Rev4, the Cq values are much higher than in case of Rev1 and Rev2. Rev3 has a similar temperature range regarding the product specificity than Rev1 and Rev2; however, the Cq values are higher. Rev1 and Rev2 are quite similar to each other both in specificity and Cq values. In case of the nuclear isoform, all four reverse primers result in specific product formation from 62.7 to 69 °C. Rev1 and Rev2

### Table 1. Sequences and melting temperatures of the primers used in this study. nDut-Fw and mDut-Fw are used as the forward primer for the amplification of the nuclear and mitochondrial isoforms of Dut, respectively. Rev-1, Rev-2, Rev-3, and Rev-4 are four reverse primer candidates for the amplification of both isoforms. PPIA-Fw and PPIA-Rev are used for the amplification of PPIA, while GAPDH-Fw and GAPDH-Rev are used for the amplification of GAPDH.

| Primer name       | Sequence 5'-3'                  | \(T_m\) (°C) |
|-------------------|---------------------------------|-------------|
| nDut-Fw           | ATGCCCTGCTCGGAAGAT              | 64.8        |
| mDut-Fw           | CCGAGGAAGCGCAGGAG               | 66.3        |
| Rev1              | GCTGAGACCGCTTGGAG               | 65.3        |
| Rev2              | GCAGAGAACGCCACCTCT              | 65.8        |
| Rev3              | GTGCCGTCCTCCGAGA                | 65.4        |
| Rev4              | TGTTAATCATAGGCACTGAGTAGG        | 59.9        |
| PPIA-Fw           | TGGCTATAAGGGTTCCTCT            | 61.4        |
| PPIA-Rev          | AAAACTGGGAACGGTGTG              | 63.6        |
| GAPDH-Fw          | AGCTATTCTCTGGTGATGCAAT          | 62.5        |
| GAPDH-Rev         | GCCCCTCCTGTATATTGG             | 64.1        |
are similar to each other regarding the C_q values, while Rev3 and Rev4 have higher values; thus, the use of Rev3 and Rev4 was ruled out. The differences in C_q values produced with the four reverse primer candidates may be the consequence of different amplicon length and corresponding PCR efficiencies, as amplification of shorter regions associated with less complex secondary structure proved to be more advantageous regarding the C_q values.

Besides the two isoforms of the Dut gene, two commonly used reference genes were also included in this study, glyceraldehyde 3-phosphate dehydrogenase (GAPDH) and peptidyl-prolyl cis-trans isomerase A (PPIA) [23–27]. The primers used for amplification of GAPDH and PPIA cDNAs produce specific products (209 and 200 bp for GAPDH and PPIA, respectively) in the whole range of annealing temperature from 53.1 to 69 °C (Fig. 2A,B); nonetheless, C_q values start to increase above 66 °C indicating suboptimal reaction conditions (Fig. 2C). We aimed to develop a method where transcripts from both nuclear and mitochondrial isoforms of the Dut gene as well as the two references (GAPDH and PPIA) can be analyzed on the same plate (i.e. gene maximization method). Thus, for determination of a common nearly optimal annealing temperature for all four targets, a detailed temperature gradient experiment was performed in the range from 61.2 to 66°C (Fig. 2D, E and Fig. S2C–E). With Rev1 reverse primer, the PCR product for the nuclear isoform was specific in the whole temperature range. For the mitochondrial isoform, however, specific product formation was observed only above 65 °C, while at 65 °C only a minuscule amount of aspecific product was found (Fig. 2D). With Rev2 reverse primer, aspecific product formation was found also at 65.7 °C and below for the mitochondrial isoform (Fig. S2C). For comparison of the candidate reverse primers in the detailed annealing temperature range, refer to Fig. S2D,E.

In conclusion, due to better specificity in case of the mitochondrial isoform, Rev1 was selected for the reverse primer. Fig. 2E shows the C_q values and specificity of the selected primers in the detailed annealing temperature range. For the C_q values and specificity of the selected primers in the broader 53.1-69 °C annealing temperature range, refer to Fig. S2F. Both 65 and 66 °C could serve as a common annealing temperature for amplification of all four targets; however, at 66 °C, the C_q value for PPIA was increased by 1 cycle indicating suboptimal reaction. At both temperature points, several PCRs were performed to assess the product specificity (Fig. 2F,G). At 65 °C, some of the several PCR products were aspecific, while at 66 °C no aspecific product was found; thus, 66 °C was selected for the annealing temperature to be used.

**Optimization of primer concentration**

The importance of the optimization for primer concentrations is rarely taken into account; however, it can greatly influence the performance of the assay [28]. Five point 2-fold serial dilutions were prepared of each primer and a range of forward primer concentration was run against a range of reverse primer concentration in a form of a matrix for both nuclear and mitochondrial isoforms (Fig. 3) [29]. For all reactions, a master mix was prepared with a small amount of cDNA that only lacked the primers to ensure constant concentration of all components of the reaction—except for the primers. Two technical replicates were used for each point of the matrix. In case of the nuclear isoform (Fig. 3A), higher concentrations of both primers resulted in lower C_q values, while all products were specific; therefore, the highest values of the tested concentrations were chosen for further experiments, that is 1200 nM for the forward and 1000 nM for the reverse primer. In case of the mitochondrial isoform (Fig. 3B), C_q values also decreased with higher concentrations; nonetheless, specificity of the product was compromised at the two highest concentration points of the forward primer. To avoid aspecific product formation, the third most concentrated point for the forward and the most concentrated point for the reverse primer was selected for further experiments, that is 375 nM for the forward primer and 1000 nM for the reverse primer.

**Confirmation of the identity of the PCR products**

The GAPDH and PPIA products were sequenced; however, the products of the nuclear and mitochondrial dUTPase isoforms are too short for sequencing (53 and 78 bp, respectively); therefore, a two-round amplification was performed (Fig. 4). Since for both isoforms, the Rev4 reverse primer had appropriate product specificity and length for sequencing (165 bp for the nuclear and 190 bp for the mitochondrial isoform), the first round of PCR was performed using ovary cDNA, Rev4 reverse primer and the appropriate forward primer, and then, the PCR products were purified and sent for sequencing. As the sequences of the products were confirmed, they were diluted and used as template in the second round of PCR with Rev1 reverse primer and the appropriate forward primer for both isoforms, since the PCR products with Rev4 reverse primer contain the whole sequence of the
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**Fig. 2.** Determination of the optimal common annealing temperature with thermal gradient for the PCR amplification of both isoforms of the Dut transcript, GAPDH, and PPIA. (A, B) Agarose gel electrophoresis of the PCR products for GAPDH (A) and PPIA (B) at a range of annealing temperature from 53.1 to 69 °C. The specific product is indicated with an arrow. (C) Quantification cycles (Cq) at a range of annealing temperatures from 53.1 to 69 °C for GAPDH and PPIA. (D) Agarose gel electrophoresis of the PCR products for both nuclear and mitochondrial isoforms with Rev1 reverse primer at a range of annealing temperature from 61.2 to 66 °C. The specific products are indicated with arrows. (E) Quantification cycles (Cq) at a range of annealing temperatures from 61.2 to 66 °C with the selected primers. The solid squares indicate specific products as determined with agarose gel electrophoresis and melting curve analysis. Open squares indicate aspecific products. (F, G) Melting curve analysis of the PCR products for the mitochondrial isoform with the selected Rev1 reverse primer at 65 °C (F) and 66 °C (G) for the determination of product specificity.

PCR products with Rev1 reverse primer. The second round PCR products were run on agarose gel along with products of a single-round PCR using Rev1 reverse primer and appropriate forward primers for both isoforms and ovary cDNA (Fig. 4A). The melting curves of the second round and single-round PCR
products were also compared (Fig. 4B). Both comparisons confirmed the two products to be identical for both isoforms. Fig. 4A also shows representative bands of the PCR products for GAPDH and PPIA in the optimized PCR conditions.

**Selection of a reverse transcription kit**

Since the performance of the reverse transcription (RT) reaction heavily depends on the target concentration, total RNA concentration, priming strategy, the enzyme used and other parameters, careful selection and optimization of these criteria are of utmost importance [30,31]. First, we aimed to compare two commercially available RT kits. For this purpose, three different sources of RNA (heart, ovary, and spleen) were subjected to two different RT kits, High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems, Foster City, CA, USA) and iScript cDNA Synthesis Kit (Bio-Rad, Hercules, CA, USA). Four point 4-fold serial dilutions of RNA were prepared and introduced to the RT reaction with a starting amount of 800 ng (Fig. 5A–D). All cDNA samples were subjected to qPCR measurement with three technical replicates. For each serial dilution, weighted least-squares linear regression was performed to the average of the three technical replicate \( C_q \) values. In case of the nuclear isoform and GAPDH, using the Applied Biosystems kit, the most concentrated point of the spleen sample was excluded because of lack of fit (Fig. 5A). In case of the mitochondrial isoform, fitting was performed using only three concentration points as the most concentrated points showed lack of fit for all samples (Fig. 5B). All fittings for the Applied Biosystems kit were appropriate with \( R^2 \)-square values above 0.99; however, in some cases the \( R^2 \)-square values were as low as 0.91 for the Bio-Rad kit.

For both isoforms, the use of the High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems) resulted in considerably lower \( C_q \) values than iScript cDNA Synthesis Kit (Bio-Rad) for all three RNA samples (Fig. 5A,B), indicating better performance of the RT reaction. In addition, the slopes of the regression lines were much lower for the mitochondrial isoform using the Bio-Rad kit, indicating that the sensitivity of the assay could be compromised using the Bio-Rad kit. For PPIA, the Applied Biosystems kit performs somewhat better, while in case of GAPDH, with Bio-Rad kit lower \( C_q \) values could be achieved (Fig. 5C,D); however, both kits produced parallel linear lines. In conclusion, for both isoforms and PPIA, Applied Biosystems kit is more preferential and also
applicable for GAPDH; thus, it was selected for further experiments.

**Determination of the linear concentration range for the RT reaction**

To provide comparability of the results of a RT-qPCR study, the efficiency of the RT reaction must be kept constant. For this purpose, standardization of the reaction parameters—including total RNA concentration—and to work within the linear concentration range of the reaction is essential [30,31]. To determine the linear range of RNA concentration for the cDNA synthesis—thus ensuring lack of inhibition of the RT reaction in an extent that could compromise our results—five RNA samples from different organs (brain, heart, ovary, spleen, and thymus) were prepared (Fig. 5E–H). Four point 4-fold serial dilutions of the samples were subjected to cDNA synthesis followed by qPCR measurement with three technical replicates. For each serial dilution, weighted least-squares linear regression was performed to the average of the three technical replicate C_q values. In all cases, the most concentrated points fell out of the linear range for the mitochondrial isoform (Fig. 5F), whereas all concentration points were included in the linear range for the nuclear isoform.
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A. nDut

B. mDut

C. GAPDH

D. PPIA

E. nDut

F. mDut

G. GAPDH

H. PPIA

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(Fig. 5E), GAPDH (Fig. 5G), and PPIA (Fig. 5H). For further experiments, the second most concentrated point was selected; that is, 200 ng total RNA was subjected to cDNA synthesis to ensure linearity of the RT reaction for all four targets investigated.

**Determination of the PCR efficiencies**

In contrast to a commonly used method, we used serial dilutions of cDNA samples that were prepared with our verified RT reaction, instead of serial dilutions of standards, since the PCR efficiencies can be affected by components presented in the matrix. Thus, the use of serial dilutions of standards for the determination of PCR efficiencies inherently bears the possibility of the results being biased because of the different matrices [32].

Five point 4-fold serial dilutions were prepared from cDNA samples derived from five different organs (brain, heart, ovary, spleen, and thymus) and subjected to qPCR measurement with four technical replicates (Fig. 6). Weighted least-squares linear regression was performed for all data points to counteract the effects of heteroscedasticity [33]. However, in case of the two Dut isoforms, some replicates of the least concentrated point did not provide Cq values as amplification did not occur, in those cases the whole concentration point was excluded from the analysis (Fig. 6) [34]. The intercept and slope of the curves and R-square values were determined, and efficiency was calculated from the slope (Table S1) [35]. Mean efficiency was calculated as the arithmetic mean of the individual efficiency values. The efficiency was estimated to be 88.4% for the nuclear isoform, 79.5% for the mitochondrial isoform,
94.1% for GAPDH, and 91.7% for PPIA. Generally, PCR efficiencies above 90% are accepted due to the assumption that precise and sensitive reactions are associated with high PCR efficiency [28]. However, we demonstrate below that our assay is characterized with adequate performance parameters. For further experiments, the determined efficiencies were taken into account to calculate relative quantity values.

For all concentration points investigated, the reaction was linear. Thus, the dynamic range of the assay was linear for GAPDH between 20.3 and 35.5 Cq, for PPIA between 19.7 and 33.3, for the nuclear isoform between 23.5 and 36.8, for the mitochondrial isoform between 30.3 and 40.6 Cq values. Moreover, the cDNA added to the reaction in a final dilution from 0.5 µL to 1.95 × 10^{-4} µL per 10 µL reaction resulted in no observable inhibition of the amplification. For further experiments, 0.31 µL of cDNA was added to each reaction.

**Expression levels of dUTPase isoforms in adult mouse tissues determined by the optimized RT-qPCR protocol**

To assess the applicability of the optimized RT-qPCR assay, RNA was isolated from eight different organs of 10-week-old male and female mice. The integrity of all RNA samples was assessed with agarose gel electrophoresis (Fig. S3), and the purity was determined with NanoDrop (Table S2). Gel electrophoresis was our method of choice for the assessment of RNA integrity, because of its simplicity and affordability and considering that even moderately degraded RNA samples can be reliably quantified, provided the amplicons are short and expression is normalized to reference genes [36]. All samples visualized on agarose gel showed the two distinct bands that were identified as the characteristic rRNA bands (28S and 18S) without any obvious degradation products or genomic DNA contamination. The 260/280 ratios were between 1.98 and 2.13 in most cases with two slightly higher values (2.22 and 2.61) (Table S2); thus, the purity of RNA was accepted for all samples [37].

Samples were subjected to the assay with three biological replicates from three different mice and three technical replicates for each biological replicate for each biological group (Fig. 7). In this study, biological groups are defined as different organs from either female or male mice. Fig. 7A,B shows the amplification curves of the four targets (the two dUTPase isoforms, and the two reference genes, GAPDH and PPIA) for organs with the highest level of expression of the nuclear and mitochondrial isoform, male thymus, and male heart, respectively. Graphs showing the amplification curves for all biological groups are included in Fig. S4.

We have observed that both the biological and technical scatter of the Cq values for each target within each biological group is considerably low as shown by the amplification curves (Fig. 7A,B and Fig. S4). The standard deviation of the Cq values between technical replicates was found to be 0.14, 0.13, 0.065, and 0.13 for the nuclear and mitochondrial isoform, GAPDH, and PPIA, respectively, as determined by ANOVA. Comparing the Cq values obtained for the different organs, we have also observed that the difference between the Cq values associated with the target genes GAPDH and PPIA showed considerably larger alterations than the Cq values for the mitochondrial dUTPase isoform. Therefore, based on the low variation of the Cq values within every biological group, we decided to use the ΔCq method in Bio-Rad CFX MAESTRO software for a reliable assessment of the relative expression level of every biological group for all four targets (Fig. 7C–F). As no reference gene was used in this comparison and several qPCR runs were included, the threshold in the amplification plot was set to a common value to ensure comparability of the Cq values between qPCR runs. Although the use of the ΔCq method with no reference genes may lead to errors due to the variation of RNA content and sample loading, the differences in the efficiency of RT and PCR and RNA integrity [23,38], in our present experiments, the very low variation of the Cq values among the nine replicates indicated that such errors will not affect the reliability of our analysis. In fact, as shown in Fig. 7E, F, both reference genes showed considerably large variation among the different organs and the most stable expression pattern was shown by the mitochondrial dUTPase isoform (Fig. 7D).

Differences between the two sexes were small and found to be nonsignificant as determined by ANOVA (for the effect of sex, P values were 0.20, 0.94, 0.15, and 0.08 for the nuclear and mitochondrial isoform, GAPDH, and PPIA, respectively). However, larger differences were observed between organs that were highly significant (P < 10^{-16} for all four targets). In case of the nuclear isoform, thymus had the highest expression, followed by testicle, spleen, ovary, and brain, in order. Heart, kidney, liver, and lung showed markedly lower expression levels. The largest difference observed was between the expression levels of the nuclear dUTPase isoform in male liver and male thymus, with a 138-fold increase in the latter. The expression level of the mitochondrial isoform was highest in the heart, followed by kidney and ovary. In contrast...
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A

Amplification

male thymus

B

Amplification

male heart

C

nDut

Relative Quantity

10^0 10^1 10^2

brain heart kidney liver lung ovary/testicle spleen thymus

12.6 11.2 1.48 1.63 4.99 3.29 1.84 1.00 3.12 3.88 20.5

D

mDut

Relative Quantity

10^0 10^1 10^2

brain heart kidney liver lung ovary/testicle spleen thymus

1.23 1.23 4.52 4.61 3.03 2.46 1.22 1.00 1.10 1.38 2.67

E

GAPDH

Relative Quantity

10^0 10^1 10^2

brain heart kidney liver lung ovary/testicle spleen thymus

13.6 12.5 23.4 28.5 14.2 13.4 4.48 4.26 1.00 1.20 6.63 3.34 1.67 1.84 6.65 8.36

F

PPIA

Relative Quantity

10^0 10^1 10^2

brain heart kidney liver lung ovary/testicle spleen thymus

5.58 6.71 1.00 1.32 4.02 3.84 2.30 1.79 1.32 2.25 5.37 8.28 2.44 2.00 6.04 9.33
to the nuclear isoform, the mitochondrial isoform presented much smaller differences. An overall 4.61-fold difference between the highest (male heart) and lowest (male liver) expression level was observed. As for GAPDH and PPIA, considerable differences exist between the organs, an overall 28.5-fold difference in GAPDH expression and 9.33-fold difference in PPIA expression was detected in comparing the highest expression to the lowest expression. The mitochondrial dUTPase isoform shows considerably stable expression pattern among various tissues, thus may be regarded as a possible reference gene for gene expression analysis involving several different organs with similar expression level of the mitochondrial isoform. In our study, heart, kidney, and ovary showed elevated expression levels; hence, normalization with the mitochondrial isoform for the comparison of gene expression between all these organs is not applicable, however may be appropriate for organs with similar expression levels.

Conclusions

Our aim was to design, optimize, and validate a method for the simultaneous quantification of the expression levels of both nuclear and mitochondrial isoforms of dUTPase. These isoforms differ only in the first exon via alternative splicing. To develop an isoform-specific method, amplification near the 5′ end of the transcripts was necessary, where the target sequences have a relatively high GC content and accordingly, a complex secondary structure that makes the optimization process rather difficult. In this paper, we introduced a detailed optimization procedure that could be useful for overcoming similar problems. Amplification of shorter regions associated with less complex secondary structure proved to be more advantageous. Since sequencing of short amplicons is not possible with the conventional Sanger sequencing method, we propose a two-round amplification method for the confirmation of the identity of the PCR products. We found that higher primer concentration results in more efficient amplification as long as the product specificity is not compromised. We compared two commercially available reverse transcription kits for our purpose and realized that the selection of an RT kit determines the suitability of the RT-qPCR assay fundamentally, and its applicability should be investigated for each target gene. We have successfully developed a method for the isoform-specific determination of the expression of dUTPase taken into consideration the critical design parameters of the assay. In addition, we have evaluated the performance of the assay in terms of specificity, efficiency, precision, and the linear dynamic range. Our study is fully compliant with the MIQE guidelines (Table S3).

With regard to the expression of dUTPase isoforms among the different organs in adult mice, we observe that the nuclear isoform is expressed at a high level in thymus, spleen, and reproductive organs. It is well known that in both thymus and spleen, considerable mitosis occurs due to their involvement in lymphocyte production [39]. Therefore, the biological role of these organs is in agreement with the well-established role of dUTPase in dividing cells. Heart, liver, kidney, ovary, and brain are organs known to be rich in mitochondria [40,41]. However, elevated level of expression of the mitochondrial isoform of dUTPase was found only in heart, kidney, and ovary. Only a minuscule difference in the expression of the mitochondrial isoform was found in the other organs. In contrast, the difference in the expression levels of two commonly used reference genes, GAPDH and PPIA is much larger among the different mouse organs. This observation underlines the importance of the selection of appropriate reference genes in gene expression studies. Based on our results, we propose the use of the mitochondrial isoform of dUTPase as a potential reference gene for studies involving comparisons between different organs in mice. For gene expression studies involving mitochondrial proteins, normalization is usually carried out to expression levels of traditional reference genes [42,43]; however, the use of transcripts for mitochondrial proteins as reference genes, such as the mitochondrial isoform of dUTPase, may be beneficial for gene expression studies of mitochondrial proteins as well.

Materials and methods

Animals

Mice used in the experiments were produced and maintained in the Animal Care Facility at the NAIK.
Agricultural Biotechnology Center (FVB/N background, Envigo, Huntingdon, UK). Animals were housed in groups of 2–5 with free access to food and water. Animals were kept under standard light–dark cycle (06.00–18.00 h) at 22 °C. This study was carried out in strict accordance with the recommendations and rules in the Hungarian Code of Practice for the Care and Use of Animals for Scientific Purposes. The protocol was approved by the Animal Care and Ethics Committee of the NAIC-Agricultural Biotechnology Institute and the Pest County’s governmental office (permission number: PEI/001/329-4/2013). The method used for euthanasia: cervical dislocation. All efforts were made to minimize suffering.

RNA isolation
Mice were euthanized and organs were prepared by macrodissection and were quick-frozen in liquid nitrogen and stored in –70 °C until processing, which was carried out typically within a month. RNA isolation was carried out as quickly as possible using RNeasy Plus Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. The organs were kept in liquid nitrogen, and approximately 20 mg of samples was excised. Before homogenization, samples were kept on ice. Mini beadbeater 96 (Biospec Products, Bartlesville, OK, USA) was used for homogenization with 0.9–2.0 mm RNase-free stainless-steel beads (Next Advance, Averill Park, NY, USA) for 1 min. DNase treatment was carried out with RNase-Free DNase Set (Qiagen) according to the manufacturer’s instructions. RNA was eluted in 50 µL of RNase-free water (Ambion, Austin, TX, USA), and the concentration and purity were determined with NanoDrop ND-1000 (Table S2). Samples were diluted to 24 ng µL⁻¹ final concentration. The integrity and genomic DNA contamination of all RNA samples were tested using nondenaturing gel electrophoresis with 1% agarose gel in TBE buffer stained with ethidium bromide (Fig. S3). GeneRuler 1kb plus DNA ladder (Thermo Scientific, Waltham, MA, USA) was loaded as marker. Gel Doc XR+ Imager (Bio-Rad) was used for imaging.

cDNA synthesis
All RNA samples—except those that were used to assess the applicability of the iScript cDNA Synthesis Kit (Bio-Rad)—were subjected to cDNA synthesis using the High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems) according to the manufacturer’s recommendations in reactions of 20 µL total volume with random hexamer primers. Thermal cycling conditions were set in ProFlex PCR System (Applied Biosystems). For the determination of optimal conditions for cDNA synthesis, four point 4-fold serial dilutions of RNA were prepared and introduced to the RT reaction with a starting amount of 800 ng. For all other purposes, 200 ng of RNA was used for each individual cDNA synthesizing reaction, unless otherwise noted. cDNA samples were kept at –20 °C until further processing.

Design of primers
For the design of primers, the sequences of the transcripts of interest were downloaded from the RefSeq database. The gene Dut has two known splice variants that differ in the first exons (accession numbers: NM_023595.6 and NM_001159646.1, for the nuclear and mitochondrial isoform, respectively). For our experiments, we have confirmed the corresponding genomic DNA sequence of Dut by sequencing and isoform-specific primers were designed. Two reference genes were also used, GAPDH and PPIA. GAPDH has two known splice variants in mice (accession numbers: NM_001289726.1 and NM_008084.3) differing in the first exons; thus, primers were designed to amplify the 3‘ common region of the mRNA. For GAPDH, an exon junction spanning forward and an exonic reverse primer were designed. PPIA has only one splice variant (NM_008907.2). For PPIA, exonic forward and reverse primers were designed. The design of primers and the prediction of the specificity were carried out with the Primer-BLAST tool [22]. The oligonucleotides were ordered from Sigma–Aldrich (St. Louis, MO, USA) with desalting purification.

qPCR method
For the amplification, MyTaq HS Mix (Bioline, London, UK), Evagreen dye (Biotium, Hayward, CA, USA), RNase-free water (Ambion), cDNA template, and appropriate primers were mixed in 10 µL final volume reactions. Clear Hard-Shell 96-Well PCR Plates (Bio-Rad) and Microseal ‘B’ PCR Plate Sealing Film (Bio-Rad) were used. For the temperature gradient optimization, cDNA reverse-transcribed from 200 ng of RNA isolated from 10-week-old ovary sample was used to result in a final dilution of 0.05 µL/10 µL reaction. Final dilution in these cases was reached in a series of dilutions. For the primer concentration optimization, cDNA reverse-transcribed from 200 ng of RNA isolated from 10-week-old ovary sample was used to result in a final dilution of 0.055 µL/10 µL reaction. For the determination of optimal conditions for cDNA synthesis, RNA was isolated from 10-week-old female brain, heart, and ovary and 4-week-old male spleen and thymus with three biological replicates from different mice for each organ. Equal amount of each replicate RNA sample was pooled and cDNA was prepared—with either High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems) or iScript cDNA Synthesis Kit (Bio-Rad)—using 800, 200, 50, and 12.5 ng of total RNA. The resulting cDNA samples were subjected to qPCR in a final dilution of 0.31 µL/10 µL reaction.
For determining the PCR efficiencies, cDNA samples with three biological replicates from different mice for each organ were reverse-transcribed from 200 ng of RNA isolated from 10-week-old female brain, heart, and ovary and 4-week-old male spleen and thymus. Equal amount of each replicate cDNA sample was pooled and used to prepare fourfold serial dilutions. cDNA was subjected to qPCR in final dilutions of 0.5 µL, 0.0125 µL, 3125 × 10^{-3} µL, 7.81 × 10^{-4}, and 1.95 × 10^{-4} µL per 10 µL reaction.

For the assessment of the optimized RT-qPCR assay, cDNA from 200 ng of RNA isolated from 8 different organs (brain, heart, kidney, liver, lung, ovary/testicle, spleen, and thymus) of 10-week-old male and female mice were subjected to qPCR in a final dilution of 0.31 µL/10 µL reaction. Each organ from either female or male mice forms a biological group, which includes three biological replicates from different mice.

The primers used for amplification are summarized in Table 1. GAPDH-Fw and GAPDH-Rev primers were used in a final concentration of 315 nM; PPIA-Fw and PPIA-Rev primers were used in a final concentration of 425 and 93.75 nM. For all other experiments, 1200 nM final concentration was used for Rev1 primer. For the temperature gradient optimization, nDut-Fw and mDut-Fw primers were used in final concentrations of 300 and 375 nM, respectively. For the primer concentration optimization, Rev1, Rev2, Rev3, and Rev4 primers were used in final concentrations of 1000, 500, 250, 125, and 62.5 nM. For all other experiments, 1000 nM final concentration was used for Rev1 primer. For the temperature gradient optimization, nDut-Fw and mDut-Fw primers were used in final concentrations of 300 and 375 nM, respectively. For the primer concentration optimization, nDut-Fw primer was used in final concentrations of 1200, 600, 300, 250, and 125 nM, while mDut-Fw primer was used in final concentrations of 1500, 750, 375, 187.7, and 93.75 nM. For all other experiments, 1200 nM final concentration was used for nDut-Fw primer and 375 nM final concentration was used for mDut-Fw primer.

A gene maximizing plate design was used with manual reaction setup. Each plate contained four replicates of no template controls (NTC) for each target and three technical replicates of no reverse transcriptase (NRT) controls for each target and each sample. All other reactions were carried out in triplicates, unless otherwise noted.

PCR cycling was performed at 95 °C for 5 min followed by 50 cycles of denaturation at 95 °C for 30 s, annealing and extension at 66 °C for 30 s, and melting curve analysis from 60 to 95 °C at a rate of 0.5 °C/5 s. CFX96 real-time PCR detection system (Bio-Rad) was used for thermal cycling and detection.

**Assessment of the specificity of PCR**

The specificity of the PCR products was tested first using gel electrophoresis with 2% agarose gel in TBE buffer stained with GelRed nucleic acid gel stain (Biotium). GeneRuler 100 bp plus DNA ladder (Thermo Scientific) was loaded as marker. Gel Doc XR+ Imager (Bio-Rad) was used for imaging. To confirm the identity of all PCR products—except for the PCR product for the nuclear and mitochondrial isoforms, which were too short for sequencing—DNA was isolated from agarose gel with NucleoSpin gel and PCR clean-up (Macherey-Nagel, Duren, Germany) according to the manufacturer’s recommendations and sequenced by Microsynth Seqlab, Göttingen, Germany. To confirm the identity of the PCR products for the nuclear and mitochondrial isoforms, PCR product for both isoforms with Rev4 reverse primer using ovary cDNA was isolated from agarose gel. The concentration of the DNA was determined with NanoDrop ND-2000, and the samples were used as template in a second round of PCR with 10 and 0.1 fg µL^{-1} final concentration for both isoforms. For further reactions, the specificity of the formed PCR products was confirmed with melting curve analysis. Results were accepted only with uncompromised product specificity—or indicated otherwise for the process of optimization.

**Analysis of data**

Amplification curves and melting curves were illustrated by CFX Maestro 1.1 software (Bio-Rad). Cq determination mode was set to single threshold. For primer optimization experiments, all data were used in the analysis, and none were excluded. However, in other experiments we have set several criteria to define outlier data. First, Cq values were identified as outlier and disposed of, if the final RFU value was lower by 10% than the average final RFU value for the corresponding target. Second, Cq values were identified as outlier and disposed of, if the melting curve analysis showed aspecific product formation. Approximately 5% of the data were excluded from the analysis based on these two criteria. Third, the differences in Cq values between the unknown samples and NTC/NRT controls were investigated. In every case, the difference was more than 5 cycles; however, in most cases, it was more than 8 cycles, thus all results were accepted based on this criterion. Relative quantity was calculated and illustrated by CFX Maestro software (Bio-Rad). Scatter graphs and 3D bar graphs were created with ORIGINPRO 2018 (OriginLab Corp., Northampton, MA, USA).

For the results presented in Fig. 7, ANOVA was performed with TIBCO STATISTICA 13.4 (Tibco Software Inc., Palo Alto, CA, USA). For the model, the random ‘sample’ variable was nested into the interaction term of the variables ‘organ’ and ‘sex’, and type III (orthogonal) decomposition was used. To check for the homogeneity of variances, the Bartlett chi-square test was used, and normality of the within-cell residuals was checked on normal probability plots. The residuals were also checked by plotting them against the observed values. In all cases, the assumptions for the use of ANOVA were met. The standard deviation of the Cq values was calculated as the square root of the mean sum of squares due to error.
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Conflict of interest

The authors declare no conflict of interest.

Author contributions

GR and BV planned the experiments; GR and NN performed the experiments and analyzed the data; ZG, TP, and LH were involved in sample preparation; GR, NN, and BV wrote the paper.

References

1 Vérettsy BG and Tóth J (2009) Keeping uracil out of DNA: physiological role, structure and catalytic mechanism of dUTPases. Acc Chem Res 42, 97–106.
2 Kerepesi C, Szabó JE, Papp-Kádár V, Dobay O, Szabó D, Grolmusz V and Véretissy BG (2016) Life without dUTPase. Front Microbiol 7, 1768.
3 Dengg M, Garcia-Muse T, Gill SG, Ashcroft N, Boulton SJ and Nilsen H (2006) Abrogation of the CLK-2 checkpoint leads to tolerance to base-excision repair intermediates. EMBO Rep 7, 1046–1051.
4 Kumar H, Kehrer J, Singer M, Reining M, Santos JM, Mair GR and Frischknecht F (2019) Functional genetic evaluation of DNA house-cleaning enzymes in the malaria parasite: dUTPase and Ap4AH are essential in Plasmodium berghei but ITPase and NDH are dispensable. Expert Opin Ther Targets 23, 251–261.
5 Muha V, Horváth A, Békési A, Pukánszki M, Hodosesek B, Merényi G, Róna G, Batki J, Kiss I, Jankovics F et al. (2012) Uracil-containing DNA in Drosophila: stability, stage-specific accumulation, and developmental involvement. PLoS Genet 8, e1002738.
6 Muha V, Zagvya I, Venkei Z, Szabad J and Véretissy BG (2009) Nuclear localization signal-dependent and -independent movements of Drosophila melanogaster dUTPase isoforms during nuclear cleavage. Biochem Biophys Res Commun 381, 271–275.
7 Ladner RD and Caradonna SJ (1997) The human dUTPase gene encodes both nuclear and mitochondrial isoforms. Differential expression of the isoforms and characterization of a cDNA encoding the mitochondrial species. J Biol Chem 272, 19072–19080.
8 Róna G, Marfori M, Borsos M, Scheer I, Takács E, Tóth J, Babos F, Magyar A, Erdéi A, Bozóky Z et al. (2013) Phosphorylation adjacent to the nuclear localization signal of human dUTPase abolishes nuclear import: structural and mechanistic insights. Acta Crystallogr D Biol Crystallogr 69, 2495–2505.
9 Pri-Hadash A, Hareven D and Lifschitz E (1992) A meristem-related gene from tomato encodes a dUTPase: analysis of expression in vegetative and floral meristems. Plant Cell 4, 149–159.
10 Békési A, Zagvya I, Hunyadi-Gulyás E, Pongrácz V, Kovári J, Nagy AO, Erdéi A, Medzhirdzsky KF and Véretissy BG (2004) Developmental regulation of dUTPase in Drosophila melanogaster. J Biol Chem 279, 22362–22370.
11 Wilson PM, Fazzzone W, LaBonte MJ, Deng J, Neamati N and Ladner RD (2008) Novel opportunities for thymidylate metabolism as a therapeutic target. Mol Cancer Ther 7, 3029–3037.
12 Takatori H, Yamashita T, Honda M, Nishino R, Arai K, Yamashita T, Takamura H, Ohta T, Zen Y and Kaneko S (2010) dUTP pyrophosphatase expression correlates with a poor prognosis in hepatocellular carcinoma. Liver Int 30, 438–446.
13 Kawahara A, Akagi Y, Hattori S, Mizobu T, Shirouzu K, Ono M, Yanagawa T, Kuwano M and Kage M (2009) Higher expression of deoxyuridine triphosphatase (dUTPase) may predict the metastasis potential of colorectal cancer. J Clin Pathol 62, 364–369.
14 Huntley MA, Lou M, Goldstein LD, Lawrence M, Dijkstra GJP, Kaminker JS and Gentleman R (2016) Complex regulation of ADAR-mediated RNA-editing across tissues. BMC Genom 17, 61.
15 Brawand D, Soumillon M, Nesiculea A, Julien P, Csárdi G, Harrigan P, Weier M, Liechti A, Aximu-Petri A, Kircher M et al. (2011) The evolution of gene expression levels in mammal organs. Nature 478, 343–348.
16 Keane TM, Goodstadt L, Danecek P, White MA, Kong W, Yalcin B, Heger A, Agam A, Slater G, Goodson M et al. (2011) Mouse genomic variation and its effect on phenotypes and gene regulation. Nature 477, 289–294.
17 Soumillon M, Nesiculea A, Weier M, Brawand D, Zhang X, Gu H, Barthès P, Kokkinaki M, Nie S, Grunke A et al. (2013) Cellular source and mechanisms of high transcriptome complexity in the mammalian testis. Cell Rep 3, 2179–2190.
18 Barbosa-Morais NL, Irimia M, Pan Q, Xiong HY, Gueroussov S, Lee LJ, Slobodeniuc V, Kutter C, Watt S, Colak R et al. (2012) The evolutionary landscape of
alternative splicing in vertebrate species. Science 338, 1587–1593.

19 Mamedov TG, Pienaar E, Whitney SE, TerMaat JR, Carvill G, Goliath R, Subramanian A and Viljoen HJ (2008) A fundamental study of the PCR amplification of GC-rich DNA templates. Comput Biol Chem 32, 452–457.

20 Bustin SA, Benes V, Garson JA, Hellemans J, Huggett J, Kubista M, Mueller R, Nolan T, Pfaffl MW, Shipley GL et al. (2009) The MIQE guidelines: minimum information for publication of quantitative real-time PCR experiments. Clin Chem 55, 611–622.

21 Debode F, Marien A, Janssen J, Kubista M, Mueller R, Nolan T, Pfaffl MW, Shipley GL et al. (2009) The MIQE guidelines: minimum information for publication of quantitative real-time PCR experiments. Clin Chem 55, 611–622.

22 Ye J, Coulouris G, Zaretskaya I, Cutcutache I, Rozen S and Madden TL (2012) Primer-BLAST: a tool to design target-specific primers for polymerase chain reaction. BMC Bioinformatics 13, 134.

23 Radonić A, Thulke S, Mackay IM, Landt O, Siegert W and Nitsche A (2004) Guideline to reference gene selection for quantitative real-time PCR. Biochem Biophys Res Commun 313, 856–862.

24 Eberhard M, Kurtudoğan M,average = 3.167174603174604; median = 3; standard deviation = 1.9999999999999996; variance = 3.9999999999999996; range = 6; minimum = 1; maximum = 6; mode = 3; frequency = 3; count = 6; null frequency = 0; unique count = 6; unique values = 6; null count = 0; null percentage = 0; median absolute deviation = 1; standard error = 0.16666666666666666; Skewness = 0; Kurtosis = 0; # 313, 856–862.

25 Ren S, Zhang F, Li C, Jia C, Li S, Xi H, Zhang H, Yang L and Wang Y (2010) Selection of housekeeping genes for use in quantitative reverse transcription PCR assays on the murine cornea. Mol Vis 16, 1076–1086.

26 Ali H, Du Z, Li X, Yang Q, Zhang YC, Wu M, Li Y and Zhang G (2015) Identification of suitable reference genes for gene expression studies using quantitative polymerase chain reaction in lung cancer in vitro. Mol Med Rep 11, 3767–3773.

27 Gong Z-K, Wang S-J, Huang Y-Q, Zhao R-Q, Zhu Q-F and Lin W-Z (2014) Identification and validation of suitable reference genes for RT-qPCR analysis in mouse testis development. Mol Genom 289, 1157–1169.

28 Nolan T, Hands RE and Bustin SA (2006) Quantification of mRNA using real-time RT-PCR. Nat Protoc 1, 1559–1582.

29 Gunson R, Gillespie G and Carman WF (2003) Optimisation of PCR reactions using primer chessboarding. J Clin Virol 26, 369–373.

30 Ståhlberg A, Häkansson J, Xian X, Semb H and Kubista M (2004) Properties of the reverse transcription reaction in mRNA quantification. Clin Chem 50, 509–515.

31 Miranda JA and Steward GF (2017) Variables influencing the efficiency and interpretation of reverse transcription quantitative PCR (RT-qPCR): An empirical study using Bacteriophage MS2. J Virol Methods 241, 1–10.

32 Pfaffl MW (2004) Quantification strategies in real-time PCR. In A-Z of Quantitative PCR (Bustin SA, ed.), pp. 87–112. International University Line, La Jolla, CA.

33 Almeida AM, Castel-Branco MM and Falcão AC (2002) Linear regression for calibration lines revisited: weighting schemes for bioanalytical methods. J Chromatogr B Anal Lett Technol Biomed Life Sci 774, 215–222.

34 Ellison S, English CA, Burns MJ and Keer JT (2006) Routes to improving the reliability of low level DNA analysis using real-time PCR. BMC Biotechnol 6, 33.

35 Svec D, Tichopad A, Novosadova V, Pfaffl MW and Kubista M (2015) How good is a PCR efficiency estimate: recommendations for precise and robust qPCR efficiency assessments. BiomedaupeQuantif 3, 9–16.

36 Fleige S and Pfaffl MW (2006) RNA integrity and the effect on the real-time qRT-PCR performance. Mol Aspects Med 27, 126–139.

37 Bustin SA and Nolan T (2004) Template handling, preparation, and quantification. In A-Z of Quantitative PCR (Bustin S, ed.), pp. 141–215. International University Line, 2004–2006, La Jolla, CA.

38 Hruz T, Wyss M, Docquier M, Pfaffl MW, Masanetz S, Borghi L, Verbrugghe P, Kalaydjieva L, Bleuler S, Laule O et al. (2011) RefGenes: identification of reliable and condition specific reference genes for RT-qPCR data normalization. BMC Genom 12, 156.

39 Alberts B, Johnson A, Lewis J, Morgan D, Raff M, Roberts K and Peter W (2015) The innate and adaptive immune systems. In Molecular Biology of the Cell, 6th edn (Zayatz E-B, Lewis SG and Clayton J, eds), pp. 1297–1342. Garland Science, New York, NY.

40 Veltri KL, Espiritu M and Singh G (1990) Distinct genomic copy number in mitochondria of different mammalian organs. J Cell Physiol 143, 160–164.

41 Bromfield JJ and Piersanti RL (2019) In Mammalian oogenesis: the fragile foundation of the next generation. In The Ovary, 3rd edn (Leung PCK and Adashi EY, eds), pp. 157–220. Academic Press, Cambridge, MA.

42 Tang P-A, Duan J-Y, Wu H-J, Ju X-R and Yuan M-L (2017) Reference gene selection to determine differences in mitochondrial gene expressions in phosphate-susceptible and phosphate-resistant strains of Cryptolestes ferrugineus, using qRT-PCR. Sci Rep 7, 7047.

43 Merlo Pich M, Raule N, Catani L, Fagioli ME, Faenza I, Coco L and Lenaz G (2004) Increased transcription of mitochondrial genes for Complex I in human platelets during ageing. FEBS Lett 558, 19–22.

**Supporting information**

Additional supporting information may be found online in the Supporting Information section at the end of the article.
**Fig. S1.** Possible secondary structures of the PCR products as predicted by the OligoAnalyzer tool of Integrated DNA Technologies. The introduced structures were selected based on the highest change in Gibbs free energy according to the tool. Blue circles indicate A-T interaction, red circles indicate G-C interaction, green indicates G-T interaction. (A, B) Secondary structure of the nuclear (A) and mitochondrial (B) isoform with isoform-specific forward primers and Rev4 reverse primer. (C, D) Secondary structure of the nuclear (C) and mitochondrial (D) isoform with isoform-specific forward primers and Rev1 reverse primer. (E) Table summarising the thermodynamic parameters of the four introduced structures.

**Fig. S2.** (A) Agarose gel electrophoresis of the PCR products for the nuclear isoform with Rev1 reverse primer at a range of annealing temperature from 53.1 to 69 °C. The specific product is indicated with an arrow. Vertical line indicates cropping of the image. (B, C) Agarose gel electrophoresis of the PCR products for both nuclear and mitochondrial isoforms with Rev2 reverse primer at a range of annealing temperature from 53.1 to 69 °C (B) and from 61.2 to 66 °C (C). The specific products are indicated with arrows. Vertical line indicates cropping of the image. (D, E) Quantification cycles (Cq) at a range of annealing temperatures from 61.2 to 66 °C with all four reverse primer candidates for the mitochondrial isoform (D) and nuclear isoform (E). The solid squares indicate specific products as determined with agarose gel electrophoresis and melting curve analysis. Open squares indicate aspecific products. (F) Quantification cycles (Cq) at a range of annealing temperatures from 53.1 to 69 °C with the selected primers. The solid squares indicate specific products as determined with agarose gel electrophoresis and melting curve analysis. Open squares indicate aspecific products.

**Fig. S3.** Agarose gel electrophoresis of the RNA samples used in this study. Vertical lines indicate cropping of the image.

**Fig. S4.** Amplification curves of all the biological groups. Red colour indicates the nuclear isoform, blue indicates the mitochondrial isoform, green indicates GAPDH and brown indicates PPIA. Three biological replicates and three technical replicates for each biological replicates are shown for each graph. Threshold values are set to 500 RFU. RFU, relative fluorescent unit. The biological groups of the same organs are juxtaposed with the female sex first.

**Table S1** Parameters of the fitted linear curves used to determine qPCR efficiencies. Weighted least squares linear regression was performed to the Cq values from 4-fold cDNA dilution. Accordingly, base 4 logarithm was used for the efficiency calculation.

**Table S2** Purity of the RNA samples used in this study indicated by 260/280 absorbance ratio values measured by Nanodrop.

**Table S3** MIQE checklist for our study. ✔: fulfillment of the requirements; – : not provided data; N/A : not applicable parameter.