Identification and validation of stable reference genes for quantitative real time PCR in different minipig tissues at developmental stages

Jeongah Song¹*, Jeonghee Cho¹,², Jeongsik Park¹ and Jeong Ho Hwang¹*

Abstract

Background: Quantitative real time PCR (qPCR) is a powerful tool to evaluate mRNA expression level. However, reliable qPCR results require normalization with validated reference gene(s). In this study, we investigated stable reference genes in seven tissues according to four developmental stages in minipigs. Six candidate reference genes and one target gene (ACE2) were selected and qPCR was performed. BestKeeper, geNorm, NormFinder, and delta Ct method through the RefFinder web-based tool were used to evaluate the stability of candidate reference genes. To verify the selected stable genes, relative expression of ACE2 was calculated and compared with each other.

Results: As a result, HPRT1 and 18S genes had lower SD value, while HMBS and GAPDH genes had higher SD value in all samples. Using statistical algorithms, HPRT1 was the most stable gene, followed by 18S, β-actin, B2M, GAPDH, and HMBS. In intestine, all candidate reference genes exhibited similar patterns of ACE2 gene expression over time, whereas in liver, lung, and kidney, gene expression pattern normalized with stable reference genes differed from those normalized with less stable genes. When normalized with the most stable genes, the expression levels of ACE2 in minipigs highly increased in intestine and kidney at PND28, which is consistent with the ACE2 expression pattern in humans.

Conclusions: We suggest that HPRT1 and 18S are good choices for analyzing all these samples across the seven tissues and four developmental stages. However, this study can be a reference literature for gene expression experiments using minipig because reference gene should be validated and chosen according to experimental conditions.

Keywords: Quantitative real time PCR, Reference genes, Minipig, RefFinder

Background

Quantitative real-time-polymerase chain reaction (qPCR) is a powerful tool to evaluate mRNA expression level by measuring the fluorescent dyes binding to the amplified double-stranded DNA [1]. This technique is a useful and accurate technique due to its high sensitivity and reproducibility [2]. In addition, qPCR enables high-throughput analysis with ease. However, it needs to control for errors during experiment, such as RNA quality, initial sample amount, differences in transcriptional activity of samples, and PCR efficiency [2]. To control this error, a similar amount of good-quality RNA should be used for reverse transcription, and the gene expression level should be normalized to validated reference genes. Housekeeping genes are commonly used as a reference [2, 3]. Housekeeping genes are genes that are essential...
for the maintenance of cellular function, such as cell metabolism and structural gene [4]. They are expected to be expressed at a constant level across all tissue types, regardless of developmental status and experimental conditions [4]. However, this ideal housekeeping gene has not been found [5, 6]. Well-known housekeeping genes are not constantly expressed in all tissues, which might be partly because housekeeping genes have functions other than the maintenance of cellular function [7–10]. A myriad of housekeeping genes, such as glyceraldehyde-3-phosphate dehydrogenase (GAPDH), tubulins, actins, albumins, cyclophilin, microglobulins, ribosomal units (18S or 28S rRNA), ubiquitin (UBQ), and new candidate normalization genes, which were suggested using microarray [5, 11, 12], have been described. Many researchers recommended that when performing quantitative gene expression study, reference genes need to be validated to ensure stable expression in each experimental condition [13].

Researchers have tried to develop various algorithms, such as BestKeeper, NormFinder, geNorm, delta Ct, and Reffinder, with which the expression variability or stability of candidate reference genes can be measured. Bestkeeper is an excel-based software tool (http://www.gene-quantification.org/bestkeeper.html) that determines gene expression stability based on the variation of Ct values and Pearson correlation coefficient (r), which is calculated based on the inter-gene relation of all reference pairs and the BestKeeper Index, which is calculated as the geometric mean of the remaining genes. Genes with standard deviation (SD) value higher than 1 are considered unstable [14]. Normfinder was developed by Anderson et al. (2004) (https://moma.dk/normfinder-software). This algorithm calculates a stability value by combining intragroup and intergroup variation for candidate reference genes [5]. geNorm calculates the average pairwise variation for a reference gene with all other genes and presents it as M value. The lowest M value represents the most stable gene expression [15]. Delta Ct method compares the relative expression of ‘pairs of reference genes’ within each sample, and sequentially eliminates the reference gene with a large deviation in delta Ct values. Through this process, the most stable reference genes can be selected [16]. Finally, Reffinder is a free web-based software that performs an analysis based on the delta Ct method, geNorm, Normfinder, and bestkeeper. Moreover, it provides comprehensive ranking order of the candidate reference genes based on the geometric mean of ranking values obtained from the aforementioned analysis [17, 18]. These algorithms have been widely used in the studies of identification of reference genes.

Minipigs (Sus scrofa) are useful large experimental animals, in that they share anatomical and physiological similarities with humans, such as metabolic profile, renal system, and longer lifespan. They have been developed for biomedical research purposes since the late 1940s, resulting in at least 45 breeds of minipig breeds; and since the 1980s, the use of minipigs in research has steadily increased [19, 20]. Minipigs present many advantages over other rodents, dogs, or nonhuman primates. They have a long gestation period of approximately 114 days (d) [21], while mice, rats, and dogs have a short gestation period of 19–21 d, 21–23 d, and 58–68 d, respectively [22–24]. Minipigs have relatively large litter size (5–6 pigs per litter) and their birth weight of 400–500 g makes it easy to manipulate for dosing from an early age, or for surgical intervention, or for sampling of blood [25]. Therefore, minipigs are a useful model for juvenile toxicity studies. In addition, their long gestation period, large fetal size, and fetal weight make minipig a good model for the investigation of fetal development, including the immune system, nervous system, or specific organs.

In this study, we investigated stable reference genes in minipigs. To find out the appropriate reference genes in various tissues at different developmental stages, we measured the mRNA expression of six housekeeping genes and one target gene, angiotensin-converting enzyme2 (ACE2), in the heart, lung, liver, kidneys, stomach, intestine, and spleen of fetus (gestation d 70 (GD 70) and GD85), neonates (postnatal d 1 (PND1)) and piglets. We then analyzed the expression level of ACE2 against 6 reference genes, and compared them according to the tissues and developmental stages, to confirm their suitability for gene expression study.

**Results**

**Primer validation for specificity and PCR amplification efficiency**

To verify the specificity of primer pairs, melting curves were examined and agarose gel electrophoresis was performed. A single peak was detected in all primer pairs, and a single band was observed on agarose gel (Supplementary Figs. 1 and 2). PCR amplification efficiencies ranged from 90.30 to 97.19% (Table 1).

**Expression levels of various reference genes**

The mRNA expression of the six candidate reference genes was measured in the seven tissues of fetus (GD 70 and 85), neonates (PND1) and 4-week-old piglets (PND28). Supplementary Table 1 shows the average Ct values, minimum or maximum values of each gene. The six candidate reference genes showed Ct values ranging 13.93 to 26.09. β-actin and 18S were relatively highly expressed genes, with Ct values ranging 13.93 to 18.63 and 14.86 to 18.94, respectively. HMBS and HPRT1 were
the relatively least expressed, with Ct values ranging 18.12 to 26.09 and 21.34 to 24.98, respectively.

Analysis of the Ct values distribution of each candidate reference gene across the different tissues showed that HPRT1 and 18S genes had lower standard deviation value, which means similar levels of HPRT1 and 18S were expressed in the seven tissues, regardless of the developmental stage (Supplementary Table 1 and Fig. 1). In contrast, HMBS and GAPDH genes had higher SD value (Supplementary Table 1 and Fig. 1). The Ct values of all six candidate reference genes were altered across the different tissues.

Next, the Ct values of candidate reference genes were analyzed according to the developmental stages (Fig. 2). Four candidate reference genes except for B2M and 18S showed no differences in average Ct values, which means these four genes were stably expressed at all these

**Table 1** Amplification efficiency of standard curves in candidate reference genes and target gene

| Gene symbol | Efficiency (%) | Slope   | R² values |
|-------------|---------------|---------|-----------|
| ACE2        | 94.12         | −3.471  | 0.999     |
| B2M         | 96.64         | −3.405  | 1.000     |
| β-actin     | 90.30         | −3.579  | 0.999     |
| GAPDH       | 95.32         | −3.439  | 0.997     |
| HMBS        | 92.59         | −3.513  | 0.948     |
| HPRT1       | 97.19         | −3.391  | 0.999     |
| 18S         | 93.01         | −3.502  | 0.998     |

*a* PCR amplification efficiency is calculated by the following formula: Efficiency = \((10^{-\text{slope}} - 1) \times 100\)

---

**Fig. 1** Distribution of Ct values of each reference gene across the different tissues of minipig. The expression levels are shown as median (middle line), 25th to 75th percentiles (boxes), and range (whiskers) for 15 animals of GD70 (n = 4), GD85 (n = 5), PND1 (n = 3), and PND28 (n = 3). (n = 15 per tissue)
developmental stages. The average Ct values significantly decreased over time in B2M gene and 18S showed a little bit higher Ct value on PND1, compared to the Ct values of other stages (Supplementary Table 1 and Fig. 2).

Assessment of the expression stability of reference genes using web-based analysis tools
Reference genes are expected to be consistently expressed with low variation in samples. To evaluate the expression stability of candidate reference genes, statistical algorithms, BestKeeper, delta Ct, geNorm, Normfinder, and RefFinder, were employed. Using BestKeeper algorithm, the order of gene stability in all samples was as follows: HPRT1 > 18S > β-actin > GAPDH > B2M > HMBS (Table 2). The BestKeeper algorithm showed that the five candidate reference genes are considered stably expressed, excepting HMBS with SD higher than 1 [14]. The calculated stability index of delta Ct, geNorm, and Normfinder algorithms designated HPRT1 as the most stable gene, followed by 18S, β-actin, B2M, GAPDH, and HMBS (Table 3). The recommended comprehensive ranking calculated by RefFinder, which automatically calculates the geometric mean of ranking values obtained from the aforementioned these four algorithms, was also concordant with this ranking. Only BestKeeper showed swapped place of GAPDH and B2M (Table 3)

Regarding tissue type and developmental stage, Tables 4 and 5 show the comprehensive rankings. In the case of tissues, the candidate reference genes ranked top 3 were apparently variable according to tissue. GAPDH was the most stable reference gene in liver, intestine, and kidney, and the expression of HPRT1 was the most stable in spleen and stomach. B2M was the least stable gene in heart, lung, stomach, intestine, and kidneys. Regarding developmental stage, HPRT1 exhibited the most stable expression of GD70, GD85, and PND1. On PND28, the expression of 18S was the most stable. HMBS was the least stable gene, excepting PND1. Supplementary Tables 2–5 show the detailed stability index and ranking order according to the integrative RefFinder and BestKeeper, Normfinder, CT values, and geNorm.
Assessment of the expression levels of target gene against various reference genes

The ranking order of reference genes based on the comprehensive rankings of RefFinder algorithm in all samples was HPRT1 > 18S > β-actin > B2M > GAPDH > HMBS (Table 3). To validate the stability of the candidate reference genes, we measured and analyzed the mRNA expression levels of ACE2 according to tissue type and developmental stage. Relative quantification of ACE-2 genes was performed using the $2^{-\Delta\Delta Ct}$ method.

First, the relative expression levels of ACE2 were analyzed using the candidate reference genes according to Table 3. The ranking order of reference genes based on the comprehensive rankings of RefFinder algorithm in all samples was HPRT1 > 18S > β-actin > B2M > GAPDH > HMBS. To validate the stability of the candidate reference genes, we measured and analyzed the mRNA expression levels of ACE2 according to tissue type and developmental stage. Relative quantification of ACE-2 genes was performed using the $2^{-\Delta\Delta Ct}$ method.

Table 2: Descriptive statistics and pairwise correlation of the six reference genes analyzed by BestKeeper algorithm

| Gene   | BestKeeper | Delta CT | Genorm | Normfinder | RefFinder |
|--------|------------|----------|--------|------------|-----------|
|        | Rank       | SD       | Rank   | MV         | Rank      | GM       |
| HPRT1  | 1          | 0.59     | 1      | 1.31       | 1         | 0.38     | 1        | 0.70     | 1.00     |
| 18S    | 2          | 0.62     | 2      | 1.41       | 2         | 1.02     | 2         | 1.11     | 1.68     |
| β-actin| 3          | 0.75     | 3      | 1.47       | 3         | 1.11     | 3         | 0.89     | 3.00     |
| GAPDH  | 4          | 0.89     | 5      | 1.81       | 5         | 1.38     | 5         | 1.48     | 4.73     |
| B2M    | 5          | 0.99     | 4      | 1.61       | 4         | 1.21     | 4         | 1.17     | 4.23     |
| HMBS   | 6          | 1.49     | 6      | 2.08       | 6         | 1.62     | 6         | 1.85     | 6.00     |

Table 3: Expression stability values of candidate reference genes in all samples

| Gene   | BestKeeper | Delta CT | Genorm | Normfinder | RefFinder |
|--------|------------|----------|--------|------------|-----------|
|        | Rank       | SD       | Rank   | MV         | Rank      | GM       |
| HPRT1  | 1          | 0.59     | 1      | 1.31       | 1         | 0.38     | 1        | 0.70     | 1.00     |
| 18S    | 2          | 0.62     | 2      | 1.41       | 2         | 1.02     | 2         | 0.70     | 1.68     |
| β-actin| 3          | 0.75     | 3      | 1.47       | 3         | 1.11     | 3         | 0.89     | 3.00     |
| GAPDH  | 4          | 0.89     | 5      | 1.81       | 5         | 1.38     | 5         | 1.48     | 4.73     |
| B2M    | 5          | 0.99     | 4      | 1.61       | 4         | 1.21     | 4         | 1.17     | 4.23     |
| HMBS   | 6          | 1.49     | 6      | 2.08       | 6         | 1.62     | 6         | 1.85     | 6.00     |

Abbreviations: SD standard deviation, MV M-value, SV stability value, GM geometric mean [17]
tissue type and developmental stage. ACE gene was normalized with each of six candidate reference genes. Figures 3 and 4 show that ACE2 revealed a different degree of expression change according to the reference gene used for normalization. In intestine, all candidate reference genes exhibited significant increase of ACE2 gene expression over time, though the degree of change differed among reference genes. However, in heart, liver, lung, and kidney, gene expression level using stable reference genes (HMBS, HPRT1, and GAPDH in heart; GAPDH, 18S, and HPRT1 in liver; HMBS, β-actin, and HPRT1 in lung; and GAPDH, HPRT1, and β-actin in kidney) differed significantly from those using less stable reference genes (B2M and β-actin in heart; HMBS in liver; 18S in lung; and B2M in kidney) (Figs. 3 and 4). In the case of kidney, the gene expression level of ACE2 normalized with the least stable reference gene (B2M) was 0.47 at PND28, while the relative quantity of ACE2 normalized with the most stable reference gene (GAPDH) was 2.28. When HMBS, the least stable gene in the liver, was used as reference gene, the expression of ACE2 increased at PND1 and 28, whereas the expression levels normalized with stable reference genes, GAPDH and HPRT1, significantly decreased. Generally, normalization with the three most stable genes showed similar expression pattern to ACE2.

Next, we analyzed the expression levels normalized with multiple reference genes. Two top-ranked genes (HPRT1/18S) and three low-ranked genes (HMBS/GAPDH/B2M) of all samples were chosen. Gene expression of ACE2 were normalized with their geomean values. Figures 5 and 6 show the results. Generally, normalization with two stable candidate reference genes led to a similar expression value to that with the most stable one in each tissue. Normalization with the three least stable candidate reference genes also resulted in more similar expression pattern of ACE2 to that with the most stable one, than that with normalized with each of the three least stable candidate reference genes.

**Discussion**

Quantitative real time PCR is a useful tool for quantifying gene expression levels [26]. The calculation of relative quantities of genes of interest generally requires a normalization step against reference genes. Housekeeping genes are widely used as reference, because they are ideally expected to be expressed at constant levels in all tissues under variable experimental conditions.
However, many studies revealed that these ideal reference genes do not exist [5, 6]. Therefore, choosing suitable reference genes is very important for the quantification of gene expression.

In this study, we investigated appropriate reference genes for different tissues at developmental stages in minipigs. The minipig is a valuable experimental model for developmental research. They have a long preg
period (114 d) and relatively large litter size [25]. In addition, the sizes of fetuses at each gestation day or neonates are relatively large enough to handle. Our previous studies showed that fetal weights at GD42 were approximately 8.8 g, and their tissues were large enough to collect for analysis. We evaluated six candidate reference genes from 7 different tissues of a total 15 pigs (including fetuses). Overall, β-actin, 18S, and B2M had lower Ct values, and HPRT1 and HMBS showed higher Ct values, which means that β-actin, 18S, and B2M were highly expressed, whereas HPRT1 and HMBS were less expressed. Nygard et al. reported that the order of the relative expression of candidate reference genes in 17 different tissues (from high to low) was: B2M > β-actin > GAPDH > HPRT1 > HMBS [27], while the rank of the relative expression of the reference genes in seven tissues in the current study was: β-actin > 18S > B2M > GAPDH > HMBS > HPRT1 (Table 3). The difference seems to originate from the different developmental stages of different tissue samples. The former used young female siblings, while the latter used minipigs of GD70, 85, and PND1 and 28. However, it can be generally said that β-actin, 18S, and B2M are relatively high-abundance genes, while HMBS and HPRT1 are relatively low-abundance genes [27–29].

Among the six candidate reference genes, HPRT1 showed the lowest SD of Ct values, while 18S and β-actin demonstrated the 2nd and 3rd lowest SD values in all samples. Regarding different tissues, the Ct values of HPRT1 and 18S showed lesser variability in all tissues, whereas the Ct values of GAPDH and HMBS varied across tissues, because lower Ct values of GAPDH and HMBS were detected in heart and spleen, respectively, compared to other tissues. Regarding developmental stages, B2M showed high variation of Ct values, because the Ct values of B2M decreased over time. The decrease in the average Ct value for B2M (19.44 at GD70, 19.06 at GD85, 18.19 at PND1, and 16.93 at PND28) is consistent with the other results performed by Kuijk et al. [29] and Hildyard et al. [30]. Kuijk et al. showed an increase in expression levels of B2M from oocytes to blastocysts. Hildyard et al. showed that the expression levels of B2M increased with increasing gestation days. B2M, beta-2-microglobulin, is a component of major histocompatibility (MHC) class I, and is expressed in all nucleated cells. MHC class I presents peptide derived from viral proteins of infected cells to T cell receptors, and induces adaptive immunity [31]. In addition, B2M is critical for immune cell development. B2M-knockout mice showed lack of NKT cells and mature CD8 T cells [32]. Age-associated upregulation of B2M seems to be developmentally regulated as immunity increases.

To find out stable reference genes, researchers have developed some statistical algorithms or used multiple reference in their gene expression studies. RefFinder is a valuable web-based software, which automatically calculates each stability index of delta Ct, NormFinder, geNorm, and BestKeeper and provides comprehensive

![Fig. 6](https://example.com/fig6.png)

**Fig. 6** Analysis of the expression pattern normalized with the most stable gene (purple box), the three least stable genes (yellow, light yellow, and white), and the geomean of HPRT1/18S (green box) and HMBS/GAPDH/B2M (red box) at each developmental stage. Data are expressed as mean ± SEM; *p < 0.05, and **p < 0.01 indicate significant difference compared to the relative expression normalized with the most stable gene of each tissue or developmental stage.
ranks. RefFinder designated HPRT1 as the most stable gene, followed by 18S, β-actin, B2M, GAPDH, and HMBS in all samples. The seven different tissues and four developmental stages showed different rank from the four software, while RefFinder suggested comprehensive ranks in each tissue and developmental stage by calculating the geomean of ranks derived from the four algorithms. Generally, the high ranking orders of each algorithm seemed to be similar in the same tissue or developmental stage, though there were some exceptions, such as liver and PND1 (Supplementary Table 1 or 2). Depending on the condition of the study, it is necessary to select appropriate reference genes to measure the expression levels of the target genes. For example, in analyzing the expression levels of seven different tissues at different development stages, HPRT1 and 18S were good reference genes. When expression levels are quantified at a fixed time point, HPRT1 and B2M are stably expressed across the different tissues except PND28. At PND28, 18S and B2M were good reference genes. Though the expression levels of B2M increased from fetuses to piglets, at a fixed time B2M showed constant expression levels across the seven tissues.

To validate the candidate reference genes, the expression of ACE2 was measured and analyzed according to the various tissues and developmental stages. As expected, the gene expression pattern varied against the reference genes used for normalization. In the current study, the expression patterns normalized with the three top ranked candidate reference genes seemed generally similar to each other; however, they were quite different from those normalized with the least and less stable genes. At various developmental stages, the expression patterns of ACE2 stayed constant in heart, decreased in liver, and significantly increased in kidney at PND28; whereas, ACE2 expression levels increased in heart and liver and decreased in kidney, when normalized with the least stable genes, β-actin, HMBS, and B2M, respectively. In addition, the relative quantities normalized with B2M were significantly lower than those normalized with other reference genes at PND1 and 28, because expression levels of B2M upregulated over time (Figs. 5 and 6). Relative quantification of qPCR relies on the assumption that the reference genes are stably expressed across all tested samples, regardless of cell or tissue type, age, or treatment. Therefore, normalization with unsuitable reference gene leads to biased qPCR results.

To lessen bias induced by an unsuitable reference gene, many researchers have recommended the use of two or more reference genes for qPCR study [3, 33–35]. For the developmental study in seven tissues, we chose two sets of genes, e.g., one set of HPRT1 and 18S (the two most stable genes of all samples) and another set of HMBS/GAPDH/B2M (the three least stable genes of all samples), and analyzed the expression levels of ACE2 by normalizing with the geomean of two or three genes. The expression levels of ACE2 normalized with the two most stable reference genes, and with the most stable reference gene were generally similar. The expression patterns of ACE2 normalized with the three least stable reference genes seemed, to some extent, to compensate for false results induced by one unsuitable gene. These results imply that normalization with multiple reference genes can provide more reliable qPCR results, though they are the least stable genes. In addition, the use of more than 2 reference genes gives the research an opportunity to evaluate the stability of these genes.

ACE2 is one of the metallocarboxypeptidases that cleaves Angiotensin I into angiotensin 1–9 and angiotensin II into angiotensin 1–7. It plays a key role in diabetes, renal, and cardiovascular disease by counterbalancing the ACE/renin angiotensin aldosterone system which is associated with sodium and water retention, vasoconstriction, hypertension, and accelerated thrombosis [31]. It is also a receptor for severe acute respiratory syndrome coronavirus 2. ACE2 gene is highly expressed in the kidney, heart, small intestine, and testis and is known to be expressed in all organs [36]. In this study, we analyzed the ACE2 expression levels in the seven tissues at each developmental stage. The expression levels of ACE2 in mini-pigs highly increased in intestine and kidney other than tissues at PND1 and PND 28, which is consistent with the ACE2 expression pattern in humans (https://www.ncbi.nlm.nih.gov/gene/59272) [37]. This result indirectly confirmed that our candidate reference genes were good enough for qPCR study.

Dozens of papers have evaluated and validated reference genes in pigs. Martino et al. [38] analyzed the stability of reference genes in cardiac tissues of the healthy and heart failure minipigs. HPRT-1, TBP, and peptidylprolyl isomerase A (PPIA) were the most stably expressed genes in the right and left atrium, while PPIA, GAPDH, and ACTB and HPRT-1, TBP, and GAPDH were the most stable genes in right and left ventricle, respectively. Wang et al. [39] reported that B2M HMBS/HPRT1 were the most stable gene set and GAPDH, β-actin, and 18 were the least stable genes in the gastrointestinal tract during the weaning period. Piórkowska et al. [40] evaluated the stability of 8 candidate reference genes in the adipose tissue of three pig breeds. Ornithine decarboxylase antizyme 1 (OAZ1), 60S ribosomal protein L27 (RPL27), and β-actin were the most stable genes. Nygard et al. [27] showed that β-actin, RPL4, TBP and HPRT1 were expressed at a constant level across 17 different pig tissues. Erkens et al. [28] evaluated the expression stability of 10 reference genes in the porcine backfat.
and longissimus dorsi muscle. β-actin, TBP, and TOP2B were the most stable, while RPL13A and SDHA were the least stable. Kuijk et al. [29] analyzed transcription levels in porcine oocytes and embryos using seven reference genes such as B2M, BACT, GAPDH, H2A, PGK1, S18, and UBC. GAPDH was the most stable gene, while B2M was the least stable gene. These results imply that there were no stable reference genes that encompass all experimental conditions, because researchers showed that stable reference genes differ with experimental condition, namely, strain, age, and tissues of pigs or experimental animals used for experiment.

Conclusions
In this study, we validated six candidate reference genes in seven tissues of minipigs at various stages of development. It is very difficult to find a stable reference gene across various tissues at different developing time points. We suggest that HPRT1 and 18S are a good choice for analyzing all these samples across the seven tissues and developmental stages. Also, the introduction of the multiple reference genes is strongly recommended for qPCR study. However, this study can be a reference literature for gene expression experiments using minipig, because the reference gene should be validated and chosen according to experimental conditions, as gene expression differs according to strain, tissue type, and age of minipigs. Validation of reference genes and the introduction of two or more reference genes lessen the bias, and allow more reliable qPCR data/results in gene expression studies.

Materials and methods
Animal experiments
Pregnant Yucatan miniature pigs were purchased from Optipharm Co., Ltd. (Cheongju, Korea). Sow and piglets were housed in a loose farrowing pen in a controlled environment (temperature of 19–25°C, relative humidity of 50 ± 10%, and air ventilation of 10–20 times per hour (h) with a 12 h/12 h light/dark cycle). Sows were euthanized by intravenous injection of thiopental sodium (0.5 g/15 kg) after anesthesia with xylazine (0.5–3 mg/kg, intramuscular) and ketamine (15 mg/kg, intramuscular). Fetuses at GD70 and GD85 were obtained by hysterectomy. The uterus was exposed through midline abdominal incision. Fetuses were rapidly removed from uterus, and then put in plastic dishes on ice. Gestation d was calculated from the day of mating. For sampling on PND1 (day of birth), neonates were removed from sow within 2–3 h after birth. For sampling on PND28, piglets were euthanized using the same method described above. After euthanasia, heart, lung, liver, kidneys, stomach, intestine, and spleen were removed and placed in RNA later stabilization solution (Sigma Aldrich, MO, USA). All experiments were approved by the Institutional Animal Care and Use Committee of Korea Institute of Toxicology (Approval No.: 2012–0018, 2010–0307, 2008–0250). This manuscript complies with the ARRIVE guidelines [41].

RNA extraction and cDNA synthesis
Each tissue was stored in RNA later stabilization solution at −20°C until analysis. Total RNA was extracted using RNeasy Mini kit (Qiagen, Hilden, Germany), according to the manufacturer’s protocol. RNA yield and purity were assessed using a QIAxpert® system (QIAGEN, Hilden, Germany). RNA samples with A260/280 and A260/230 > 1.8 were used. Four hundred nanogram of total RNA was transcribed using GoScript™ reverse transcription System (Promega, Madison, WI, USA), according to the manufacturer’s instructions.

Selection of reference genes and a target gene
Glyceraldehyde-3-phosphate dehydrogenase (GAPDH), β-actin, and ribosomal protein S18 (18S) are commonly used as reference genes in qPCR studies. Hypoxanthine phosphoribosyltransferase 1 (HPRT1), Hydroxymethylbilane synthase (HMBS), and β2-Microglobulin (B2M) were included in this study from the literature [4, 27, 39, 40]. Angiotensin converting enzyme 2 (ACE2) was chosen as a target gene from the literature. This gene is expressed in many tissues including kidneys, heart, gastrointestinal tract, lung, oral cavity, and liver and the expression pattern of this gene has been extensively studied in humans (https://www.ncbi.nlm.nih.gov/gene/59272) [36, 42–44]. We can compare the expression levels of ACE2 in different tissues of minipigs with those of humans, thus indirectly validating the suitability of the reference genes. The sequence of the selected reference genes and a target gene were obtained from the GenBank database (https://www.ncbi.nlm.nih.gov/genbank/).

Validation of primers and qPCR
PCR primers were designed using NCBI BLAST (https://www.ncbi.nlm.nih.gov/tools/primer-blast/index.cgi?LINK_LOC=BlastHome) or Primer Express® software v3.0.1 (Applied Biosystems, Foster City, CA, USA). Table 6 lists the information of reference genes and primers. The specificity of the primers was checked using the melting curve and agarose gel electrophoresis. PCR amplification efficiency ($E%=(10^{−1/slope}−1)×100$) was measured by means of standard curves. Adult spleen sample was used for this validation.

mRNA levels were quantified using SYBR Green on the Applied Biosystems QuantStudio 5 (Foster City, CA, USA). PCR reactions were performed in a final volume of
20 μL containing 10 μL of SYBR Green PCR Master Mix (Applied Biosystems, Woolston, Warrington, UK), 5 μM of each primer, 5 μL of diluted cDNA, and distilled water. The PCR conditions were 95 °C for 10 min, followed by 40 cycles of 15 s at 95 °C and 1 min at 60 °C. Four or five 5-fold serial dilution of a pool of cDNA samples was used to construct standard curves to evaluate the efficiency of primer pairs.

The relative expression levels of target genes were calculated using the 2-ΔΔCt method [45]. All reactions were carried out in three to five biological replicates and two technical replicates.

Analysis of the expression stability of candidate reference genes
First, we calculated the mean value and standard deviation of the threshold cycle (Ct) value of six candidate reference genes in a total 105 samples. Then, the mean and SD were recalculated according to tissue type and developmental stage.

To evaluate the stability of the candidate reference genes, we used BestKeeper, geNorm, NormFinder, and delta Ct method through the RefFinder web-based tool (https://heartcure.com.au). This statistical analysis was also performed for all samples, according to tissue type and developmental stage. The stability of candidate reference genes was analyzed from each statistical algorithm, and then ranked in most stable to least stable order.

Next, the relative expression of ACE2 was calculated in all samples according to the different tissues and developmental stages to validate the candidate reference genes. Normalization was performed using each of six candidate reference genes, the geomean of the two most stable genes, or the geomean of the three least stable genes.

Relative mRNA quantity was expressed as fold change relative to the average of GD70 or heart. Relative quantification of ACE2 genes was performed using the 2-ΔΔCt method, where

\[ \Delta \Delta Ct = [(Ct_{ACE2} - Ct_{reference gene})_{GDx} - (Ct_{ACE2} - Ct_{reference gene})_{GD70}] - [(Ct_{ACE2} - Ct_{reference gene})_{heart}] \]

Data analysis
To determine the difference of Ct values of reference genes according to the different tissues and developmental stages, one-way analysis of variance with Dunnett’s test was carried out using SPSS 12 (SPSS, Chicago, IL, USA, 2005). The statistical significance of the relative expression of ACE2 was evaluated with Student’s t test using SPSS 12 (SPSS, Chicago, IL, USA, 2005).

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12864-022-08830-z.

| Table 6 Reference genes used in this study and primer |
|-----------------------------------------------------|
| **Gene symbol (gene name)** | **GenBank Accession number** | **Primer forward (F)/reverse (R)** | **Amplicon size** |
| ACE2 (angiotensin converting enzyme 2) | NM_001123070.1 | F: TGCACAGGCCAAATGGT R: ATCTAGTGATACGT TGGGCAAT | 67 |
| B2M (beta-2-microglobulin) | NM_213978.1 | F: CGGAAACGAAATCTCA R: ATCTCTCCTCCCC GTTTTCCA | 88 |
| β-actin (beta-actin) | XM_021086047.1 | F: GATGCGAAAGGAGTACG R: ATCTGCTGG AAGTGGGACAG | 130 |
| GAPDH (Glyceraldehyde 3-phosphate dehydrogenase) | NM_001206359.1 | F: CCACCCAGAAGACTGTGGA R: AAGCAGGGA TGATGTCTGG | 81 |
| HMBS (hydroxymethylbilane synthase) | DQ845174.1 | F: AGGATGGCAACTCTACTCT G: GATGGTGGC CTGCAATGCT | 83 |
| HPRT1 (hypoxanthine phosphoribosyltransferase 1) | NM_001032376.2 | F: AAGCCTTGCTGTGAAGAAGA R: GTCAAGGGG ATAGCCACCA | 100 |
| 18S (18S ribosomal RNA) | NR_046261.1 | F: CCCAGGAATGAGAAGAGA R: TTGACGGAA GGGCAACA | 122 |

Additional file 1.
Additional file 2.

Authors’ contributions
JS designed the experiments; JS, JC, and JP performed the animal study and quantitative real time PCR; JS and JC analyzed the quantitative real time PCR data; JS wrote the manuscript; JH reviewed and editing the data. The authors read and approved the final manuscript.

Funding
We would like to acknowledge the financial support from the Korea Institute of Toxicology (KIT, Korea) grant funded by the Ministry of science and ICT (MIST, Korea) [Project number: KK-2107].

Availability of data and materials
All data generated or analyzed during this study are included in this published article and its supplementary information files.
Declarations

Ethics approval and consent to participate
All procedures performed in studies involving animals were in accordance with the ethical standards of the institution or practice at which the studies were conducted. The study protocol was reviewed and approved by the Institutional Animal Care and Use Committee of Korea Institute of Toxicology (Approval No. 2012–0018, 2010–0307, 2008–0250). This manuscript complies with the ARRIVE guidelines.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Author details
1. Animal Model Research Group, Korea Institute of Toxicology, Jeon-geup 56212, Republic of Korea. 2. Department of Bio-Non- Clinical Science, Graduate School of Konyang University of Bioconvergence, 158, Gwanjeo-dong-ro, Seo-gu, Daejeon 35365, Republic of Korea.

Received: 28 January 2022 Accepted: 28 July 2022 Published online: 13 August 2022

References

1. Bustin SA, Nolan T. Pitfalls of quantitative real-time reverse- transcription polymerase chain reaction. J Biomol Tech. 2004;15:155–66.
2. Huggett J, Dheda K, Bustin S, Zumbil A. Real-time RT-PCR normalisation; strategies and considerations. Genes Immun. 2005;6:279–84.
3. Kozera B, Rapacz M. Reference genes in real-time PCR. J Appl Genet. 2013;54:391–406.
4. Eisenberg E, Levanon EY. Human housekeeping genes, revisited. Trends Genet. 2013;29:569–74.
5. Andersen CL, Jensen JL, Orntoft TF. Normalization of real-time quantitative reverse transcription-PCR data: a model-based variance estimation approach to identify genes suited for normalization, applied to bladder and colon cancer data sets. Cancer Res. 2004;64:5245–50.
6. Zhang Y, Li D, Sun B. Do housekeeping genes exist? PLoS One. 2015;10:e0123691.
7. Bustin SA. Absolute quantification of mRNA using real-time reverse transcription polymerase chain reaction assays. J Mol Endocrinol. 2000;25:169–93.
8. Yperman JDVG, Holvoet P, Flameng W. Beta-actin cannot be used as a control for gene expression in ovine interstitial cells derived from heart valves. J Heart Valve Dis. 2004;13:848–53.
9. Bustin S. Quantification of mRNA using real-time reverse transcription PCR (RT-PCR): trends and problems. J Mol Endocrinol. 2002;29:23–39.
10. Zhu G, Chang Y, Zuo J, Dong X, Zhang M, Hu G, et al. Fudenedine, a C-termi- nal truncated rat homologue of mouse promin, is blood glucose-reg- ulated and can up-regulate the expression of GAPDH. Biochem Biophys Res Commun. 2001;281:951–6.
11. Warrington JA, Nair A, Mahadevappa M, Tsyganskaya M. Comparison of human adult and fetal expression and identification of 535 housekeep- ing/maintenance genes. Physiol Genomics. 2000;2:143–7.
12. McLoughlin KJ, Pedrini E, MacMahon M, Guduric-Fuchs J, Medina RJ. Selection of a real-time PCR housekeeping gene panel in human endothelial Colony forming cells for cellular senescence studies. Front Med (Lausanne). 2019;6:33.
13. Bustin SA, Benes V, Garson JA, Hellemans J, Huggett J, Kubista M, et al. The MIQE guidelines: minimum information for publication of quantitative real-time PCR experiments. Clin Chem. 2009;55:611–22.
14. Pfaff MW, Tichopad A, Prgomet C, Neuvians TP. Determination of stable housekeeping genes, differentially regulated target genes and sample integrity: BestKeeper–Excel-based tool using pair-wise correlations. Biotechnol Lett. 2004;26:509–15.
15. Vandesompele J, De Preter K, Pattyn F, Poppe B, Van Roy N, De Paepe A, Speleman F. Accurate normalization of real-time quantitative RT-PCR data by geometric averaging of multiple internal control genes. Genome Biol 2002;3:RESEARCH0034.
16. Silver N, Best S, Jiang T, Thein SL. Selection of housekeeping genes for gene expression studies in human reticulocytes using real-time PCR. BMC Mol Biol. 2006;7:33.
17. Zhou L, Chen F, Yu J, Pan H. Selection of reliable reference genes for RT-qPCR analysis of Bursaphelenchus muckonatus gene expression from different habitats and developmental stages. Front Genet. 2018;9:269.
18. Xie F, Xiao P, Chen D, Xu L, Zhang B. mDeepFinder: a miRNA analysis tool for deep sequencing of plant small RNAs. Plant Mol Biol. 2012;80:75–84.
19. Smith AC, Swindle MM. Preparation of swine for the laboratory. ILAR J. 2006;47:358–63.
20. Gutierrez K, Dicks N, Glanzner WG, Agellon LB, Bordignon V. Efficacy of the porcine species in biomedical research. Front Genet. 2015;6:293.
21. Cox DF. Relation of litter size and other factors to the duration of gesta- tion in the pig. J Reprod Fertil. 1964;7:405–7.
22. Murray SAMJ, Jorgensen CV, Cirera S, Friedholm M. Selection of reference genes for gene expression studies in pig tissues using SYBR green qPCR. BMC Mol Biol. 2007;8:67.
23. Erkkens AC, Hekerman TW, de Vogel JW, van Haafsten B. Influence of litter size and breed on variation in length of gestation in the dog. Vet Q. 1993;15:160–1.
24. Biggers JD, Curnow RN, Finn CA, McLaren A. Regulation of the gestation period in mice. J Reprod Fertil. 1963;6:125–38.
25. Hoberman A, Lewis EM. Pediatric non-clinical drug testing principles, requirements, and practice. New Jersey: Wiley; 2012.
26. Hellemans J, Mortier G, De Paepe A, Speleman F, Vandesompele J. qBase relative quantification framework and software for management and auto- mated analysis of real-time quantitative PCR data. Genome Biol. 2007;8:R19.
27. Nygard AB, Jorgensen CB, Cirera S, Friedholm M. Selection of reference genes for gene expression studies in pig tissues using SYBR green qPCR. BMC Dev Biol. 2006;6:41.
28. Kuijk EW, du Puy L, van Tol HT, Haagsman HP, Collembander B, Roelen BA. Validation of reference genes for quantitative RT-PCR studies in porcine oocytes and preimplantation embryos. BMC Dev Biol. 2007;7:58.
29. Hildyard J, Wells D, Piercy R. Identification of qPCR reference genes suit- able for normalising gene expression in the developing mouse embryo. Wellcome Open Res. 2021;6:197.
30. Hildyard J, Wells D, Piercy R. Selection of housekeeping genes for the normalization RT-qPCR data from the intestine of minipig heart failure model and evaluation of TNF-alpha mRNA expres- sion. Biotechnol Lett. 2020;42:1007.
31. Wieczorek M, Abualrous ET, Sticht J, Alvaro-Benito M, Stolzenberg S, Noe F, et al. Mouse hereditary hypertensive strain: a porcine model for the study of hypertension. Front Genet. 2015;6:293.
32. Bevan MJ. The earliest knockouts. J Immunol. 2010;184:4585–6.
33. Nicot N, Hausmann JF, Hoffmann L, Evers D. Housekeeping gene selection for real-time RT-PCR normalization in potato during biotic and abiotic stress. J Exp Bot. 2005;56:2907–14.
34. Kohler M, Leitsch D, Muller N, Walochnick J. Validation of reference genes for the normalization of RT-qPCR gene expression in Acanthamoeba spp. Sci Rep. 2020;10:13062.
35. Wieczorek M, Abualrous ET, Sticht J, Alvaro-Benito M, Stolzenberg S, Noe F, et al. Validation of reference genes for gene expression studies in pig tissues using SYBR green qPCR. BMC Mol Biol. 2007;8:67.
36. Hildyard J, Wells D, Piercy R. Identification of qPCR reference genes suit- able for normalising gene expression in the developing mouse embryo. Wellcome Open Res. 2021;6:197.
40. Piorkowska K, Oczkowicz M, Rozycki M, Ropka-Molik K, Piestrzynska-Kajtoch A. Novel porcine housekeeping genes for real-time RT-PCR experiments normalization in adipose tissue: assessment of leptin mRNA quantity in different pig breeds. Meat Sci. 2011;87:191–5.
41. Percie du Sert N, Hurst V, Ahluwalia A, Alam S, Avey MT, Baker M, et al. The ARRIVE guidelines 2.0: updated guidelines for reporting animal research. BMC Vet Res. 2020;16:242.
42. Xu H, Zhong L, Deng J, Peng J, Dan H, Zeng X, et al. High expression of ACE2 receptor of 2019-nCoV on the epithelial cells of oral mucosa. Int J Oral Sci. 2020;12:8.
43. Zou X, Chen K, Zou J, Han P, Hao J, Han Z. Single-cell RNA-seq data analysis on the receptor ACE2 expression reveals the potential risk of different human organs vulnerable to 2019-nCoV infection. Front Med. 2020;14:185–92.
44. Hamming I, Timens W, Bulthuis ML, Lely AT, Navis G, van Goor H. Tissue distribution of ACE2 protein, the functional receptor for SARS coronavirus. A first step in understanding SARS pathogenesis. J Pathol. 2004;203:631–7.
45. Schmittgen TD, Livak KJ. Analyzing real-time PCR data by the comparative C(T) method. Nat Protoc. 2008;3:1101–8.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.