Reliable Reference Genes for Gene Expression Assessment in Tendon-Derived Cells under Inflammatory and Pro-Fibrotic/Healing Stimuli

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Abstract: Tendon cells (TCs) are important for homeostatic maintenance in the healthy tendon and to promote tissue healing after injury. Further, resident and rare populations of tendon stem/progenitor cells, located at various sites within the tendon, contribute to tendon recovery by differentiating into repairing TCs. Gene expression analysis, through quantitative reverse-transcription polymerase chain reaction (qRT-PCR), constitutes a useful tool to study cellular responses, including the transition from initial inflammation to healing processes. A critical step required for data normalization is the choice of reliable reference genes (RGs), a process highly underestimated in tendon biology. In this study, the suitability of five commonly used RGs (ACTB, B2M, GAPDH, HPRT1, and RPLP0) was evaluated using TCs samples cultured in both standard and progenitor-enriching conditions, as well as under either inflammatory (IFNγ + TNFα) or pro-fibrotic/healing (CTGF) stimulation. The stability of the candidate RGs was computationally determined using NormFinder, geNorm, BestKeeper, and DeltaCt applets. Overall, ACTB resulted as the most stable RG on the basis of the integration of each gene weight, whereas B2M and RPLP0 performed poorly. To further validate ACTB’s optimal performance, we evaluated the expression of ICAM1, coding for an immune-related cell surface glycoprotein, and COL1A1, encoding collagen type I that is the main component of the tendon extracellular matrix (ECM), both known to be modulated by inflammation. The expression of both genes was heavily affected by the RGs used. Consequently, when analyzing gene expression in tendon-derived cells subjected to various stimulatory protocols, the use of a suitable RG should be considered carefully. On the basis of our results, ACTB can be reliably used when analyzing different TC types exposed to pathological conditions.

Keywords: tendon; tenocyte; progenitor cell; inflammation; reference gene; qRT-PCR

1. Introduction

Tendinopathies can be described as a “continuum” of disturbances historically associated with progressive “wear and tear” of the tendon resulting from aging, overuse, and other causes [1]. As a well-established paradigm, while physiologic loads are required to maintain tendon homeostasis, abnormal
Cells 2019, 8, 1188 2 of 12

loading can lead to tendon injury, through either an acute trauma or a degenerative process resulting from an accumulation of micro-damage and an altered cell/matrix balance [2]. In this frame, pioneering works completed a roadmap of cellular and molecular cascades involved in tendon mechanobiology for both healthy and injured tissues [2]. In addition, more recently, immune/inflammatory responses have also been associated with tendon pathophysiology, complementing the original degenerative descriptions [3]. In fact, tendon injuries, along with mechanical stress, disturb the homeostatic balance between specialized tendon cells (TCs), named tenocytes, and immune cell compartments within the tissue [3]. Resident TCs account for a heterogeneous population of tenocytes, approximately 5% of the total tissue volume, with their main function related to the maintenance of tissue homeostasis [2]. A rare (<1%) subpopulation of the general TCs with stem-like properties has been described as tendon stem/progenitor cells (TSPCs) [4], although a large debate is still ongoing regarding a proper definition and characterization of this subset. Various strategies have helped elucidate their identification, including cell surface marker selection (e.g., CD146 positivity) [5] and culture conditions (e.g., hypoxia and PGE2 supplementation) [6]. Furthermore, the injection of connective tissue growth factor (CTGF) has been reported to enrich this population during early phases of tendon healing and promote tissue remodeling [5,7,8].

For both general TCs and specific TSPCs, a continuously expanding body of literature is starting to shed light on cellular and molecular mechanisms, aiming at deciphering their roles during acute/chronic injuries and healing responses [9,10]. Gene expression analysis, specifically qRT-PCR, has been used for decades as an informative tool to dissect such mechanisms, as it provides information regarding the different cellular responses to stimulatory environments. To consistently evaluate genes relative expression, the choice of reliable reference genes (RGs) is mandatory, as inappropriate RGs can lead to erroneous data and misleading experimental results [11]. An ideal RG must be stably expressed in all test samples, exhibiting low variability among them [12]. Unfortunately, it has become clear that the most common housekeeping genes used for many years for normalization do not meet those criteria, showing variations in the given conditions [13,14]. Therefore, it is necessary to carefully identify stable RGs to be tested and validated with different sets of cells and in multiple conditions.

To date, despite several RGs being considered “gold standards” and used in the analysis of tendons and derived cells, there is no consensus on reliable candidates. In a recent work, the suitability of six reference genes (18S, ACTB, B2M, GAPDH, HPRT1, and TBP) using samples from rotator cuff tendons was assessed, with HPRT1 resulting the most reliable [15]. In normal and diseased horse tendons, 12 commonly used RGs were analyzed, GAPDH being the most stable followed by ACTB [16]. Regarding human TCs treated with tenogenic supplements, YWHAZ and RPL13A showed superior consistency [17]. Even though these reports provide valuable information, their intrinsic distinct nature (i.e., different organisms, tissues and isolated cells, and the presence or absence of exogenous supplements) limits their use to describe a “universal” RG to study tendon cell biology, especially when dealing with its various cellular components.

For this reason, the aim of this work was to identify stable RGs in human tendon-derived cells cultured at both high and low densities, reminiscent of the described general TCs [18] and of enriched TSPCs, respectively, as the latter culturing condition has demonstrated to increase the expression of the progenitor marker OCT4 [19] Furthermore, in order to in vitro model various aspects of tendinopathy, those cells were exposed to either inflammatory (IFNγ + TNFα) or pro-fibrotic/healing (CTGF) stimulation. To obtain reliable candidates for the different cell types and distinct culture conditions, four computational gene expression analysis packages were used for the first time on tendon cells (geNorm, NormFinder, BestKeeper, and DeltaCt). The results obtained with this systematic approach will become a useful technical tool for future studies aimed at dissecting the molecular underpinnings of tendon biology and healing by reliably assessing gene expression.
2. Materials and Methods

2.1. Tendon Dissection and Cell Isolation

Human tendon cells were isolated from discarded fragments of the semitendinosus and gracilis tendons harvested from three de-identified patients (n = 3, males, 33 ± 9 years old) who underwent elective anterior cruciate ligament (ACL) reconstruction using hamstring tendons and provided their written informed consent (M-SPER-015- Ver. 2 - 04.11.2016 for the use of surgical waste material). The protocol was reviewed and approved by IRCCS Istituto Ortopedico Galeazzi IRB. After 16 h of enzymatic digestion with 0.3% w/v type I collagenase (185 U/mg, Worthington Biochemical Corporation, Lakewood, NJ, USA) [20], the samples were filtered through a 100 µm cell strainer (Becton, Dickinson and Co., NJ, USA) and centrifuged (300×g, 5 min). The resulting cells were plated at a density of either 5000 cells/cm² (high density, HD) or 50 cells/cm² (low density, LD) in low-glucose DMEM supplemented with 10% and 20% FBS, respectively, and maintained in an incubator at 37 °C in a humidified atmosphere with 5% CO₂. Cells at passage 2 were treated with either IFNγ + TNFα (10 ng/mL and 15 ng/mL) or CTGF (10 ng/mL) for 48 h before further processing.

2.2. Immunofluorescence/Immunohistochemistry Analysis

Cells were fixed with 10% neutral buffered formalin, washed with PBS, and then treated with a blocking solution containing 2% bovine serum albumin (BSA, Sigma-Aldrich, St. Louis, MA, USA). A primary antibody for CD146 (ab134065, Abcam, Cambridge, MA, USA) was diluted in 1% BSA (1:500) and incubated overnight at 4 °C with the cells. The samples were then washed three times with 1% Tris-buffered saline (Sigma-Aldrich) and incubated with the secondary antibody Goat anti-Rabbit IgG (H + L) highly cross-adsorbed, (Alexa Fluor Plus 647, 1:1000) for 1 h at room temperature. The cells were then washed and incubated with 0.1 µg/mL DAPI (DNA stain, Abcam) for 5 min.

The negative controls were subjected to overnight incubation at 4 °C with blocking buffer without the primary antibody. Microscope images were acquired using a Leica DMi8 microscope with Leica LAS X software (Leica, Heidelberg, Germany).

2.3. RNA Extraction and qRT-PCR Validation Studies

RNA was extracted using RNeasy Mini Kit (Qiagen, Frederick, MD, USA). RNA purity was assessed by 260/280 nm and 260/230 nm absorbance ratios and resulted to be 2.16 ± 0.01 and 1.98 ± 0.13, respectively. cDNA synthesis was performed using SuperScript™ VILO™ cDNA synthesis kit (Invitrogen, Carlsbad, CA, USA). qRT-PCR was performed using RT² SYBR Green qPCR Mastermix (Qiagen) reagent and transcript-specific primers (RT² qPCR primer assay for human ACTB (Product # PPH00073G), B2M (PPH01094E), COL1A1 (PPH01299F), GAPDH (PPH00150F), HPRT1 (PPH01018C), ICAM1 (PPH00640F), and RPLP0 (PPH21138F), Qiagen), following the manufacturer’s instructions. For each sample, independent qRT-PCR were performed using a StepOnePlus real-time PCR system thermocycler (Applied Biosystems, Thermo Fisher Scientific, Waltham, MA, USA). Amplification was obtained using the following cycling conditions: 10 min at 95 °C, followed by 40 cycles of 15 s at 95 °C and 1 min at 60 °C. As a quality control, we generated a first derivative dissociation curve for each well under the following conditions: 95 °C, 1 min; 65 °C, 2 min; 65 °C to 95 °C increasing by 2 °C/min. No more than one peak appeared in each reaction well, confirming amplification specificity. ACTB, B2M, GAPDH, HPRT1, and RPLP0 were analyzed as reference genes, and the most stable (ACTB) and the most variable (B2M and RPLP0) genes were used to score COL1A1 and ICAM1 modulation across samples with the ∆∆Ct (cycle threshold) method.

2.4. Data Analysis

RGs expression stability was estimated using four computational gene expression analysis packages: NormFinder, geNorm, BestKeeper, and DeltaCt. The raw Ct values were used directly for stability calculations in BestKeeper analysis and DeltaCt method and converted into relative quantities before
being imported into the geNorm and Norm-Finder applets. geNorm scores the average pairwise variation of an RG versus all other genes in the given samples [13]; NormFinder calculates the expression stability value based on inter- and intra-group variation [21]; the stability ranking of a candidate reference gene is determined by the CV (coefficient of variation) and SD (standard deviation) values in BestKeeper [22]; the DeltaCt method compares the relative expression of “pairs of genes” within each sample to confidently identify useful RGs [23]. Each algorithm generates the ranking of the RG according to their stability, assigning a series of continuous integers starting from 1. To improve the outcome, the geometric mean (geomean) of each gene weight generated by the four approaches was further calculated. The RG with the lowest geomean was considered to be the most stable.

Hierarchical clustering and principal component analysis (PCA) were obtained with the ClustVis webtool after row centering under the following settings for both rows’ and columns’ clustering distance and method: correlation and average, respectively [24].

2.5. Statistical Analysis

Statistical analysis was performed using GraphPad Prism v5.0 software (GraphPad Software Inc., La Jolla, CA, USA). COL1A1 and ICAM1 normal distribution of values was assayed by Kolmogorov–Smirnov normality test prior to a Student’s t-test. Differences were considered significant at p-value < 0.05.

3. Results

3.1. Tendon Cells Isolation and Characterization

Tendon-derived primary cells seeded at $5 \times 10^3$ cells/cm$^2$ (LD) showed isolated colonies with small cobblestone-like cells, while at $5 \times 10^3$ cells/cm$^2$ (HD), a homogeneous growth surface with fusiform and fibroblast-like morphology was visible (Figure 1A). These distinct cellular morphologies were complemented phenotypically with a clear difference in the expression of the surface antigen marker CD146. As expected, and on the basis of TSPCs enrichment in low-density conditions, LD cells exhibited a marked positivity for CD146 (87% ± 6 positive). Contrarily, HD conditions, reminiscent of heterogeneous TCs, allowed only a very low and rare CD146 expression (8% ± 3) (Figure 1B). These results reinforced the idea of selecting by culture at different cell densities two tendon-derived cell populations, supporting the subsequent gene expression examination.

![Figure 1](image-url). Tendon cells’ (TCs) morphology and CD146 expression. (A). Representative bright-field pictures of low-density (LD) and high-density (HD) TCs. (B). Immunofluorescence staining of LD and HD TCs for CD146 (red) and DAPI (blue). Negative control samples were subjected to overnight incubation at 4 °C with blocking buffer without the primary antibody.
3.2. Expression of Candidate RGs

Considering all TC samples, with and without IFNγ + TNFα or CTGF treatment, ACTB had the highest expression (lowest Ct values), whereas HPRT1 had the lowest (Figure 2 and Table 1). Setting a Ct cut-off of 25 for weak expression, with the only exception of HPRT1, all candidates RGs were detected as highly to moderately expressed. Moreover, to reduce the likelihood of including co-regulated RGs in the study, we performed a contiguity analysis, which evidenced that none of them resides within the same gene cluster. ACTB is located on chromosome (chr) 7, B2M on chr15, HPRT1 on chrX, and GAPDH and RPLP0 on chr12, spatially separated at the two extremities of the chromosome (position 6,534,517–6,538,371 for GAPDH and 120,196,699–120,201,211 for RPLP0; total chr12 length is 133,275,309 bp, with 998 genes, centrosome position 35.5 Mbp) and, therefore, unlikely to reside in the same cluster, usually encompassing less than 10 genes or, in rare cases, up to 100 [25].

Figure 2. Ct values of tested reference genes (RGs) across tendon cell samples.

Table 1. Ct values of tested RGs across tendon cell samples.

|                 | ACTB | B2M  | GAPDH | HPRT1 | RPLP0 |
|----------------|------|------|-------|-------|-------|
| TC 1 LD naive  | 15.04| 18.68| 16.78 | 26.07 | 19.47 |
| TC 2 LD naive  | 15.12| 18.41| 17.05 | 26.06 | 19.32 |
| TC 3 LD naive  | 15.58| 18.16| 17.19 | 26.20 | 19.22 |
| TC 1 LD IFNγ + TNFα | 15.94| 15.55| 17.34 | 26.53 | 22.96 |
| TC 2 LD IFNγ + TNFα | 15.12| 15.08| 17.32 | 26.39 | 22.47 |
| TC 3 LD IFNγ + TNFα | 14.68| 15.10| 16.66 | 26.12 | 22.27 |
| TC 1 LD CTGF   | 15.38| 19.82| 17.19 | 26.61 | 19.28 |
| TC 2 LD CTGF   | 14.64| 18.20| 15.73 | 26.73 | 18.45 |
| TC 3 LD CTGF   | 15.71| 18.48| 17.36 | 26.73 | 19.05 |
| TC 1 HD naive  | 15.46| 18.74| 16.99 | 25.64 | 19.08 |
| TC 2 HD naive  | 15.26| 18.99| 16.99 | 25.74 | 18.98 |
| TC 3 HD naive  | 15.26| 18.94| 16.82 | 25.47 | 19.67 |
| TC 1 HD IFNγ + TNFα | 14.40| 14.98| 16.59 | 25.70 | 22.65 |
| TC 2 HD IFNγ + TNFα | 14.41| 15.26| 16.80 | 25.88 | 22.12 |
| TC 3 HD IFNγ + TNFα | 14.61| 15.80| 16.54 | 25.91 | 22.62 |
| TC 1 HD CTGF   | 15.61| 18.75| 17.15 | 25.74 | 19.03 |
| TC 2 HD CTGF   | 15.08| 19.48| 17.05 | 25.97 | 18.79 |
| TC 3 HD CTGF   | 14.86| 18.80| 16.76 | 25.21 | 18.97 |

3.3. Stability Analysis of RGs

The stability rankings of the selected RGs in LD and HD TCs, under both resting (naive) conditions and IFNγ + TNFα or CTGF exposure is shown in Table 2. Each category represents the whole dataset under the indicated treatment or seeding density, namely, 27 samples for “ALL” and 9 samples for all the other groups. Since the used approaches generated few differences in outcomes, integration and normalization of the data were applied, generating the comprehensive geomean value. In LD, the best RG resulted to be ACTB (geomean of 1.32), and the least stable one was B2M (4.73), whereas considering HD cells, GAPDH performed very well (1), whereas RPLP0 showed great variability (4.73). Further, focusing on the culturing condition, in naive (non-primed) tendon cells, GAPDH (1) was the most
stable, and B2M (4.73) was the least stable. Under inflammatory conditions (IFNγ + TNFα), HPRT1 performed good (1.32), and B2M again scored last (3.98). When CTGF was supplied, RPLP0 (1.19) behaved as a reliable RG, and HPRT1 was the least reliable (5). Eventually, since both seeding densities and culturing conditions gave rise to slightly different outcomes, all samples were scored together. In this analysis, ACTB (1.32) resulted the best performer, and RPLP0 (4.73) the less reliable RG.

| Geomean | DeltaCt | BestKeeper | NormFinder | geNorm |
|---------|---------|------------|------------|--------|
| ACTB    | 1.32    | ACTB       | ACTB       | ACTB   |
| GAPDH   | 1.41    | GAPDH      | ACTB       | ACTB   |
| HPRT1   | 2.71    | HPRT1      | ACTB       | ACTB   |
| B2M     | 4.23    | RPLP0      | RPLP0      | RPLP0  |
| RPLP0   | 4.73    | RPLP0      | RPLP0      | RPLP0  |

For BestKeeper and DeltaCt SD, for NormFinder SV, and for geNorm M values are shown; geomean value is the geometric mean. The lower the value, the higher the stability. NB: geNorm applets released a couple of best RGs.

Table 2. Stability rankings of tested RGs.

Then, the samples were analyzed by PCA and hierarchical clustering (HC) to score major differences between populations (Figure 3). PCA showed that the expression pattern of the IFNγ + TNFα samples significantly differed, with all inflammation samples grouped apart. Further, HC confirmed the PCA data and emphasized how this outcome may be associated with the consistently higher B2M (low Ct values) and lower RPLP0 (high Ct values) levels under inflammation, as observed in Figure 3. Notably, with both approaches, CTGF supplementation was not able to influence TCs signature, which indicated the absence of major effects on transcriptional regulation of the selected RGs.
3.4. Impact of RGs Choice on the Expression Levels of Target Genes

qRT-PCR assays on ICAM1 and COL1A1 were performed in order to further evaluate the reliability of the selected candidate RGs in the TCs samples (LD and HD). ICAM1 and COL1A1 expression levels were normalized using the most stable (ACTB) and the two least stable (B2M and RPLP0) RGs. First, we evaluated the modulation of selected genes between the different seeding densities using LD values as a reference. On the basis of ACTB expression, we found statistical significant change (p-value = 0.0128 with ratio of 0.30) for ICAM1 in HD samples. For this condition, also the less reliable RGs showed statistically significant modulation (fold < 0.5, p-value < 0.05). When comparing the fold change given by the different RGs, no significant (ratio lower than 0.5 or higher than 2, p-value < 0.05) changes were found, meaning that with respect to ICAM1 expression in the absence of external stimuli, all three RGs behave with a similar trend. Regarding COL1A1, no differences (fold > 2, p-value < 0.05) were scored between both samples and the different RGs.

When cytokine supplementation was assessed, the choice of the RGs became crucial for IFNγ + TNFα samples, whereas again for CTGF, no differences were scored. Under inflammatory conditions, on the basis of ACTB expression, a marked increase of ICAM1 and a decrease of COL1A1 were observed in all seeding conditions (Figure 4A, Table 3). The gain of ICAM1 was maintained also when considering the least stable RG, although the use of B2M resulted in an underestimation and that of RPLP0 led to an overestimation of mRNA abundance, always significantly different from (ratio lower than 0.5 or higher than 2, p-value < 0.05 or < 0.07 for B2M with respect to ACTB in LD) and therefore erroneous for ACTB values. For COL1A1 (Figure 4B, Table 3), the effect of unreliable RGs was the opposite, with apparently higher downregulation observed in inflammatory conditions using B2M and almost an absence of modulation using RPLP0, being all these data again significantly different with respect to ACTB (with the only exception of B2M with respect to ACTB in HD cells, p-value of 0.06). Therefore, when IFNγ + TNFα were added to the culture medium, the effect of RGs choice gets decisive for the correct evaluation of target mRNA abundance, in term of both modulation and amount of variation.
ICAM1 (A) and COL1A1 (B) modulation under IFNγ + TNFα or connective tissue growth factor (CTGF) supplementation is dependent on the selected RGs. Increase in ICAM1 and decrease in COL1A1 expression with respect to untreated cells are significantly different depending on which RG (B2M or RPLP0) is used. Values are expressed as fold-change mean ± SEM, n = 3 independent cell isolates; ns: non-significant, * for p-value < 0.05, ** for p-value < 0.01, *** for p-value < 0.001.

Table 3. ICAM1 and COL1A1 modulation in single samples and their general patterns.

| ACTB | B2M | RPLP0 |
|------|-----|-------|
| ICAM1 | COL1A1 | ICAM1 | COL1A1 | ICAM1 | COL1A1 |
| TC 1 LD | 175.243 | 0.058 | 10.770 | 0.004 | 1061.283 | 0.353 |
| TC 2 LD | 53.996 | 0.105 | 5.438 | 0.011 | 480.558 | 0.929 |
| TC 3 LD | 78.577 | 0.072 | 17.617 | 0.016 | 1219.303 | 1.121 |
| TC LD | 102.605 | 0.079 | 11.275 | 0.010 | 920.382 | 0.801 |
| TC 1 HD | 175.755 | 0.065 | 27.045 | 0.010 | 4347.781 | 1.602 |
| TC 2 HD | 154.498 | 0.077 | 20.960 | 0.010 | 2445.748 | 1.221 |
| TC 3 HD | 368.332 | 0.149 | 65.354 | 0.027 | 4462.196 | 1.805 |
| TC HD | 232.862 | 0.097 | 37.786 | 0.016 | 3751.908 | 1.543 |
| TC 1 LD | 1.117 | 1.248 | 1.697 | 1.908 | 0.674 | 0.752 |
| TC 2 LD | 0.631 | 0.912 | 0.768 | 1.103 | 0.481 | 0.691 |
| TC 3 LD | 0.936 | 1.008 | 1.077 | 1.159 | 0.764 | 0.822 |
| TC LD | 0.895 | 1.056 | 1.181 | 1.390 | 0.640 | 0.755 |
| TC 1 HD | 0.948 | 1.122 | 0.859 | 1.016 | 0.827 | 0.978 |
| TC 2 HD | 0.824 | 0.985 | 1.307 | 1.562 | 0.816 | 0.975 |
| TC 3 HD | 0.866 | 1.032 | 1.041 | 1.250 | 0.704 | 0.839 |
| TC HD | 0.880 | 1.047 | 1.069 | 1.276 | 0.782 | 0.930 |

TC 1/2/3 LD or HD stand for the single cell isolates, whereas TC LD or HD stands for the general average, as per Figure 4A,B.

4. Discussion

Herein, a rigorous method to determine and validate RGs in tendon cells identified ACTB as the most stable gene, whereas B2M and RPLP0 performed poorly. Incorrect RGs choice affected the reliable evaluation of the expression of the target genes ICAM1 and COL1A1.

In this work, five candidate genes were selected for identifying the best reference among 18 TCs samples, grown in both standard (HD) and progenitor-promoting (LD) density seeding, under different experimental conditions, mimicking both inflammation (IFNγ + TNFα) and pro-fibrotic/healing phases (CTGF) in vitro. Four computational gene expression analysis packages (geNorm, NormFinder, BestKeeper and DeltaCt) were used to reliably determine the best candidate(s) for all conditions. The selection of the packages was based on their alternative approaches: i) the average pairwise variation of a reference gene versus all other genes among the given samples is scored by geNorm [13];
was normalized with respect to RPLP0 with the role of GAPDH. The results revealed that their expression pattern was significantly over- or underestimated when previously reported [5]. This may be due to both the di
time that RGs in tendon cells have a similar performance both in vitro and in vivo [15,16]. In this frame, optimal candidates out of a similar panel of candidates, therefore herein demonstrating for the first time that RGs in tendon cells have a similar performance both in vitro and in vivo [15,16]. In this frame, for isolated tenocytes, the only published report encompassed a di
time-dependent relationship for CTGF-dependent COL1A1. Notably, in our experimental setting, CTGF was not able to induce significant upregulation as 
ACTB, using ACTB as a RG, we were able to confirm ICAM1 increase and COL1A1 downregulation in both TCs and TSPC-enriched cultures. The results revealed that their expression pattern was significantly over- or underestimated when using the unstable genes for normalization (B2M/RPLP0), or even completely abolished when COL1A1 was normalized with respect to RPLP0, suggesting that the selection of appropriate, stable RGs is crucial for the correct normalization of qRT-PCR data in inflamed tenocytes. On the contrary, when CTGF was supplemented, no major RGs modulation, at least for the assayed candidates, was observed. Notably, in our experimental setting, CTGF was not able to induce significant COL1A1 upregulation as previously reported [5]. This may be due to both the difference in cytokine concentration, 10 ng/mL vs. 100 ng/mL, and the stimulation time, 2 days vs. 14 days. This suggests a concentration- and time-dependent relationship for CTGF-dependent COL1A1 increase. Further studies will be crucial to shed light on this molecular mechanism that, to date, has been poorly deciphered in tendon cells.
5. Conclusions

To the best of our knowledge, this study constitutes the first systematic evaluation of a set of candidate RGs as normalization factors in qRT-PCR analysis for tendon-derived cells, also under inflammatory- and pro-fibrotic/healing-like environments in vitro. The results of a comprehensive ranking order showed that ACTB displayed the highest stability across the set of samples in all conditions. The expression analysis of ICAM1 and COL1A1 emphasized the importance of suitable RGs to get accurate and reliable quantitation by qRT-PCR. These results are of great importance for further gene expression analyses in tendon-derived cells, facilitating the dissection of underlying molecular mechanisms related to tendon stress and healing responses. In fact, several molecular cascades have been postulated as critical, controlling local modulation by both TCs and TSPCs, and thus require detailed descriptions. In this frame, gene expression still constitutes an important assessment tool when dissecting molecular mechanisms. Therefore, the selection of reliable RGs for normalization in human tendon-derived cells warrants analytical accuracy, and the herein presented results are of crucial importance for future studies.

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