Quantitative PCR analysis of laryngeal muscle fiber types

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Abstract

Voice and swallowing dysfunction as a result of recurrent laryngeal nerve paralysis can be improved with vocal fold injections or laryngeal framework surgery. However, denervation atrophy can cause late-term clinical failure. A major determinant of skeletal muscle physiology is myosin heavy chain (MyHC) expression, and previous protein analyses have shown changes in laryngeal muscle fiber MyHC isoform with denervation. RNA analyses in this setting have not been performed, and understanding RNA levels will allow interventions better designed to reverse processes such as denervation in the future. Total RNA was extracted from bilateral rat thyroarytenoid (TA), posterior cricoarytenoid (PCA), and cricothyroid (CT) muscles in rats. Primers were designed using published MyHC isoform sequences. SYBR Green real-time reverse transcription-polymerase chain reaction (SYBR-RT-PCR) was used for quantification. The electropherogram showed a clear separation of total RNA to 28S and 18S subunits. Melting curves illustrated single peaks for all type MyHC primers. All MyHC isoforms were identified in all muscles with various degrees of expression. Quantitative PCR is a sensitive method to detect MyHC isoforms in laryngeal muscle. Isoform expression using mRNA analysis was similar to previous analyses but showed some important differences. This technique can be used to quantitatively assess response to interventions targeted to maintain muscle bulk after denervation.

Learning outcomes: (1) Readers will be able to describe the relationship between myosin heavy chain expression and muscle contractile properties. (2) Readers will be able to separate myosin heavy chain isoforms into slow and fast twitch phenotypes. (3) Readers will be able to describe differential muscle isoform expression between different laryngeal muscles. (4) Readers will be able to compare this study to other modalities of determining muscle fiber type.

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1. Introduction

Effective communication relies on intact voice and speech, and as such, laryngeal muscle activity has a dramatic impact on voice quality. Paralysis of one or more laryngeal muscles can lead to poor voice quality, breathing difficulties, or dysphagia. The most common causes for vocal fold paralysis include surgery in the head and neck divided into the following categories neck surgery not otherwise specified (18.8%), thoracic disease/surgery (15.5%), thyroid surgery (13.3%), and skull base surgery (12.1%). Systemic or peripheral neurologic (2.6%) or inflammatory disease (2.2%) makes up a minority of causes, but in many patients, a definitive cause is not determined (21.4%) (Netterville & Billante, 2003). Current treatment for laryngeal paralysis consists of surgical manipulation of the

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paralyzed vocal fold to place it into a better position to coapt the mobile vocal fold for phonation via injection laryngoplasty or laryngeal framework surgery. Such procedures provide adequate voice but additional treatments can be required over time to address the atrophy that occurs with lack of neural input (Andrews et al., 2008; Umeno, Chitose, Sato, & Nakashima, 2008).

In addition to paralysis, laryngeal incompetence related to aging is increasingly being recognized as a contributor to poor voice (Belafsky, Postma, Reulbach, Holland, & Koufman, 2002; Reulbach, Belafsky, Blalock, Koufman, & Postma, 2001). Treatment options for aging is similarly limited to static vocal fold augmentation with injection (Kwon, An, Ahn, Kim, & Sung, 2010) or framework surgery (Omori et al., 1997). Selected patients can have dramatic improvement in voicing after intervention, but there are no clear treatment guidelines (Behrman, 2004). More importantly, just as with intervention for paralysis, the intervention is symptomatic only. It does not address the underlying cause of the insufficiency. In animal models, age related changes have been shown to be a result of both muscle alterations and soft tissue degradation (Abdelkafy et al., 2007; McMullen & Andrade, 2005; McMullen & Andrade, 2009).

To expand clinical progress in this area, better understanding of the muscle physiology of the larynx is needed. There are traditionally, two different groups of muscles in the larynx. The adductor group which closes and tenses the glottis includes the intrinsic muscles of the larynx including the thyroarytenoid (TA), lateral cricoarytenoid (LCA) and interarytenoid (IA) as well as the intrinsic cricothyroid (CT) muscle. The sole abductor muscle that opens the glottis to allow for full respiration is the posterior cricoarytenoid (PCA). The predominant myosin heavy chain (MyHC) fiber type has a great deal of influence on the contraction speed, endurance, and strength of a given muscle fiber and therefore the muscle bundle. At least five types of MyHC (types I, IIA, IIB, IIX, and IIL; see Table 1) can be identified in the rat larynx by immunohistochemistry and SDS-PAGE (DelGaudio & Sciote, 1997; Li, Lehar, Nakagawa, Hoh, & Flint, 2004; Rhee, Lucas, & Hoh, 2004; Shiotani, Coleman, Alila, & Flint, 1998; Shiotani, Nakagawa, & Flint, 2001; Wu, Baker, Crumley, & Caiozzo, 2000; Wu, Baker, Marie, Crumley, & Caiozzo, 2004).

In mammals including both the rat and human, the TA muscle has very short contraction times thus has the fastest MyHC profile (in the range of extraocular muscles for same species) with no slow phenotype fibers identified (Hoh, 2005; Horton, Rosen, Close, & Sciote, 2008; Shiotani et al., 1998). In the rat, the CT muscle contraction times are two to four times longer than the TA similar to limb muscles. CT muscle therefore has smaller proportions of the fastest MyHC fibers (Hoh, 2005). In the rat, PCA muscle contraction times are intermediate between the TA and CT (Hoh, 2005; Shiotani et al., 1998). One study from the human has demonstrated that the three compartments of the PCA muscle (horizontal, oblique, vertical) may have different functions and speeds in that the fast MyHC higher in the vertical fibers of the PCA vs. horizontal fibers of the PCA (Horton et al., 2008).

Multiple immunochemistry, SDS-PAGE, and proteomic studies have shown denervation leads to detrimental changes in MyHC isoform expression (Caiozzo, Wu, Baker, & Crumley, 2004; DelGaudio & Sciote, 1997; Li, Lehar, Samlan, & Flint, 2005; Rhee et al., 2004; Shiotani & Flint, 1998). Immunochemistry and SDS-PAGE are at best semiquantitative techniques, however, that may over-represent predominant proteins and under-represent small amounts of protein. Quantification of RNA expression may be a more valuable technique to identify changes in MyHC expression (Jankala, Harjola, Petersen, & Harkonen, 1997). SYBR Green real-time RT-PCR RNA analysis demonstrates a high degree of sensitivity when applied to the expression of selected genes from small amounts and limited tissue samples such as rat intrinsic laryngeal muscle and human muscle biopsy samples (Kubota et al., 2003; Overbergh et al., 2003; Shariat et al., 2003). Although muscle fiber type is regulated by multiple mechanisms (Goldspink et al., 1992; Gunning & Hardeman, 1991), studies have suggested MyHC expression is predominately regulated transcriptionally. In particular, recently in young humans, MyHC protein levels were shown to be correlated

Table 1

<table>
<thead>
<tr>
<th>Isoform</th>
<th>Contraction rate</th>
<th>Primary cellular metabolism</th>
</tr>
</thead>
<tbody>
<tr>
<td>I (aka beta)</td>
<td>Slow</td>
<td>Oxidative (Red)</td>
</tr>
<tr>
<td>IIA</td>
<td>Fast</td>
<td>Mixed</td>
</tr>
<tr>
<td>IIB</td>
<td>Faster</td>
<td>Glycolytic (White)</td>
</tr>
<tr>
<td>IIX</td>
<td>Fastest</td>
<td>Glycolytic (White)</td>
</tr>
<tr>
<td>IIL (aka EO)</td>
<td>Fastest</td>
<td>Glycolytic (White)</td>
</tr>
</tbody>
</table>
with mRNA transcript levels (Short et al., 2005) which underscores the importance of better understanding the transcriptional regulation of fiber types.

The purpose of this study is to establish the use of quantitative PCR in laryngeal muscles. This study serves as a basis for additional studies investigating changes resulting in the larynx after denervation, radiation treatment, etc.

2. Materials and methods

This protocol was approved by the University of Iowa Institutional Animal Care and Use Committee (IACUC). Twelve Sprague–Dawley rats, approximately 280 g, experimental rats were used for this study. After euthanasia, the thyroarytenoid (TA), posterior cricoarytenoid (PCA), cricothyroid (CT), and soleus muscles were removed using a microscopic technique. Total RNA was extracted using a TRIZol® (Invitrogen) extraction protocol. Muscle fibers were loaded into 1.5 ml Eppendorf tubes and 1 ml TRIZol® reagent added then the fibers homogenized. Cell lysate was added to the pre-spun phase lock gel-heavy tubes and incubated 5 min at room temperature. Chloroform at 0.2 ml per 1 ml TRIZol® reagent initially used was added. Tubes were capped and shaken vigorously for 15 s. Samples were centrifuged at 12,000 × g for 10 min at 4 °C. The clear, aqueous phase containing RNA was transferred to a fresh Eppendorf tube. For precipitating RNA, 0.5 ml isopropyl alcohol was added. Samples were incubated at room temperature for 10 min and the centrifuged for 10 min at 12,000 × g, 4 °C. The supernatant was decanted from the RNA pellet on bottom of tubes. The RNA pellet was washed with ml 75% ethanol and centrifuged at 7500 × g, 4 °C. The supernatant was decanted from the carefully decanted supernatant was briefly air-dried for 5 min. DNase–RNase free water (60 μl) was added to dissolve RNA pellet then incubated at 55–60 °C for 10 min to facilitate dissolution. RNA concentration was determined using a spectrophotometer.

Various methods including spectrophotometry and gel electrophoresis have traditionally been used to analyze RNA quality. The bioanalyzer (Agilent Technologies 2100 Bioanalyzer, DNA core, University of Iowa) provides a very sensitive qualitative analysis of total RNA in small quantities compared to traditional methods. The samples were electrophoretically separated into peaks of 18S and 28S ribosomal RNA. Since the intrinsic laryngeal muscle samples were small (0.5–1 mg), the peaks in the bioanalyzer signal were difficult to resolve. To verify our method of RNA extraction, total RNA was extracted from the soleus muscle and diluted 1:10, 1:100 and 1:500 to the same concentration of our intrinsic laryngeal muscles samples.

Prior to RT-PCR, DNase treatment (Promega) was performed. Each reaction contained 1–8 μl RNA in water, 1 μl 10× buffer, 1 μl/μg RNA RNase-free DNase and DNase-RNase free water to a final volume of 10 μl. The samples were incubated at 37 °C for 30 min after which time 1 μl of stop solution was added to terminate the reaction and the samples incubated at 65 °C for 10 min to inactivate DNase. All samples were normalized to a common concentration.

Oligonucleotide primers were designed using NCBI published nucleotide sequences complementary to selected regions of the rat gene encoding MyHC with NCBI accession numbers detailed in Table 2. Since the original search was performed, regular NCBI automation has collapsed some of the sequences into accession numbers NM_001135158.1 and NM_001135157.1 (National Center for Biotechnology Information [NCBI]). Commonly used housekeeping genes for real-time quantitative PCR normalization include glyceraldehyde-3-phosphate dehydrogenase (GAPDH), 18S and 28S ribosomal RNA, β-actin, β-2-microglobulin. A recent report regarding the viability of using said genes as housekeeping genes recommended GAPDH as the most stable in response to exercise in slow- and fast-twitch human muscle fibers (Jemiolo & Trappe, 2004). Glyceraldehyde-3-phosphate dehydrogenase (GAPDH, TaqMan Rodent GAPDH) primer was purchased from Applied Biosystems to be used as a control housekeeping gene.

<table>
<thead>
<tr>
<th>Isoform</th>
<th>Forward primer</th>
<th>Reverse primer</th>
<th>Accession No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>GAA TGG CAA GAC GGT GAC TGT</td>
<td>GGA AGC GTA CCT CTC CTT GAG A</td>
<td>x 15939</td>
</tr>
<tr>
<td>IIA</td>
<td>ATG ACA ACT CCT CTC GTT TTG G</td>
<td>TTA AGC TGG AAA GTG ACC CGG</td>
<td>xm_340817</td>
</tr>
<tr>
<td>IIB</td>
<td>GAA CAC GAA GCG TGT CAT CCA</td>
<td>AGG TTT CGA TAT CTG CCG AGG</td>
<td>xm_340818</td>
</tr>
<tr>
<td>IIL</td>
<td>AAG AGA TGA TCT ACC AGG CCG A</td>
<td>GCC TGC CTC TTG TAG GAC TTC A</td>
<td>xm_340820</td>
</tr>
<tr>
<td>IIX</td>
<td>CCA ATG AGA CTA AGA CGC CTG G</td>
<td>GCT ATC GAT GAA TTG TCC CTC G</td>
<td>xm_213345</td>
</tr>
</tbody>
</table>
Superscript III reverse transcriptase (Invitrogen) was used to generate the first cDNA strand from RNA. A two-step RT was performed in a total reaction volume of 20 μl. The first reaction mix contained: 1 μl Random hexomer, 1 μl Oligo (dT)18, 1 μl 10 mM dNTP mix (Biosciences, Clontech), 8 μl extracted total RNA, and 2 μl DNase-RNase free water. The mixture was incubated 65 °C for 5 min, chilled on ice for at least 1 min, and added the reaction mixture of the second step which contained: 4 μl 5× First strand buffer, 1 μl 0.1 M DDT, 1 μl Superscript III RT, 1 μl RNaseOUT (Invitrogen). The final mixture was incubated at 25 °C for 10 min and then was incubated at 42 °C for 50 min, followed by reaction termination at 85 °C for 5 min. To remove RNA complementary to the cDNA1 μl of 2 U of Escherichia coli RNase H (New England, BioLabs) was added and incubated at 37 °C for 20 min then stored at −20 °C until use.

Quantitative PCR was performed using ABI-Prism qPCR machine and Platinum SYBR green qPCR Super Mix UDG (Invitrogen). Each reaction contained: 12.5 μl of Platinum SYBR green qPCR Super Mix-UDG, 0.5 μl ROX reference dye, 0.33 μl 100 μM forward and reverse primer, 5 μl cDNA and 6.34 μl DNase-RNase free water to a final volume of 25 μl. Amplification was placed in 96-well optical reaction plate and caps (ABI Prism). PCR parameters were as follows: 2-step cycling: 50 °C for 2 min and then initial denaturation at 95 °C for 2 min followed by 50 cycles of 15 s at 95 °C, and 30 s at 60 °C. Following the final cycle, melting curve analysis (dissociation curve) was performed for each run, and data from dissociation curves that showed evidence for primer-dimers or mirrored the negative control were eliminated (Vandesompele, De Paepe, & Speleman, 2002). The cycle number where the amplification curve crossed the threshold of the standard curve was noted as critical threshold (Ct) and used for all further analysis.

3. Results

To examine the quality of extracted RNA, a dilution (10×) of soleus muscle (50 mg) total extracted RNA using the aforementioned protocol showed 28S and 18S peaks with low noise between peaks as well as minimal low molecular weight contamination (Fig. 1A). The soleus total RNA was then further diluted (500×) to a low concentration at approximating the concentration of total RNA extracted from a laryngeal muscle. The soleus muscle showed a small peak of 28S and 18S ribosomal RNA again with minimal contamination (Fig. 1B). The 500× dilution sample then showed good results when run using the SYBR green RT-PCR (Fig. 2).

To confirm primer quality, the primers were subjected a dissociation analysis following the final PCR cycle. Fig. 3 illustrates a typical melting curve for MyHC type IIX PCR products with only a single dissociation peak at 86.5 °C. Dissociation of PCR constantly produced single peaks for MyHC type I at 86.2 °C, type IIA at 83.1 °C, type IIB at 85.5 °C, type IIL at 85.1, and the housekeeping gene (GAPDH) at 87.6 °C suggesting the presence of only one product of each PCR primer. Results were verified in triplicate on the same plates, and in duplicate on different plates for all isoforms of the PCA muscle.

Type I had a higher Ct value than other MyHC isoform in the TA and PCA muscles and type IIL had a higher Ct value than other MyHC isoform in CT muscle. The TA muscle showed expression of all MyHC isoforms with type I form showed relatively low levels of expression compared to isoform IIL due to the higher Ct values for type I compared to type IIL (Table 3). The PCA muscle showed expression of all MyHC isoform expression with all type II isoforms showing similar levels of expression to that of GAPDH, but type I was expressed at low levels compared to

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Fig. 1. Bioanalyzer output from total RNA extracted from soleus muscle diluted 10× (A) and 500× (B).
the type II isoforms as a whole (Table 3). The CT muscle showed expression of all MyHC isoform in a pattern opposite of TA (Table 3). The type I form showed relatively high levels of expression compared to isoform IIL due to the higher Ct values for type IIL compared to type I.

4. Discussion

While previous analyses have focused on SDS-PAGE and immunochemical analyses, this study establishes the ability to reliably extract high-quality myosin heavy chain (MyHC) mRNA from laryngeal muscles, the thyroarytenoid (TA), posterior cricoarytenoid (PCA), and cricothyroid (CT) in control animals. Similar to previous studies, isoform IIL expression was relatively higher in TA and PCA muscles compared to CT muscle. However unlike other studies, CT muscle did demonstrate low levels of IIL expression. Type I expression has previously only been reported in CT muscles without any in TA or PCA muscle. This ability to detect low levels of expression based upon reliable standard primers points to the improved sensitivity of the quantitative PCR technique. This finding that all MyHC isoforms are expressed in all laryngeal muscles has significance. It indicates the basic framework is present in all muscle and that transcriptionally based modulation is possible. If no isoform is being transcribed even at low levels, it may be more difficult to drive the cells to express that particular isoform with exogenous stimuli (either chemical, mechanical, or electrical).
This study is limited in that we obtained all muscle fibers contained in an individual muscle and analyzed them in bulk. Previous animal studies have shown three-dimensional differences in the dog thyroarytenoid muscle (Bergrin, Bicer, Lucas, & Reiser, 2006). The current study did not account for intra-muscular differences. However in the future, laser capture microdissection and single fiber analysis can overcome this limitation to more clearly define the molecular characteristics of this hybrid muscle (Wu et al., 2000a, 2000b). In particular, the sensitivity of this analysis will allow for more comprehensive evaluation of the natural state of these muscles as well as changes.

This study was performed in rats which limits its application to mouse models of muscular or systemic disorders. Rats were selected primarily because of the size of the larynx muscle fibers and extensive previous literature regarding muscle fiber type profiles. The underlying methods of this study can be easily applied to a mouse model by simply substituting mouse primers for rat. In addition to use in mouse models, it has become clear that microRNA regulation is important in muscle physiology and may possibly have a role in reinnervation (Williams et al., 2009). This study provides a crucial foundation to be able to study microRNA models in a way we have not been able to in the past.

The findings of this study have limited direct clinical utility. As stated in the introduction, the most direct application of this data is for future research in denervation associated with laryngeal paralysis or atrophy with aging. However, it is likely that the fibrosis and atrophy and resultant poor voice, swallowing, etc., associated with radiation treatment for head and neck cancers also have a molecular basis. Very little is known about the details of those changes (Lidegran, Forsgren, Dahlqvist, Franzen, & Domeij, 1999; Nagler, Baum, Miller, & Fox, 1998), but the techniques in this study are sensitive enough to first quantify those changes as well as measure the effect of interventions designed to reduce the effect. For instance, if inflammation plays a significant role, might treatment with anti-inflammatory agents limit long-term complications?

In summary, this study has demonstrated the ability to sensitively extract RNA from rat laryngeal muscle in a reproducible fashion. Application of this technique to future studies will allow better characterization of changes associated with disease.

Appendix A. Continuing education

1. The myosin heavy chain expression profile has an influence the following phenotypic characteristics of muscle contraction:
   a. Speed
   b. Endurance
c. Strength  
d. All of the above  

2. The myosin heavy chain isoform with the fastest contraction times is:  
a. I  
b. IIA  
c. IIX  
d. IIL/EO  

3. The thyroarytenoid muscle has the highest proportion of which of the following myosin heavy chain isoform:  
a. I  
b. IIA  
c. IIX  
d. IIL/EO  

4. Quantitative PCR is more sensitive than other methods in detecting _____ levels of RNA and protein expression.  
a. High  
b. Low  
c. Variable  
d. Stimulated  

5. Previous studies have shown muscle fiber type protein expression is _____ regulated.  
a. Transcriptionally  
b. Translationally  
c. Post-Translationally  
d. Cytoplasmically  

References  


