Effect of Controlled Muscle Activation in a Unilateral Vocal Fold Polyp Setting on Vocal Fold Vibration

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Abstract: Unilateral vocal fold polyps can lead to incomplete glottal closure and irregular vocal fold vibration. Depending on polyp size and resulting dysphonia severity, voice therapy or surgery may be recommended. As part of voice therapy, patients may learn how to optimize intrinsic and extrinsic laryngeal muscle use to mitigate benign lesion effects, increase vocal efficiency, and improve voice quality. In this study, we used a low-dimensional mass model with a simulated unilateral vocal fold polyp and varied intra-laryngeal muscle activity to simulate vocal fold vibration across varied conditions. Differing muscle activation has different effects on frequency, periodicity, and intensity. Accordingly, learning how to optimize muscle activity in a unilateral polyp setting may help patients achieve the best possible periodic and most efficiently produced voice in the context of abnormal vocal fold morphology.

Keywords: vocal fold polyp; vocal fold vibration; vocal fold vibration simulation; vocal fold vibration model; aeroacoustics; larynx

1. Introduction

Vocal fold polyps are benign mass lesions arising from phonotrauma [1]. They may be hemorrhagic or gelatinous and are typically unilateral. Microscopically, vocal fold polyps exhibit fibrosis, fibrin deposition, hemorrhage, vascular thrombosis, and dilatation of blood vessels [2]. Ongoing phonotraumatic behavior will result in repetitive injury to the true vocal fold with potential for resulting dysphonia, reduced vocal range, odynophonia, and occasionally risk of scar [1]. In addition, vocal fold polyps have the potential to become cancerous. It is estimated that there will be 12,470 new cases of laryngeal carcinoma and 3820 deaths from laryngeal carcinoma in the USA in 2022. Laryngeal carcinoma is one of the most common forms of head and neck cancer, with an estimated number of 12,370 new cases and 3750 deaths in the USA in 2020 [3]. Accordingly, timely treatment of vocal fold polyps is necessary. Depending on polyp characteristics, patient vocal demands, patient preference, and degree of dysphonia, treatment may include voice therapy and/or surgery [4]. However, this type of surgery has some technical difficulties and may lead to post-operative complications [5,6]. In either case, voice therapy is useful in helping the patient identify less phonotraumatic methods of voice use to increase vocal efficiency, improve vocal quality, decrease stress on the true vocal folds, and decrease probability of recurrence if procedural intervention is pursued. Identifying optimal intra-laryngeal muscle use patterns may be a part of therapy. Computational modeling of vocal fold vibration offers the opportunity to manipulate degree of muscle activity and examine effects on vibratory characteristics and acoustic output. It could be used to better understand pathological vocal fold vibration and potentially guide therapy [7,8].

There are multiple recent studies on pathological vocal fold vibration using vocal fold models. The first low-dimensional mass model of the vocal fold was developed by Ishizaka...
and Flanagan [9] and later modified by Steinecke and Herzel [10]. Much work has been
done to couple various flow models to these two-mass models to better match physiological
data. For this study, two aspects of previous models were of particular interest. One was
the use of a nonlinear model to study chaotic vibration in the unilateral polyp setting,
first proposed by Zhang et al. [11]. Nonlinear dynamic methods can be used to describe
disordered voice production when vocal fold vibration is aperiodic. The influence of
tension asymmetry on phonation onset, intensity, and fundamental frequency has been
experimentally observed and simulated with this nonlinear model by Xue [12]. Mehta [13]
used high-speed videoendoscopy and modeling to investigate the phase asymmetry of
vibration and found that vibratory asymmetries have no direct relationship with acoustic
spectral tilt measures.

The second relevant aspect is intrinsic laryngeal muscle activation during phonation. The
intrinsic muscles interconnect the cartilages of the larynx, such as the thyroarytenoid muscle,
cricoarytenoid muscle, lateral cricoarytenoid muscle, and posterior cricoarytenoid muscle.
Previous studies have shown direct correlations between intrinsic muscle activation and
characteristics of sound production [14–16]. Chhetri et al. [14] showed the influence of various
combinations of thyroarytenoid muscle, lateral cricoarytenoid muscle, and cricothyroid muscle
activation on acoustic output in an in vivo canine model. Dewan et al. [15] also used an
in vivo canine model to assess possible neuromuscular compensation mechanisms that could
potentially be used in the treatment of vocal fold pathology. Azar et al. [16] evaluated
combinations of superior and recurrent laryngeal nerve activation to explore fundamental
relations between nerve activation and laryngeal muscle activation during phonation.

There are many differences between normal vocal fold and pathological vocal fold
vibration. The natural frequencies of the vocal fold entail the vibration of the two lips of vocal
fold to the same frequency on phonation [17,18]. It is not clear if such entrainment occurs
when a unilateral polyp is present, nor if the whole vocal fold will have the same effect when
a vocal polyp is present. For a pathological vocal fold, the natural frequency differs from the
normal one. Vocal polyps may affect the vocal fold vibrations and flow dynamics.

Prior studies have focused more on the manipulation of normal mode frequencies,
effective mass in vibration, vocal fold length, stiffness, and vocal fold thickness, which
are not directly controlled by the speaker [19]. Greater attention to muscle activation may
yield new insights into the physiology of disordered voice production. Current modeling
guidelines for vocal fold muscle activation are mostly based on normal vibration. Titze [19]
described an empirically derived set of rules that converts laryngeal muscle activities into
physical quantities and used these rules to investigate a shear mode and two compressional
modes of vibration in a self-oscillating model of vibration. Zhang [20] investigated the
implication of stiffness on fundamental frequency (F0) and glottal flow rate through a
three-dimensional model. Evidence for a direct correlation between muscle activation and
vocal fold vibration in this model, though, is still limited.

To better understand the effect of unilateral polyps on vibration, we have concentrated
more on muscle activation in a computational pathological vocal fold model. This allows
for the study of vocal fold oscillatory characteristics in perceptually and physiologically
realistic controlled spaces. Compared to animal models, computational models based
on human vocal folds are more customizable and generalizable. We have developed a
vocal fold model with a unilateral polyp to simulate irregular vocal fold vibrations. We
could also study aerodynamics in this asymmetric configuration, which obeys a set of rules
that transform muscle activation into geometrical and viscoelastic parameters of lumped
element models. Our pathological model may shed light on the effect of muscle activation
on vocal fold vibration to help patients learn how to optimize muscle activity for different
phonation situations.
2. Materials and Methods

2.1. Vocal Fold Model with a Unilateral Polyp

The two-mass model is a classic model for vocal fold vibration. The model used here is based on Steinecke and Herzl [18]. Their model regards each vocal fold as two masses that are connected by springs. In our study, we added a unilateral mass representing a polyp to this model, following previous work [11].

Figure 1 shows a schematic of the three-mass model in which the indices $a = l, r$ denote the left and right vocal fold and $i, j = 1, 2$ are the upper and lower masses or vocal fold lips. $x_{ia}$ is the displacement of the mass $m_i$ with length $d_i$. $k_{ia}$ is the stiffness constant, and $k_{ca}$ is the coupling stiffness between $m_{1a}$ and $m_{2a}$. $r_{ia}$ is the damping constant. The mass $m_{1a}$ and $m_{2a}$ has length $l_0$ and depth $D_0$. The height of $m_{1a}$ is $d_1$, and $d_2$ is the height of $m_{2a}$. The height of the vocal fold is $d_0$, $d_0 = d_1 + d_2$.

![Figure 1. Schematic of vocal fold model with a polyp. Vertical plane represents midline.](image)

A unilateral vocal polyp is connected with the lower mass $m_{1r}$ of the right vocal fold by the spring $k_{3r}$ and the damper $r_{3r}$. The vocal polyp has mass $m_{3r}$, length $l_3$, depth $D_3$, and height $d_3$. $x_{3r}$ is the displacement of the polyp $m_{3r}$. In this model, the vocal folds and polyp have the same density. Collisions occurs between $m_{1l}$ and $m_{1r}$, $m_{1l}$ and $m_{3r}$, and $m_{2l}$ and $m_{2r}$. $c_{ia}$ is the spring constant, $c_{ia} = 3k_{ia}$. All masses are only allowed to move in the medio-lateral direction.

Using these parameters, we first began to develop muscle activation rules for the pathological model.

2.2. Development of Rules for Muscle Control

In reality, the value of some parameters will vary with the movement of the laryngeal muscles during phonation, which is called the effective value of vocal fold vibration [21]. For a more realistic simulation, we improved the rules for controlling geometric and elastic parameters with muscle activation proposed by Titze [19] to fit our model. These rules take the place of the physics of deformable bodies for vocal fold posturing. To reduce the number of control parameters, only the cricothyroid muscle activity $a_{CT}$ (ranging from 0.05 to 0.9), the thyroarytenoid muscle activity $a_{TA}$ (same range), and the lateral cricoarytenoid muscle activity $a_{LC}$ (same range) were considered. The reason for setting the different muscle activities in this range is that the muscle cannot actually relax and contract completely. The length of the vocal fold is the primary geometric variable for adjustment of $F_0$, and it changes during phonation. It is therefore important to identify what is meant by effective
vocal fold length. Titze [22] showed that the fractional length change (vocal fold strain) due to rotation can be written as

$$\varepsilon = G(Ra_{CT} - a_{TA}) - Ha_{LC},$$

(1)

where \(\varepsilon\) is the longitudinal vocal fold strain, \(G\) is a gain factor, \(R\) is a torque ratio, and \(H\) is the adductory strain factor. The range of \(\varepsilon\) for human must be higher to achieve a larger pitch range than canines. To obtain a larger maximum elongation, we needed to increase the values of the above parameters (\(G, R, H\)) compared to the canine model. For this study, we set \(G = 0.2, R = 3.0,\) and \(H = 0.2,\) as in a previous study [22]. When a polyp develops on one vocal fold, it can be approximated that the length and tension of the vocal folds remain constant, while the vocal fold linear density \(\rho\) increases slightly in the local area around the polyp [21]. In the present model, we treated the entire vocal fold cover as one with the polyp but still follow this formula. Using the above rule, the vocal fold length can be expressed in the form

$$l = l_0(1 + \varepsilon) = l_0[1 + G(Ra_{CT} - a_{TA}) - Ha_{LC}],$$

(2)

where \(l_0\) is the resting length, for which the average value in humans is 1.4 cm.

Vocal fold depth increases with vocal fold shortening during vocal fold vibration. We developed the following rules for thickness:

$$D = D_{muc} + 0.5D_{lig} + 0.2\varepsilon$$

(3)

where \(D_{muc}\), standing for depth of mucosa is 0.2 cm, and \(D_{lig}\), standing for depth of ligament, is 0.2 cm. Thickness is described as

$$d = \frac{d_0}{1 + 0.8\varepsilon}$$

(4)

where \(d_0\) is the vibrating thickness at resting length. We defined the ratio of the effective vibration parameter to the original parameter as the effective vibration ratio \(\omega\). The effective vibration ratio of length \(\omega_l\), depth \(\omega_D\), and thickness \(\omega_d\) can be written as

$$\omega_l = l/l_0 \quad \omega_D = D/D_0 \quad \omega_d = d/d_0$$

(5)

In addition to the above parameters, the stresses acting on the vocal fold also change with muscle activation during vibration. The cover layer has only a passive stress and can be expressed as

$$\sigma_c = \left[\sigma_{muc}D_{muc} + 0.5\sigma_{lig}D_{lig}\right]/D$$

(6)

where \(\sigma_{muc}\) is the mucosa stress, and \(\sigma_{lig}\) is the ligament stress. According to previous work [23,24], the passive stress of mucosa and ligament has been modeled with a combination of a linear and an exponential function and matched to vocal fold stress-strain curves. The form of this function can be expressed in the form

$$\sigma_p = \begin{cases} 0 & \text{for } \varepsilon < \varepsilon_1 \\ -\sigma_0/\varepsilon - \varepsilon_1 & \text{for } \varepsilon_1 \leq \varepsilon \leq \varepsilon_2 \\ [\sigma_0/\varepsilon_1(\varepsilon - \varepsilon_1)] + \sigma_2[e^{c(\varepsilon - \varepsilon_2)} - C(\varepsilon - \varepsilon_2) - 1] & \text{for } \varepsilon < \varepsilon_2 \end{cases}$$

(7)

\(\sigma_0\) is the stress when \(\varepsilon = 0\), \(\varepsilon_1\) is the strain where the linear portion goes to zero, \(\varepsilon_2\) is a scale factor for the exponential portion, and \(\varepsilon_2\) is the strain where the exponential portion begins. The numerical constants used in Equation (7) for mucosa are \(\varepsilon_1 = -0.5, \varepsilon_2 = 0.35, \sigma_0 = 0.5\) kPa, \(\sigma_2 = 30\) kPa, \(C = 4.4\); for ligament are \(\varepsilon_1 = -0.5, \varepsilon_2 = 0, \sigma_0 = 0.4\) kPa,
\( \sigma_2 = 1.393 \text{ kPa}, \ C = 17; \) and for TA muscle are: \( \varepsilon_1 = -0.5, \ \varepsilon_2 = -0.05, \ \sigma_0 = 1.0 \text{ kPa}, \ \sigma_2 = 1.5 \text{ kPa}, \ C = 6.5. \) Hence, the effective vibration ratio of cover stress is

\[ \omega_{\sigma} = \sigma_c / \sigma_0 \] (8)

where \( \sigma_c \) is stress on the cover when the vocal folds are at rest. The above rules consider the polyp and vocal fold as a whole.

2.3. Effective Parameters of Vocal Fold Vibration

Muscle activation affects the mass and stiffness by changing the above parameters, which have an impact on the vibration \([19]\). In our low-dimensional model, the mass of the vocal folds can be calculated by

\[ m = \rho ldD \] (9)

Changes in muscle activation do not affect density of the vocal fold. Therefore, we only consider the effective length, depth, and width of the vibration, such that the effective mass of the vocal fold now become the following:

\[ m_{ia} = (\omega_D \omega_d \omega_l) \cdot m_{ia}m_{3r} = (\omega_D \omega_d \omega_l) \cdot m_{3r} \] (10)

where \( m_{ia} \) and \( m_{3r} \) on the right side of the equation are the default values of vocal fold mass.

For vocal fold stiffness, the following relationship is given by the definition of fundamental frequencies:

\[ k = \pi^2 \sigma dD / l \] (11)

Likewise, for our model, we can similarly write,

\[ k_{ia} = (\omega_D \omega_d \omega_l) \cdot k_{ia}k_{3r} = (\omega_D \omega_d \omega_l) \cdot k_{3r} \] (12)

where \( k_{ia} \) and \( k_{3r} \) on the right side of the equation are the vocal fold stiffness when no vibration occurs. In addition, the coupling stiffness \( k_{ca} \) is different from others. According to the mass distribution rule of a previous study \([19]\), the effective coupling stiffness becomes

\[ k_{ca} = \left[ \left( \frac{18lD_c}{7d} - \frac{2ld}{D} \right) / \left( \frac{18lD_0}{7d_0} - \frac{2ld_0}{D_0} \right) \right] \cdot k_{ca} \] (13)

The default parameters in the deterministic vocal fold models with steady airflow used in Table 1 are considered as common physiological conditions \([19,25]\). The units of parameters are cm, g, ms, and their corresponding combinations.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Symbol</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lower mass</td>
<td>( m_{1a} )</td>
<td>0.125</td>
</tr>
<tr>
<td>Upper mass</td>
<td>( m_{2a} )</td>
<td>0.05</td>
</tr>
<tr>
<td>Lower stiffness constant</td>
<td>( k_{1a} )</td>
<td>0.08</td>
</tr>
<tr>
<td>Upper stiffness constant</td>
<td>( k_{2a} )</td>
<td>0.008</td>
</tr>
<tr>
<td>Coupling stiffness constant</td>
<td>( k_{ca} )</td>
<td>0.025</td>
</tr>
<tr>
<td>Lower mass thickness</td>
<td>( d_1 )</td>
<td>0.25</td>
</tr>
<tr>
<td>Upper mass thickness</td>
<td>( d_2 )</td>
<td>0.05</td>
</tr>
<tr>
<td>Thickness of the vocal fold</td>
<td>( d_0 )</td>
<td>0.3</td>
</tr>
<tr>
<td>Length of the vocal fold</td>
<td>( l_0 )</td>
<td>1.4</td>
</tr>
<tr>
<td>Depth of the vocal fold</td>
<td>( D_0 )</td>
<td>0.3</td>
</tr>
<tr>
<td>Lower mass spring constant during vocal fold collisions</td>
<td>( c_{1a} )</td>
<td>0.24</td>
</tr>
<tr>
<td>Upper mass spring constant during vocal fold collisions</td>
<td>( c_{2a} )</td>
<td>0.024</td>
</tr>
<tr>
<td>Air density</td>
<td>( \rho )</td>
<td>0.00113</td>
</tr>
<tr>
<td>Subglottal pressure</td>
<td>( P_s )</td>
<td>0.006</td>
</tr>
</tbody>
</table>
The damping constant of the vocal folds can be calculated as

\[
\begin{align*}
r_{1a} &= 2\zeta_1 \sqrt{m_1k_1} = 0.1 \\
r_{2a} &= 2\zeta_2 \sqrt{m_2k_2} = 0.6
\end{align*}
\]

where \(\zeta_1, \zeta_2\) refer to damping ratios for the viscous resistances \(r_{1a}\) and \(r_{2a}\). Based on previous studies [11], we selected the following parameters for the vocal fold polyp: \(m_{3r} = m_{2a}l_3D_3/lD_0d_1\), \(k_{3r} = 0.014\), \(c_{3r} = 3k_{3r}\), \(r_{3r} = r_{3r}(m_{3r}k_{3r}/m_{1r}k_{1r})^{0.5}\), \(l_3 = l_0/3\), \(D_3 = 0.1D_0\), \(d_3 = (d_1 + d_2)/3\).

### 2.4. Equations of Motion for the Three-Mass Model

The differential equations for the pathological model are

\[
m_{ia}\ddot{x}_{ia} + r_{ia}\dot{x}_{ia} + k_{ia}x_{ia} + \Theta(\text{a}_i)c_{ia}\frac{a_i}{2l_i} + k_{ca}(x_{ia} - x_{ja}) + T_{ia} = F_{ia},
\]

and for the equation of the right vocal fold polyp,

\[
m_{3r}x_{3r} - T_{1r} + T_{1l} = F_3
\]

The function \(\Theta(x)\) describing the collision is approximated by

\[
\Theta(x) = \begin{cases} 
\tanh\left(\frac{30x}{50}\right) & \text{for } x > 0 \\
0 & \text{for } x \leq 0 
\end{cases}
\]

The polyp \(m_{3r}\) and the right lower mass \(m_{1r}\) are connected by nonlinear spring and the damping, which can be described as

\[
T_{1r} = r_{3r}(\dot{x}_{1r} - \dot{x}_{3r}) + k_{3r}[x_{1r} - x_{3r} + \eta_{k3}(x_{1r} - x_{3r})^3],
\]

while the effective nonlinear spring forces of the left lower mass \(m_{1l}\) can be written as

\[
T_{1l} = \Theta(-a_3)c_{3r}\left[\frac{a_3}{2\omega l_3} + \eta_{k3}\left(\frac{a_3}{2\omega l_3}\right)^3\right],
\]

for which the parameters were set as \(\eta_{k3} = 100\) and \(\eta_{k3} = 500\). There is no coupling or collision effect between the right vocal fold polyp and the upper mass, so \(T_{2l} = 0\) and \(T_{2r} = 0\).

To obtain the glottal airflow, it is necessary to calculate the minimal glottal area, using

\[
a_{min} = \min(a_{1l}, a_{2l}) + \min(a_{1r}, a_{2r}, a_{3r}'),
\]

and the glottal areas of the lower mass pair \(m_{1a}\) and the upper mass pair \(m_{2a}\), using

\[
a_l = a_{0l} + l_0(x_{1l} + x_{1r}) = a_{il} + a_{ir}
\]

The area from the midline to \(m_{ia}\) is \(a_{ia}\), and \(a_{01} = a_{02} = 0.05\). Similarly, the equation of glottal area for the polyp can be written as:

\[
a_{3r}' = a_{01}/2 - l_3D_3 + (l_0 - l_3)x_{1r} + l_3x_{3r}
\]

\[
a_3 = a_{03} + l_3(x_{1l} + x_{3r})
\]

\[
a_{03} = l_3(a_{01}/l_0 - D_3)
\]

\[
a_{3r}' = a_{01}/2 + l_0x_{1l} + a_{3r}'.
\]
The volume flow velocity $U$ can be described by the Bernoulli equation as

$$P_1 = P_s[1 - \Theta(a_{min})(\frac{a_{min}}{a_1})^2] \Theta(a_1)$$

$$P_2 = 0$$

$$P_3 = P_s[1 - \Theta(a_{min})(\frac{a_{min}}{a_3})^2] \Theta(a_3)$$

$$U = \sqrt{\frac{2P_s}{\rho} a_{min} \Theta(a_{min})}$$

The effective driving forces $F_{1a}$ acting on $m_{1a}$, $F_{2a}$ acting on $m_{2a}$, and $F_{3r}$ acting on $m_{3r}$ can thus be deduced as

$$F_{1a} = (P_1 l_0 (d_1 - d_3) + P_3 l_0 d_3) \cdot (\omega_d \omega_1), F_{2a} = 0$$

$$F_{1r} = (P_1 l_0 (d_1 - d_3) + P_3 (l_0 - l_3) d_3) \cdot (\omega_d \omega_1), F_{2r} = 0$$

$$F_{3r} = P_3 l_3 d_3 \Theta(a_3) \cdot (\omega_d \omega_1).$$

The equations in this section give the vibratory equations of the vocal folds coupled with a right vocal polyp, while the vocal fold model with a unilateral polyp is reducible to the normal two-mass model when $D_3 = 0, l_3 = 0, d_3 = 0$, and $T_{3a} = 0$.

### 3. Results and Discussion

#### 3.1. Qualitative Features of Pathological Vocal Fold

The glottal volume flow velocity $U$ of the normal and pathological vocal folds is shown in Figure 2. As can be seen from the figure, the mass block displacement in both normal and pathological vocal fold models vibrates periodically and produces a valley-cut sinusoidal shape for airflow. However, the amplitude of volume flow velocity $U$ decreases when the polyp is present. In the closing phase of the whole cycle, the pathological vocal fold can not close entirely, and the duration of the closed phase is reduced. As we can see, vocal fold polyps have an impact on the quality and intensity of phonation. Hence, the study focused on simulating muscle activations that may reduce the adverse effect of polyps on phonation.

Previous work has identified that for $a_{LC} > 0.5$, the vocal processes are approximated, and for $a_{LC} < 0.5$, the vocal processes are separated. Oscillation occurs approximately along the constant horizontal line of $a_{LC} = 0.5$ [19]. Therefore, we fixed $a_{LC}$ at 0.5 to better understand the implication of muscle activation on adduction and abduction. When $a_{CT} = 0.25, a_{TA} = 0.25$, and $a_{LC} = 0.5$, the longitudinal strain of the vocal fold $\varepsilon$ will be 0, and all laryngeal muscles are activated. Hence, fixing $a_{CT}$ at 0.25 for the study of $a_{TA}$ was seen as the best choice. The same method used for fixing $a_{TA}$ was used for the study of $a_{CT}$.

Figure 3 shows glottal volume flow velocity waveforms of different cricothyroid muscle activities ($a_{CT}$). $a_{CT}$ now increases from 0.05 to 0.9 with $a_{TA}$ constant at 0.25 and $a_{LC}$ constant at 0.5. Note that the effect of increasing $a_{CT}$ is an initial increase in fundamental frequency. The average amplitude of the glottal airflow first increases and then decreases with increasing $a_{CT}$. It can be seen the vocal polyp cannot change the status of the glottal cycle on phonation. However, the figure shows that the vocal folds can maintain a more periodic vibration only when $a_{CT}$ is at a moderate value. When the level of CT muscle activity is too low, there is low stiffness of the vocal fold cover, causing the vocal fold to vibrate with little speed and amplitude. However, when $a_{CT}$ increases, the effect of the polyp on vibration becomes obvious, and there is abnormal glottal airflow. As the stiffness increases with $a_{CT}$ to the maximum of the range, the stabilized waveforms of glottal airflow tend to be a straight line, and vibration is severely impaired.
Figure 2. Glottal volume flow velocity rate. (a) The normal vocal fold. (b) The vocal fold with a unilateral polyp.

Figure 3. Glottal volume flow velocity $U$ for a unilateral polyp vocal fold model across differing levels of cricothyroid muscle activity, $a_{CT}$. In each case, $a_{TA}$ was fixed at 0.25 and $a_{LC}$ was fixed at 0.5.

Figure 4 shows glottal volume flow velocity waveforms for different levels of thyroarytenoid muscle activity ($a_{TA}$). The waveform is nearly sinusoidal at small values of $a_{TA}$ but decreases in amplitude when $a_{TA}$ increases. The waveforms in Figures 3a and 4d are similar, showing the effect of low stiffness on vocal fold vibration. The fundamental frequency slightly decreases with increasing values of $a_{TA}$. This supports the function of the thyroarytenoid and cricothyroid muscles in controlling fundamental frequency [21].
Figure 4. Glottal volume flow velocity $U$ for a unilateral polyp vocal fold model with different $a_{TA}$. In each case, $a_{CT}$ was fixed at 0.25 and $a_{LC}$ was fixed at 0.5.

Figure 5 shows glottal volume flow velocity waveforms for different levels of lateral cricoarytenoid muscle activity ($a_{LC}$). There is a decrease in glottal airflow amplitude with increasing $a_{LC}$. Unlike for the thyroarytenoid muscle, the glottal airflow waveform is almost sinusoidal when $a_{LC}$ is around 0.5. Irregular vocal fold vibration occurs with increasing or decreasing $a_{LC}$. Furthermore, there is a glitch in the glottal airflow waveforms for large values of $a_{LC}$. This indicates that in the vocal fold polyp setting, high levels of lateral cricoarytenoid activity affects voice quality. Of note, increasing $a_{LC}$ can also result in a small decrease in $F_0$.

Evaluating the above three sets of waveforms, we obtain an initial understanding of the muscle activation ranges that will optimize voice production in the polyp setting. Cricothyroid activity should be greater than 0 but less than 0.5, and both $a_{TA}$ and $a_{LC}$ should be kept at a moderate value. Under these conditions, periodic vocal fold vibration can occur.

3.2. Glottal Flow and Implications for Phonation

Phonation is essentially an energy-transfer process, transforming aerodynamic energy into acoustic energy. Because the glottal flow is the main determinant of voice production, further studying its properties can help us learn the effect of muscle activation on
onset, effort, and quality. Following the experimental method in III.A and referring to the experimental design from Xue [12], we plotted the variation of four airflow parameters affecting sound quality with respect to different levels of muscle activation (Figure 6). Figure 6a shows the root-mean-square fluctuation of the airflow velocity ($U_A$). We defined the intensity of fluctuation in glottal flow rate as $U_A$. The effect of increasing $a_{CT}$ was to obtain an initial increase in $U_A$, followed by a decrease. $U_A$ reaches a maximum when $a_{CT}$ ranges from 0.4 to 0.5. In contrast to $a_{CT}$, glottal flow fluctuation decreases monotonically with $a_{TA}$ and $a_{LC}$. For small values of $a_{LC}$, however, $U_A$ is always at a high level.

Likewise, we plotted the time rate of glottal flow-rate change, $U_B$, in Figure 6b, which is related to the sound strength of human [12]. This plot shows several important factors that have direct impact on the onset and intensity of vocal fold vibration in phonation ($U_B$). Compared with $U_A$, the behavior of $U_B$ is similar, which means that the variation in glottal flow velocity with different muscle activation is approximately the same as the variation in acceleration of change of the airflow. $U_B$ also provides an understanding of efforts to improve intensity of voice production. Changes in laryngeal muscle activities, such as increased CT muscle activities and decreased TA and LC muscle activities, will result in an increase in the amplitude of vocal fold vibration, thus increasing intensity of the voice. The cricothyroid muscle plays the primary role in controlling $F_0$. However, in the pathological vocal fold, the process of changing cricothyroid muscle activity to make $F_0$ reach the medium-high frequency is accompanied by irregular glottal airflow (large airflow velocity fluctuations and large airflow acceleration fluctuations). This suggests that stable voice production cannot be obtained in the vocal fold with a unilateral polyp. Thus, humans usually increase subglottal pressure to match the muscle activation to reduce laryngeal muscle fatigue and increase sound intensity. When the value of $a_{CT}$ is large enough to create a high frequency, $U_A$ and $U_B$ decrease significantly. According to Section 3.1, this is because the stiffness of the vocal fold is high at this point, making it almost impossible for the vocal folds to vibrate effectively. Hence, it is difficult for a patient with a polyp to phonate even with small fluctuations in the amplitude of $U_A$ and $U_B$. Similarly, for the thyroarytenoid and lateral cricoarytenoid muscles, low activity leads to unstable sound intensity. The values of $U_A$ and $U_B$ decrease monotonically with $a_{TA}$ and $a_{LC}$.

Two figures related to glottal flow rate are shown here that evaluate the effect of muscle activation on sound quality. Figure 6c shows the mean value of volume glottal flow rate ($U_C$) which is indicative of flow impedance. Figure 6d shows the leakage flow rate ($U_D$), which is the average of minimum volume flow rate over cycles, reflective of glottal gap size. This is a method for measuring the leakage of glottal flow during maximal glottic closure. High values of each parameter indicate a breathy voice quality.

Figure 6c shows that the mean flow rate decreases monotonically with $a_{CT}$ and increases monotonically with $a_{TA}$ and $a_{LC}$, a result that matches clinical results. Stretching the CT muscle leads to elongation of the vocal fold, but stretching the TA muscle or LC muscle leads to contraction of the vocal fold. The elongation and contraction of vocal fold affects the glottal area. The variation of $U_C$ is related to vocal fold vibration. Vibration at low frequency often requires high glottal volume flow velocity. The opposite is true at high frequency. This may seem counter-intuitive, but it is difficult for pathological vocal folds to vibrate regularly even if the muscle activity achieves the conditions for producing high-frequency sound. Because of the unilateral polyp, the amplitude of vocal fold vibration is small, and the glottal area has little variation. Increased vocal fold stiffness prevents normal vibration. Figure 6d can be regarded as a supplement to other figures. Leakage flow is not zero, which means that there is incomplete glottal closure in each cycle for the pathological vocal fold. Note that the fluctuation in the amplitude of volume flow rate increases as the minimum volume flow rate decreases. That is, in contrast to normal vocal folds, low leakage flow in pathological vocal folds may not result in a good voice quality. Vibrator periodicity should still be the primary consideration.
Figure 6. Comparison of effects of important airflow parameters related to laryngeal muscle activation on voice quality. (a) The fluctuation amplitude of volume flow rate, which is the root mean square of the time shift of glottal flow during the test. (b) The time-rate of change of the volume flow rate, which is the root mean square of the time shift of glottal flow velocity during the test. (c) The mean value of volume flow rate during the test time. (d) The average of minimum volume flow rate over cycles during the test time.
3.3. Fundamental Frequency and Related Perturbation Parameters

Other properties of the pathological vocal fold, such as natural frequency, probably have some differences from the normal vocal fold. To determine the effect of differing muscle activation on phonatory frequency and perturbation parameters, we fixed the activity of two muscles at a certain value and varied the value of the other to create plots of frequency and related perturbation parameters versus muscle activity (Figure 7).

![Figure 7](image-url)

**Figure 7.** Effect of CT, TA, LC activity on fundamental frequency and perturbation parameters. (a) Mean fundamental frequency. (b) Mean jitter (variation in frequency). (c) Mean shimmer (variation in amplitude).

Figure 7a shows that the CT muscle is most highly correlated with fundamental frequency. Similar to normal people, increasing CT muscle activity will always increase
$F_0$ in people with vocal fold polyps. However, increasing TA and LC muscle activity only causes a decrease in $F_0$. The deformation of the polyp counteracts the increase in stiffness. Activation of the TA and LC muscle under this condition may only shorten the vocal fold and reduce the stiffness. These examples illustrate the antagonistic roles that the CT and adductor muscles play in phonation onset parameters. As pointed out in previous work, these muscle activities could be interpreted as a requirement for stability in the laryngeal framework. When the vocal folds are being tensed, restraining forces may be needed to counteract gross movement of the larynx. In our pathological model, the control range of the vocal fold on frequency may be wider than that seen in the normal physiological situation. The reason is that we amplified the influence of muscle activation on vocal fold vibration. Our model aims to shed light on the effect of muscle activation on frequency to help find adapted muscle activities for different phonation situations. In fact, it should be noted that the actual muscle activation range of polyp vocal folds is much smaller than that of normal vocal folds. Hence, there will be a decrease in the upper frequency range for actual pathological vocal folds.

Variations in frequency (jitter) and amplitude (shimmer) have been commonly used to describe pathological voice quality. For our model, jitter is calculated by the average absolute difference between consecutive periods, divided by the average period, as

$$\text{Jitter} = \frac{1}{N-1} \sum_{i=1}^{N-1} |T_i - T_{i-1}| \frac{1}{N} \sum_{i=1}^{N} T_i$$

where $T_i$ is the extracted period length, and $N$ is the number of periods [26]. The mean value of jitter during phonation is depicted in Figure 7b. Overall, there was not a strong correlation with activity of any muscle. For large values of $\alpha_{CT}$, the jitter was high, reflective of aperiodic vibration. For small values of $\alpha_{TA}$, a decrease in $\alpha_{TA}$ will generally cause an increase in jitter.

Shimmer is the cycle-to-cycle variation in amplitude. Similar to jitter, shimmer is measured as the average absolute difference between the amplitudes of consecutive periods, divided by the average amplitude:

$$\text{Shimmer} = \frac{1}{N-1} \sum_{i=1}^{N-1} |A_i - A_{i-1}| \frac{1}{N} \sum_{i=1}^{N} A_i$$

where $A_i$ is the extracted amplitude and $N$ is the number of extracted periods [26]. Figure 7c shows the mean value of shimmer on phonation. Shimmer seems to have a strong link with CT and LC activity. The waveform of $U_A$ is similar to the waveform for shimmer. In general, increasing $\alpha_{CT}$ will cause an initial increase in shimmer, followed by a decrease. Shimmer is high when $\alpha_{CT}$ ranges from 0.3 to 0.6. When $\alpha_{CT}$ is between 0.3 and 0.4, shimmer is at its maximum. For LC muscle, shimmer increases monotonically when $\alpha_{LC} < 0.5$. In particular, for $\alpha_{LC} < 0.2$, the value of shimmer is much higher than at other activity levels.

Figure 7 narrows the range of muscle activation for normal phonation by frequency and perturbation parameters. Similar to the results in the previous section, in addition to the difficulty with high-frequency vibration, patients with a vocal fold polyp should pay attention to increasing their TA and LC muscle activity while decreasing CT muscle activity to produce a voice in a medium-high frequency range. One of the reasons for limited pitch range may be a decreased ability to vary vocal fold length. As shown in the figure, vocal fold vibration is irregular when $\alpha_{CT} > 0.4$, so CT activity will be restricted to a small range. The results also provide insights into the increase in fundamental frequency in response to effort. For patient with a vocal fold polyp, obtaining a better balance between CT and TA muscle activity is critical. As the results in Figures 6 and 7 show, when the value of $\alpha_{CT}$ is not small, TA muscle activity has little influence on voice quality. Hence, TA muscle activity can be increased appropriately to control $F_0$ while avoiding the generation
of high-frequency sound. LC activity should be kept greater than 0.3 even when the pitch is in the low-frequency range.

4. Conclusions

In this paper, a low-dimensional mass model is proposed to study the effects of muscle activation in a unilateral polyp setting. The simulations showed that different combinations of muscle activation have different effects on frequency, quality, and intensity. Based on the experimental results, we can draw conclusions about the optimal degree of muscle activation: $a_{CT}$ should be around 0.3, $a_{LC}$ should be fixed at greater than 0.3, and $a_{TA}$ can assume most values. Model simulations demonstrate that, in a unilateral polyp setting, combined TA and LC muscle activity will produce a more periodic voice at a higher frequency compared to high CT activity alone. The upper frequency range is also limited.

Overall, our computational model produced a substantial amount of significant information about the human vocal fold during phonation. However, the model is still simplistic compared with in vivo vocal fold geometry [27]. Future studies will focus on more realistic three-dimensional models.

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