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Abstract. Subglottic stenosis (SGS) is a challenging disease to diagnose in neonates. Long-range optical coherence tomography (OCT) is an optical imaging modality that has been described to image the subglottis in intubated neonates. A major challenge associated with OCT imaging is the lack of an automated method for image analysis and micrometry of large volumes of data that are acquired with each airway scan (1 to 2 Gb). We developed a tissue segmentation algorithm that identifies, measures, and conducts image analysis on tissue layers within the mucosa and submucosa and compared these automated tissue measurements with manual tracings. We noted small but statistically significant differences in thickness measurements of the mucosa and submucosa layers in the larynx ($p < 0.001$), subglottis ($p = 0.015$), and trachea ($p = 0.012$). The automated algorithm was also shown to be over 8 times faster than the manual approach. Moderate Pearson correlations were found between different tissue texture parameters and the patient’s gestational age at birth, age in days, duration of intubation, and differences with age (mean age 17 days). Automated OCT data analysis is necessary in the diagnosis and monitoring of SGS, as it can provide vital information about the airway in real time and aid clinicians in making management decisions for intubated neonates. © The Authors. Published by SPIE under a Creative Commons Attribution 4.0 Unported License. Distribution or reproduction of this work in whole or in part requires full attribution of the original publication, including its DOI. [DOI: 10.1117/1.JBO.24.9.096001]

Keywords: diagnostic imaging; intubation injury; neonate; optical coherence tomography; subglottic stenosis; texture analysis.

1 Introduction

Subglottic stenosis (SGS) in the neonatal population poses a significant diagnostic and management challenge. Injury to the delicate subglottic epithelium in neonates can result from a myriad of causative factors, including cyclical micromotion of the ventilation circuit, bacterial infection, biofilm formation, and direct interfacing of the endotracheal tube (ETT) with the subglottic mucosa. Following acute mucosal inflammation, edema and fibrosis can lead to SGS, which in some cases can be progressive and life threatening. SGS has a contemporary incidence between 0% and 2% per year in these patients, with the mean total charge in discharges from hospitalization being upward of $110,000.2,4 While direct laryngoscopy and bronchoscopy remain the diagnostic gold standard for SGS, this procedure requires general anesthesia and poses considerable risk of airway compromise.5 Hence, there exists a critical need for a less invasive and practical means for neonatologists and otolaryngologists to image the upper airway in intubated neonates.

Optical coherence tomography (OCT) is a high-speed, micrometer-resolution, cross-sectional diagnostic imaging modality that has been previously described to characterize changes in the subglottic mucosa following endotracheal intubation in animals and humans.6–8 Long-range OCT (LR-OCT) is an advanced adaptation of this technology, with an imaging range up to 20 mm.9,10 A practical limitation of OCT imaging is the massive volume of images recorded per airway scan (amounts to 1 to 2 Gb). OCT image processing and objective analysis can also be challenging and highly time-consuming, precluding real-time assessment of the airway and clinical decision-making.11,12 This limitation has also been observed in intravascular OCT imaging, where each scan can generate up to 1000 individual images and requires hours of offline manual analysis.13 If OCT were adopted in the neonatal intensive care unit (NICU) for SGS, thousands of images per patient would need to be analyzed, reconstructed, visualized, and then reviewed by an expert. Like other diagnostic studies widely used as screening measures, such as pap smears or complete blood counts, OCT image analysis would require a large component of automation for broad adoption in clinical medicine.14,15
Texture analysis of tissue images is a viable and automated process that has been previously utilized in tissue identification, pathologic quantification, and cancer detection.\textsuperscript{16-18} Texture properties represent tissue changes within an image and can be evaluated via spatial distributions of pixel intensities. Texture analysis can be investigated through gray-level occurrence matrices (GLCM) or by statistical analysis of texture properties by examining the spatial distributions of pixels.\textsuperscript{17} In 2017, we described the dependence of two texture properties, correlation and homogeneity, on intubation time in a rabbit model of SGS.\textsuperscript{7} This was the first step in the long-term objective of automating analysis of thousands of OCT images per patient and optimizing airway management to avoid SGS in intubated neonates.

In this study, we examined two key issues relevant to the automated analysis of OCT images of the subglottis. First, we evaluated software aimed at segmenting the subglottic airway wall into two distinct layers and directly compared these results with manual segmentation. Second, to these segmented regions, we applied basic texture matrix analysis to quantitatively define tissue characteristics and compared this data to patient clinical parameters, such as gestational age (GA) or duration of intubation.

2 Methods

2.1 Study Design

Analysis was conducted on data acquired from a previous study of intubated neonates, which examined the use of OCT on the neonatal laryngotracheal airway.\textsuperscript{19} Each neonate was imaged in the NICU at either the University of California Irvine (UCI) or Children’s Hospital of Orange County (CHOC). Clinical parameters including GA at birth and at time of imaging, total duration of intubation, ETT size, and patient weight were all recorded for each patient. Each subject’s family provided written informed consent for participation, and imaging was performed following the protocol approved by the Institutional Review Boards at UCI and CHOC.

2.2 Optical Coherence Tomography System and Imaging

Data sets for analysis were acquired through an LR-OCT system that has been previously described.\textsuperscript{19,20} In brief, a 1310-nm swept source laser with a repetition rate of 50 kHz (Axsun Technologies, Massachusetts) was utilized as the light source of the OCT engine, and an acousto-optic modulator (Brimrose Corp., Massachusetts) was incorporated to generate a career frequency of 150 MHz in the reference arm to achieve a 20-nm working distance. Flexible, side-view endoscopic OCT probes with an outer diameter of 0.7 mm and length of 65 to 70 cm were used for imaging the neonatal subglottis. A sterilized, distally sealed sheath encasing the probe was inserted into the subglottic region through the ETT in an intubated patient. To acquire images, the probe was rotated at a speed of 1.56 mm/s. This continuous helical scanning scheme generated 300 to 600 360-deg images in total, with each image comprised of 2000 axial scans (A-lines) and each A-line storing 2048 12-bit pixels. Images were compressed into 8-bit bitmap for analysis.

2.3 Data Collection and Selection

Details of the methodology of OCT image acquisition have been previously described.\textsuperscript{19} Spiral OCT scans through the larynx, subglottis, and proximal trachea were obtained of each subject. A total of 58 OCT data sets were obtained, based on data from 49 different patients. Nine patients who were intubated for extended durations were serially imaged on different days. Each data set was evaluated for image quality prior to segmentation and texture analysis. Image quality was determined by the extent to which background noise affected the automated segmentation algorithm and the thickness measurements of the upper airway (mucosa and submucosa layers) that were produced. If the speckle noise or ghost images could not be initially cropped out of the OCT images or the noise significantly impacted the process of edge detection so that the anatomical structures of the image could not be accurately segmented [Figs. 1(e)–1(f)], the data set was excluded from analysis. Figures 1(a)–1(d) illustrate data with adequate image quality, and Figs. 1(e)–1(f) demonstrate examples of discarded data with image artifacts that precluded analysis. Many images had either minimal background noise [Figs. 1(a)–1(b)] or noise that was either filtered out of the image or did not significantly impact the process of segmentation [Figs. 1(c)–1(d)]. If the tissue contours of the mucosa and submucosa layers were faint or distorted, the data set was also excluded from analysis. In both circumstances, tissue segmentation by a human observer would also not be feasible.

Seventeen out of the 58 OCT data sets were excluded due to the aforementioned reasons. Out of the remaining 41 data sets, select trachea (3), subglottis (2), and larynx (2) subsets of data were discarded from within their respective data set for the same reasons. In those cases, the remaining analyzable images within the data set were included in the analysis. Each data set was divided into three airway segments, larynx, subglottis, and trachea, based on well-defined anatomic markers (e.g., true vocal fold margin, tracheal or cricoid cartilage, and laryngeal ventricles), which were identified in the OCT images. OCT has previously been used in the recognition of tissue structures.\textsuperscript{21,22}

Both automated and manual segmentations were performed by one trained study member (K.K.). Manual segmentation was performed on most cross-sectional images in 28 out of the total 41 data sets, whereas automated segmentation and texture analysis were performed on almost every one of the 41 data sets (38 trachea, 39 subglottis, and 39 larynx data sets). A single OCT data set ranged from 100 to 400 (mean 200) individual images, depending on how distal the OCT probe was positioned in the trachea at the beginning of the airway scan. The trachea typically included a greater number of frames compared to the larynx and subglottis, with a total count depending on how distal the OCT probe was positioned in the trachea at the beginning of the airway scan. Given the proportionally higher amount of trachea data, approximately every fifth tracheal image was segmented to achieve a proportionally sized data subset and to conserve time; at least 20 trachea images were analyzed per data set. In contrast, each and every image within the subglottis and larynx subsets was segmented; on average, there were approximately 40 data frames in each of these two airway segments. Approximately 100 total images were manually and autosegmented per data set. The amount of time required for the program to segment each data set was also recorded.
2.4 Manual Segmentation

In manual segmentation, tracing of the airway wall layers was performed at user discretion with a computer mouse in a program written in MATLAB with graphical user interface. Anatomical OCT images of the airway are displayed in polar coordinates [Fig. 2(a)], though representation in Cartesian form facilitates better identification of key morphological features during manual segmentation [Figs. 2(b) and 2(c)]. Layers of airway wall microstructure (e.g., epithelium, submucosa, and cartilage) were distinguished based on differences in grayscale (pixel) intensity, as each tissue layer has its unique optical composition.23 The first layer of the airway wall bordering the ETT is the mucosa, which consists of epithelium, basement membrane, and lamina propria. Underlying the mucosa is the submucosa, which is followed by cartilage tissue, when present. Two tissue interfaces were identified: the interface between the airway lumen and the mucosal epithelium, and the interface between the submucosa and cartilage. Both boundaries were traced and the mucosa–submucosa boundary was delineated by manual segmentation [Fig. 2(d)]. The layers identified were the same as those identified by autosegmentation, as described below.

2.5 Automated Segmentation

The automated segmentation algorithm that was developed for imaging bovine airway OCT images following smoke inhalation injury was adapted for use in this study.23 This graph theory-based segmentation algorithm was used on successive OCT images in each data set via a three-step process: a preprocessing step, followed by an edge detection algorithm, and lastly a thickness measurement heuristic (Fig. 3).

In the preprocessing step, the airway wall, including the mucosa and submucosa, was distinguished from the background (e.g., air, cartilage, ETT, OCT probe sheath, artifact, and noise). This step also addressed problems posed by speckle noise generated during imaging, ghost objects produced by internal interference of the optics, and mirror image/objects induced by Fourier transformation.24–26 These obstacles were compensated for with a series of low-level procedures, such as speckle noise...
suppression, area filtering, dilation, or bridging. The different
thresholds that controlled each of these procedures had to be
manually set. Additionally, because the imaging probe was
protected by a plastic sheath, it was necessary for this sheath
to be identified and excluded from segmentation. Images were
manually cropped to further eliminate any possible artifacts
that could adversely affect the edge delineation step of the seg-
mentation algorithm. Following frame cropping and threshold
adjustment to suppress speckle noise, segmented grayscale
images with reliable airway structures for edge detection were
generated. These image stacks were subsequently rapidly ana-
yzed by the program in real time on our computer workstation.

The edge detection step took previously generated localized
airway regions and performed the segmentation task using a
dynamic programming algorithm. The algorithm used graph
construction and recursive solution finding to obtain the optimal
path for the edge detection process. It treated every pixel in
the OCT image as a node in a graph to find the shortest path
(the edge) for the graph, based on the pixel intensity. The edge
was refined, or smoothed, by averaging neighboring pixels. The
airway lumen that was extracted was used as the first reference
dynamic edge (epithelial surface) and the outer boundary of the airway
wall, at the submucosa–cartilage junction represents the second
reference edge. The middle edge, or the boundary between the
mucosa and submucosa layers, was found by repeating this
process in the closed region between the epithelium and carti-
lage edges. The average thickness of the mucosa and submucosa
layers could then be determined after the mucosa and submu-
cosa layers were accurately delineated.

2.6 Texture Analysis

Each autosegmented image was evaluated using texture
analysis. This was done using the GLCM—a statistical
method of examining the spatial distributions of pixels. Four
different tissue texture properties (correlation, homogeneity,
contrast, and energy) were analyzed at four different angles
of the GLCM (0 deg, 45 deg, 90 deg, and 135 deg), resulting
in 16 unique texture variables for each image. Texture analysis
was performed on each frame in a data set in all 41 data sets.
Automatic and manual segmentation measurements were
compared using pairwise t-tests with significance level 0.05.
Associations between patient clinical parameters (e.g., GA at
birth and duration of intubation) and texture variables were
explored using Pearson correlations. As this was an exploratory
analysis, significance levels were not adjusted for multiple
comparisons.

3 Results

The automated and manual segmentation methods were found to
have very small but statistically significant differences in thick-
ness measurements. Sample sizes were 28, 27, and 23 data sets
for the trachea, subglottis, and larynx, respectively (Table 1).
Statistical analysis was performed after the segmentation of each
data set, and airway segmentation plateaued with statistical sig-
nificance around 28 data sets. Hence, manual segmentation was
performed on only 28 of the 41 total data sets. We noted that
the automated segmentation method tended to consistently
but minimally underestimate the mean airway wall thickness
(mucosa and submucosa) when compared to manual measure-
ments, measured in pixels. Manual segmentation of each image
stack (~100 frames) required 30 to 40 min, depending on the
total number of images, and this time was noted. Automated
segmentation, compared to manual, was consistently faster
\( p < 0.01 \) for analysis of the trachea (9 times faster), subglottis
(9 times faster), and larynx (6.7 times faster) (Fig. 4).

Analysis of the 16 different texture variables did not show
any statistically significant associations with thickness measures
but did demonstrate modest correlations with select patient
variables. In the subglottis, there were moderate correlations
between the contrast, energy, and homogeneity texture variables
and the age in days of the neonate (Table 2). In addition, in the
subglottis, the energy texture variables had a moderate correla-
tion with the patient’s GA at birth and the contrast texture

| Table 1 Comparison between the airway wall thickness values (mea-
| measured in pixels) obtained by automated and manual segmentations
| (paired t-test). |
|---------------|------------|--------|
|               | Mean thickness | P-value |
| Autosegmentation | Manual segmentation |
| Trachea       | 38.013       | 38.852  | 0.012 |
| Subglottis    | 44.793       | 46.657  | 0.015 |
| Larynx        | 52.183       | 55.069  | <0.001 |
variables had a moderate correlation with the number of days the neonate was intubated (Table 2). Lastly, in the subglottis, there were consistent differences by age when categorized as below and above the median (≤17 days versus > 17 days) for the energy and homogeneity texture variables. Although these differences are consistent, due to the small sample size they would not reach statistical significance when adjusted for multiple comparisons (Table 3). Other clinical variables, including race, gender, or weight, showed no correlations with the texture variables.

![Fig. 4 Timewise comparison of manual and automated segmentations.](image)

**Table 2** Comparison between texture variables and clinical parameters in subglottis (t-test).

<table>
<thead>
<tr>
<th>Texture variables</th>
<th>Age in days</th>
<th>GA at birth</th>
<th>Days intubated</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Correlation (p-value)</td>
<td>Correlation (p-value)</td>
<td>Correlation (p-value)</td>
</tr>
<tr>
<td>Contrast (0D)</td>
<td>0.317 (0.049)</td>
<td>-0.276 (0.089)</td>
<td>0.283 (0.081)</td>
</tr>
<tr>
<td>Contrast (45D)</td>
<td>0.316 (0.05)</td>
<td>-0.285 (0.079)</td>
<td>0.282 (0.082)</td>
</tr>
<tr>
<td>Contrast (90D)</td>
<td>0.333 (0.039)</td>
<td>-0.286 (0.077)</td>
<td>0.279 (0.085)</td>
</tr>
<tr>
<td>Contrast (135D)</td>
<td>0.316 (0.05)</td>
<td>-0.285 (0.079)</td>
<td>0.282 (0.082)</td>
</tr>
<tr>
<td>Correlation (0D)</td>
<td>0.184 (0.262)</td>
<td>-0.305 (0.059)</td>
<td>0.203 (0.215)</td>
</tr>
<tr>
<td>Correlation (45D)</td>
<td>0.181 (0.27)</td>
<td>-0.28 (0.084)</td>
<td>0.207 (0.206)</td>
</tr>
<tr>
<td>Correlation (90D)</td>
<td>0.25 (0.125)</td>
<td>-0.319 (0.048)</td>
<td>0.268 (0.099)</td>
</tr>
<tr>
<td>Correlation (135D)</td>
<td>0.178 (0.277)</td>
<td>-0.28 (0.084)</td>
<td>0.203 (0.216)</td>
</tr>
<tr>
<td>Energy (0D)</td>
<td>-0.347 (0.031)</td>
<td>0.33 (0.04)</td>
<td>-0.318 (0.049)</td>
</tr>
<tr>
<td>Energy (45D)</td>
<td>-0.344 (0.032)</td>
<td>0.332 (0.039)</td>
<td>-0.315 (0.051)</td>
</tr>
<tr>
<td>Energy (90D)</td>
<td>-0.347 (0.031)</td>
<td>0.332 (0.039)</td>
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<td>Energy (135D)</td>
<td>-0.344 (0.032)</td>
<td>0.332 (0.039)</td>
<td>-0.315 (0.051)</td>
</tr>
<tr>
<td>Homogeneity (0D)</td>
<td>-0.329 (0.041)</td>
<td>0.288 (0.075)</td>
<td>-0.293 (0.07)</td>
</tr>
<tr>
<td>Homogeneity (45D)</td>
<td>-0.328 (0.042)</td>
<td>0.297 (0.067)</td>
<td>-0.293 (0.071)</td>
</tr>
<tr>
<td>Homogeneity (90D)</td>
<td>-0.343 (0.032)</td>
<td>0.298 (0.065)</td>
<td>-0.29 (0.074)</td>
</tr>
<tr>
<td>Homogeneity (135D)</td>
<td>-0.328 (0.041)</td>
<td>0.296 (0.067)</td>
<td>-0.293 (0.07)</td>
</tr>
</tbody>
</table>

4 Discussion

OCT is a high-resolution cross-sectional imaging modality that is widely used in ophthalmologic, coronary vasculature, and dermatologic imaging. While OCT has been shown to reliably measure and characterize airway wall morphometry, it has limited practical applicability to airway monitoring due to the vast amount of data acquired with each scan. We aimed to address this need by constructing automated OCT data analysis methods, with the goal to ultimately provide clinicians with objective OCT data in a timely and practical manner.

4.1 Automated Segmentation

Unlike a manual approach, the autosegmentation code may erroneously classify noise in the OCT images as tissue structure, as it does not remove all noise during the preprocessing step of the algorithm. This occurs commonly with speckle noise at the lateral margins of the image and if it is not addressed, can lead to unreliable measurements during segmentation, resulting in an overestimation of wall thickness. To compensate, images must be analyzed with the selection of a specific and usually smaller than normal window size to find regions of pixel density that do not include speckle noise at the lateral margins of the image. The criteria used here thus tend to minimally but consistently underestimate the thickness of the mucosa and submucosa layers of the upper airway of patient data sets. More importantly, this minimal error comes with a critical gain in the consistency of autosegmentation program, as evident by our findings illustrated in Table 1.

Due to the necessity of the image preprocessing steps, the autosegmentation process was indeed semiautomated as opposed to fully automated, as some level of screening had to be performed. However, once the preprocessing steps were completed, the image stack was rapidly analyzed. More importantly, this entire semiautomated process, which encompassed the manual preprocessing and automated segmentation steps, was still significantly faster than a purely manual approach.

There is great clinical value in the rapid detection and segmentation of the upper airway wall. Edema can develop very quickly in the airway wall, and the state of the mucosa can change day by day. Therefore, clinicians need real-time data

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**Table 3** Comparison between texture variables and age of patient (≤17 days versus > 17 days) in subglottis (t-test).

<table>
<thead>
<tr>
<th>Subglottis</th>
<th>Unadjusted p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Energy (0D)</td>
<td>0.025</td>
</tr>
<tr>
<td>Homogeneity (0D)</td>
<td>0.038</td>
</tr>
<tr>
<td>Energy (45D)</td>
<td>0.026</td>
</tr>
<tr>
<td>Homogeneity (45D)</td>
<td>0.044</td>
</tr>
<tr>
<td>Energy (90D)</td>
<td>0.028</td>
</tr>
<tr>
<td>Homogeneity (90D)</td>
<td>0.037</td>
</tr>
<tr>
<td>Energy (135D)</td>
<td>0.028</td>
</tr>
<tr>
<td>Homogeneity (135D)</td>
<td>0.044</td>
</tr>
</tbody>
</table>
to determine whether to, for example, extubate or perform a tracheostomy on a patient, as other extubation readiness tests are of low value, and this type of real-time data can be obtained much more feasibly with an automated method of data analysis.36 Other studies have previously reported automated measurement of images, including that of CT and MRI images.37–39 However, OCT images have a much higher resolution and clarity compared to CT and MRI images, and this combined with our segmentation algorithm and texture analysis separates our image analysis from other studies.

4.2 Texture Analysis

Texture properties serve as an objective proxy for microscopic changes within native airway wall tissue.40,41 The optical properties of the airway wall differ between patients, presumably because airway wall microanatomy changes with physiologic states such as edema and fibrosis. Previous OCT studies have described texture analysis for the purpose of tissue classification.42,43–44 However, the correlation of texture properties with specific physiologic changes is not well understood. Ajose-Popoola et al.7 described texture analysis following OCT imaging of the rabbit subglottis following controlled brush injuries, in which significant correlations were noted between texture variables homogeneity and correlation with time from injury. These animal studies provided the motivation for the current analysis of in vivo human data sets.

We used Pearson correlations to identify associations between the texture variables and clinical neonate variables, such as GA at birth, intubation time, and ETT size. We noted moderate correlations between subglottic texture properties and GA at birth, age, and duration of intubation. Second, texture analysis of the subglottis showed consistent differences with age (median age of 17 days) at or near the significance level of p < 0.05 before adjustment for multiple comparisons. Findings here suggest that texture analysis of OCT images may potentially be used to correlate physiologic changes in tissue composition with these specific patient metrics. For example, two neonates with respective intubation durations of 3 days and 3 weeks may indeed have similar subglottic wall thicknesses. However, the optical microstructure of their respective airway walls can be characterized by texture properties, which may help differentiate healthy tissue from edema in recently intubated neonates and fibrosis following lengthy or repeated intubations.

4.3 Study Limitations

As all imaging was performed in vivo, direct comparison of airway wall measurements and microanatomic composition with tissue histology was not possible, as laryngeal biopsy is generally never indicated for SGS management and postmortem evaluation of the neonate is not widely practiced as the emotional burden for families is often insurmountable. However, given prior animal OCT studies that demonstrate correlation of OCT and histology, our study offers a means to attain objective, structural information on the human subglottic airway.45 Texture analysis is a simple feature extraction technique that can be used as a reliable deterministic statistical process. Although it has its limitations, as various factors can affect the reliability, reproducibility, and robustness of texture features, we were able to use this rudimentary feature classification technology to show moderate correlations nearing significance between clinical neonate variables and certain texture variables, even with a small patient sample.46 While we obtained small effect sizes with this small patient sample, this is an early result that warrants further study. A larger sample size and future improvements in the segmentation algorithms may lead to stronger associations between clinical neonate variables and texture variables, resulting in a larger effect size. Although texture analysis is a well-established albeit rudimentary pattern recognition approach, other pattern recognition and feature extractions methods, such as fuzzy clustering, could also be also used to find evidence of changes in tissue structure.47 Fuzzy clustering is a process that sorts specific elements of tissue into different classes so that elements in a class are similar to each other. It has already been used to analyze nerve fibers in glaucoma patients as well as to evaluate breast cancer nuclei as malign or benign and advance breast cancer diagnosis.48,49 As OCT is a high data volume imaging modality with multispecialty applications, this underscores the critical need to develop automated techniques and advanced pattern recognition approaches, possibly by incorporating machine learning, to analyze all this data.

4.4 Future Steps

Many of the issues that result in more of a semiautomated approach are technical factors in the OCT system which make the preprocessing step necessary, such as the speckle noise and image artifacts that appear in the resulting OCT images. Current research efforts at our institution have largely eliminated these factors, and future OCT image analysis could potentially eliminate all preprocessing steps and become a fully automated process. We are also designing additional algorithms to automatically both segment airway wall layers more efficiently and analyze tissue composition, such as three-dimensional tissue segmentation.50 This would simplify the process of recognizing upper airway tissue morphology changes and dramatically speed up diagnosis of upper airway diseases. We expect that our work will be reproducible and of benefit to other OCT imaging teams. With minimal in-person training, we foresee other medical centers being able to utilize our segmentation algorithms. Moreover, we believe that with minimal alterations, if any, our algorithm will be translated to measure other tissue types.

The integration of vertical-cavity surface-emitting lasers (VCSEL) technology with OCT systems is a rapidly developing arena, as these imaging systems feature vast improvements in image quality with refined optics and probe design.51 VCSEL OCT imaging is an active area of interest in our group, and we aim to perform further imaging in intubated neonates and gain improved quality image sets to better understand relationships between OCT data and clinical parameters.

5 Conclusion

OCT is a high-resolution diagnostic imaging modality that has tremendous potential as a means to objectively and serially analyze the airway in intubated neonates. The automated image analysis algorithm described in this report offers an efficient and precise solution to analyzing large-volume OCT data stacks. With automated image analysis, OCT may ultimately offer clinicians real-time information about the health of the intubated neonatal airway and aid in airway management.

Disclosures

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Biographies of the authors are not available.