Laryngotracheobronchial papillomatosis: findings on computed tomography scans of the chest*

Papilomatose laringotraqueobrônquica: aspectos em tomografia computadorizada de tórax

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Abstract

Objective: To present the findings of computed tomography (CT) scans of the chest in patients with laryngotracheobronchial papillomatosis.

Methods: We retrospectively analyzed CT scans of eight patients, five males and three females, ranging from 5 to 18 years of age with a mean age of 10.5 years. Images were independently reviewed by two radiologists. In discrepant cases, a consensus was reached.

Results: The most common CT findings were intratracheal polypoid lesions and pulmonary nodules, many of which were cavitated.

Conclusions: In patients with laryngotracheobronchial papillomatosis, the most common tomographic finding was the combination of intratracheal polypoid lesions and multiple pulmonary nodules, many of which were cavitated.

Keywords: Tomography, X-ray computed; Papilloma; Tracheal neoplasms.

Introduction

Recurrent respiratory papillomatosis, previously known as childhood laryngeal papillomatosis is typically a benign and self-limiting disease.1 It is caused by infection of the upper respiratory tract with the human papillomavirus (HPV),2,3 resulting in the formation of papillomas, which are the most common benign tumors occurring in the respiratory tract during childhood.4,5 Although the disease is more common in children, it can also occur in adults.3 It is typically restricted to the larynx but can also invade the tracheobronchial tree and the pulmonary parenchyma.6,6 Central airway involvement is seen in 2% to 5% of patients with laryngeal papillomas, whereas small airway or alveolar involvement occurs in less than 1%.1,7 The disseminated form is also known as laryngotracheobronchial papillomatosis.

The definitive diagnosis of laryngotracheobronchial papillomatosis is made based on the results of transbronchial biopsy (bronchoscopy) of laryngeal or tracheal lesions, although the

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diagnosis can be suspected on the basis of computed tomography (CT) findings. Here, we present the tomographic aspects observed in eight patients with laryngotracheobronchial papillomatosis.

**Methods**

This was a retrospective study analyzing CT scans of the chest related to eight patients diagnosed with laryngotracheobronchial papillomatosis at five medical facilities in three Brazilian states (Rio de Janeiro, São Paulo and Bahia). Five patients were male, and 3 were female. Ages ranged from 5 to 18 years (mean, 10.5 years). In all of the cases, the diagnosis had been made some years prior based on findings of papillomas in the upper airways. Five of the eight patients had been less than two years old at the time of diagnosis. All of the patients had undergone multiple previous resections. In the four cases in which the resections had been quantified, the number ranged from 15 to 40 (mean, 25). In one patient, malignant degeneration, to squamous cell carcinoma, was reported.

The most common clinical complaint was hoarseness, which occurred in six cases. Other findings

**Figure 1** - a) Slice at the level of the trachea. Tracheal wall with irregular borders and a polypoid formation on the left. Homogeneous nodule in the left upper lobe. b) Slice at the level of the main bronchi. Multiple, thick-walled cavitated nodules in the right lung.

**Figure 2** - a) Slice at the level of the upper lobes. Trachea with irregular walls. Voluminous cystic formation in the right upper lobe. b) Slice at a slightly lower level than that shown in a. Note the better view of the cystic lesion, which has irregular internal walls. c) Slice at the level of the main bronchi, showing a thin-walled cystic lesion in the lingula.
included bronchospasm (in two), dyspnea (in two), recurrent respiratory infections (in two), respiratory failure (in two), dry cough (in one) and purulent expectoration (in one). Seven of the eight patients had been submitted to tracheostomy, at some point during the evolution of the disease, to remove the obstruction.

In all cases, the CT scans were performed when the patients presented involvement of the upper airways, especially of the larynx. Seven patients had presented tracheal and pulmonary involvement; one had presented tracheal involvement alone.

The CT scans of the chest were performed using different various tomography scanners, axial slices ranging from 5 to 10 mm in thickness, in 10-mm increments, during a deep inhalation, from the apices to the lung bases. Some patients were also submitted to high-resolution CT, with slices of 1 mm or 2 mm in thickness. Tests were performed using a parenchymal window, with

![Figure 3](image1.png)

**Figure 3** - a) Slice with mediastinal window, upper lung fields. Polypoid lesion on the posterior wall of the trachea. b and c) Slices at the level of the lower lobes. Multiple, multilobulated nodular lesions, many cavitated, some with thick, irregular walls and others with thin walls. Note the signs of air trapping.

![Figure 4](image2.png)

**Figure 4** - a) Multiple areas of flow impairment affecting the entire circumference of the trachea. b) Slice at the level of the bronchial bifurcation. Various cavitated nodular lesions in both lungs, with a tendency toward confluence on the left. Note the mass with irregular borders in the right lung, as well as the nodular formation in the right main bronchus.
patients presented involvement of the pulmonary parenchyma. In all eight cases, there were cavitated polypoid lesions, with irregular internal borders and walls of various thicknesses, multilobulated in six cases and presenting confluence of the lesions in five (Figures 1 through 5). Solid nodules were seen in six patients. An air-fluid level was observed in only one patient. The lesions were predominantly in the lung bases in three patients and in the middle thirds in two, whereas they were evenly distributed in two. In one patient, the lesions were accompanied by a mass, and, in another patient, they were accompanied by consolidation. In the latter case (Figure 5), there was malignant degeneration in multiple lesions. Signs of air trapping were also observed in one case. There were no identified instances of lymph node enlargement or pleural effusion.

Discussion

Laryngotracheobronchial papillomatosis is a rare, benign disease in children and is even more rarely seen in adults. The histological presentation is benign squamous epithelial stratification, and the disease is typically restricted to the larynx. However, it can occasionally become aggressive, resulting in persistent or recurrent involvement of the nasopharynx, larynx and tracheobronchial tree. Dissemination to the pulmonary parenchyma occurs in less than 1% of cases. Such dissemination can occur years after the diagnosis of laryngeal papilloma. There have been no reports of the disease affecting the tracheobronchial tree without prior upper airway involvement. All of the patients evaluated in the present study had been diagnosed years earlier through tracheobronchial biopsy (via bronchoscopy).

The etiological agent of laryngotracheobronchial papillomatosis is HPV, which is a virus that presents great genetic variation, various types and subtypes having been genetically defined. The types most often associated with laryngotracheobronchial papillomatosis are HPV-6 and HPV-11, followed by HPV-16 and HPV-18. These last two types have more often been associated with malignant transformation, particularly with transformation to squamous cell carcinoma. Infection occurs most frequently at birth and is related to passage through the birth canal in mothers with vulvar papillomatosis lesions. However, infection can also occur later.
in life, possibly by sexual transmission [oral contact
with infected external genitalia]. The forms of laryngotraceobronchial papil-
ломатosis are classified as childhood or adult. The
childhood form occurs in patients less than 20 years
of age, presenting multiple lesions that are unpre-
dictable in terms of their response to treatment,
and the rates of recurrence tend to be high. In this
population, solitary papillomas are rare. Among
adults, (patients over 20 years of age at diagnosis),
the disease is more common in men, and the papil-
ломas are typically solitary, present a high degree
of inflammatory reactivity, do not usually dissemi-
nate and recur less frequently than those seen in
the childhood form. Various hypotheses have been
formulated in order to explain distal dissemination
of laryngeal papillomatosis: contiguous spread;
diffuse viral contamination; multicentric origin of
the papillomas; and canalicular dissemination of
fragments. Iatrogenic factors, such as laryn-
goscopy, bronchoscopy, tracheostomy and surgical
manipulation can also cause dissemination of papil-
ломas to the distal portion of the tracheobronchial
tree. In our study, seven patients had undergone
tracheostomy, at some point during the evolution
of the disease, in order to remove an obstruction.

In children, the clinical profile characteristic of
laryngeal papillomatosis is the triad of progressive
hoarseness, stridor and difficulty in breathing. In
adults, hoarseness is the most common finding. In
the trachea, the most common symptoms frequently
mimic obstructive pulmonary diseases such as
asthma and chronic pulmonary obstructive disease,
which can impede the diagnosis and lead to errors
in the treatment strategy. The physical examina-
tion can reveal wheezing, stridor, tachypnea and
accessory muscle recruitment during respiration.
Peripheral dissemination can lead to recurrent
pneumonia, as well as obstructive atelectasis and
malignant degeneration. Clinical symptoms can
include fever, cough, hemoptysis and progressive
dyspnea. In our sample, the most common
symptom was hoarseness.

The ideal method for diagnosing lesions in the
central airways is bronchoscopy, which makes it
possible to obtain biopsy samples of such lesions in
order to perform histopathological evaluation and to
plan therapeutic interventions. In most cases, laryn-
gotraceobronchial papillomatosis is easily diagnosed,
since the pulmonary presentation is typically preceded
by a rich clinical history and a previous diagnosis of
laryngeal papillomatosis. In the bronchoscopy, the
papillomas are seen as whitish polypoid lesions in the
larynx, trachea or bronchi. Bronchoscopy remains
the diagnostic method of choice, since it is both
diagnostic and therapeutic, allowing the resection of
the lesions, which can then be submitted to anato-
mopathological study.

Chest X-rays can occasionally produce findings
suggestive of the disease, such as the combina-
tion of solid or cavitated pulmonary nodules and
vegetative nodular lesions in the trachea or in the
main bronchi. Unlike the pulmonary nodules, which
are often identified on simple chest X-rays, intra-
lumen papillomas in the trachea or bronchi are
rarely visible on X-rays. In practice, therefore, the
disease is rarely diagnosed on the basis of chest
X-ray findings. The nodules are typically multiple,
well defined, of various dimensions, cavitated and
thick-walled, being more numerous in the basal and
posterior lung regions. The cavitated nodules can
be air-filled or, when infected, can present an air-
fluid level.

Chest X-rays are inferior to CT scans of the chest,
especially those performed using spiral CT (volum-
metric acquisition), in the initial phases of pulmonary
dissemination, due to the fact that CT better char-
acterizes and allows better visualization of nodular
tracheobronchial vegetation. Tomographic findings
include focal or diffuse airway narrowing caused
by the nodules. The nodules form on the mucosal
surface, and their invasion into the lumen is best
evaluated using CT. These alterations are easily
related to the presence of infec-
tions with papillomatosis. Other findings related
to airway obstruction and accompanying infections
are atelectasis, consolidations, air trapping and
bronchiectasis. In our sample, polypoid forma-
tions in the trachea were observed on all of the CT
scans. Seven patients presented involvement of the
pulmonary parenchyma, characterized by cavitated
polypoid lesions with irregular internal borders
and walls of various thicknesses, with a multilob-
ulated aspect and a tendency toward confluence.
Solid nodules were seen in six patients. None of
the patients presented lymph node enlargement or
pleural effusion.

Papillomas appear as masses or nodules (single
or multiple) that are exophytic (sessile or peduncu-
lated), soft and friable, most often seen on the vocal
cords, ventricular folds, subglottis and laryngeal surface of the epiglottis. Histologically, papillomas present as projections of the stratified squamous epithelium with fibrovascular centers. Hyperplasia of basal cells and large vacuolated epithelial cells with clear cytoplasm are typical findings. When the lesion invades the tracheobronchial tree, the epithelium can be squamous or ciliated and cylindrical. The pulmonary lesions are foci of squamous epithelium that grow circumferentially within the alveoli, depending on them for vascularization. Near the central portion of a solid or cavitated lesion, there are areas necrosis and regeneration, containing fragments. In the periphery, squamous cells invade the adjacent alveoli by direct extension of the principal cell mass, lymphocytes and macrophages being identified in the alveolar content. These cells grow, coalesce and destroy the parenchyma, forming cavities.

Malignant degeneration to squamous cell carcinoma is reported in 1% to 10% of all cases of laryngotracheobronchial papillomatosis, typically occurring after radiotherapy or chemotherapy with bleomycin, as well as in patients with a history of smoking. It can occur in childhood or even decades after the diagnosis of benign papillomas. In most cases, it occurs in patients having previously presented dissemination of the disease to the tracheobronchial tree. Malignant degeneration rarely occurs in patients with the laryngeal form of the disease. In one of the cases evaluated here, there was malignant degeneration reported based on biopsy findings in three different lesions.

Small lesions provoking minimal symptoms can be treated with corticosteroids and antibiotics. Larger, symptomatic lesions extirpated through bronchoscopic procedures such as curettage, laser ablation, electrocautery and cryosurgery, surgical intervention (thoracotomy or sternotomy) rarely being necessary. Radiotherapy, autogenic vaccination and chemotherapy have proven unsuccessful. Antiviral therapy has also been employed. None of the various treatment protocols has been shown to be truly efficacious. Recurrence of papillomas is common, regardless of the treatment given. In practice, it is necessary to submit such patients to frequent bronchoscopic examinations.

This disease takes on greater importance due to the severity of its evolution and the fact that it affects children and adolescents, resulting in considerable morbidity and having a profound impact on patient quality of life, requiring multiple surgical excisions and tracheostomies; the prognosis is unfavorable, and the disease can even evolve to death. Although the diagnosis is typically made in childhood, based on a finding of papillomas in the upper airways, a subsequent finding of intrathoracic polypoid lesions, with or without pulmonary nodules (cavitated or not), should raise the suspicion of the disease.

References