Auditory and vestibular dysfunctions in systemic sclerosis: literature review

ALTERAÇÕES AUDITIVAS E VESTIBULARES NA ESCLEROSE SISTÊMICA: revisão de literatura

ABSTRACT

Purpose: To describe the prevalence of auditory and vestibular dysfunction in individuals with systemic sclerosis (SS) and the hypotheses to explain these changes. Research strategy: We performed a systematic review without meta-analysis from PubMed, LILACS, Web of Science, SciELO and SCOPUS databases, using a combination of keywords “systemic sclerosis AND balance OR vestibular” and “systemic sclerosis AND hearing OR auditory.” Selection criteria: We included articles published in Portuguese, Spanish or English until December 2011 and reviews, letters, and editorials were excluded. We found 254 articles, out of which 10 were selected. Data analysis: The study design was described, and the characteristics and frequency of the auditory and vestibular dysfunctions in these individuals were listed. Afterwards, we investigated the hypothesis built by the authors to explain the auditory and vestibular dysfunctions in SS. Results: Hearing loss was the most common finding, with prevalence ranging from 20 to 77%, being bilateral sensorineural the most frequent type. It is hypothesized that the hearing impairment in SS is due to vascular changes in the cochlea. The prevalence of vestibular disorders ranged from 11 to 63%, and the most frequent findings were changes in caloric testing, positional nystagmus, impaired oculocephalic response, changes in clinical tests of sensory interaction, and benign paroxysmal positional vertigo. Conclusion: High prevalence of auditory and vestibular dysfunctions in patients with SS was observed. Conducting further research can assist in early identification of these abnormalities, provide resources for professionals who work with these patients, and contribute to improving the quality of life of these individuals.

RESUMO

Objetivo: Descrever a prevalência das alterações auditivas e vestibulares em indivíduos com Esclerose Sistêmica (ES) e as hipóteses elencadas para explicar essas alterações. Estratégia de pesquisa: Revisão sistemática, sem meta-análise, a partir das bases de dados PubMed, LILACS, Web of Science, SciELO e SCOPUS, utilizando a combinação das palavras-chave “systemic sclerosis AND balance OR vestibular” e “systemic sclerosis AND hearing OR auditory”. Critérios de seleção: Foram incluídos artigos publicados em Português, Espanhol ou Inglês até dezembro de 2011 e excluídos os artigos de revisão de literatura, cartas e editoriais. Foram localizados 254 artigos e selecionados dez. Análise dos dados: Foi realizada a descrição do delineamento dos estudos e elencadas as características e frequência das alterações auditivas e vestibulares. Após, investigaram-se as hipóteses formuladas pelos autores para explicar o comprometimento auditivo e vestibular na ES. Resultados: A perda auditiva foi o achado mais comum, com prevalência de 20 a 77%, sendo o tipo sensorineural bilateral o mais frequente. Hipotetiza-se que o prejuízo auditivo na ES é decorrente de alterações vasculares na cóclea. A prevalência das alterações vestibulares variou de 11 a 63%, e os achados mais frequentes foram alterações na prova calórica, nistagmo de posicionamento, resposta oculocefálica anormal, alterações nos testes clínicos de integração sensorial e vertigem postural paroxística benigna. Conclusão: Elevada prevalência de alterações auditivas e vestibulares em pacientes com ES. A condução de novas investigações pode colaborar na identificação precoce dessas alterações, fornecer subsídios para os profissionais que atuam junto a esses pacientes e contribuir para a melhoria da qualidade de vida desses indivíduos.

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INTRODUCTION

Systemic sclerosis (SS), also known as systemic scleroderma, is an autoimmune connective tissue inflammatory disease, characterized by fibrosis of the skin, blood vessels, and visceral organs. It is a rare disease affecting 4–12 individuals per 1,000,000 inhabitants/year. It is more common among persons aged between 30 and 50 years, and it is at least three times more frequent among women, especially those at fertile age. The prevalence of the disease among women suggests the involvement of sex hormones on the development of SS. Despite being known since the 18th century, and even though it has been recognized as a multisystemic pathology since the 19th century, its etiology is unknown, and its pathogenesis is not clear. Several chemical agents (vinyl chloride and benzene), drugs (bleomycin and toxic oil syndrome), or viral infections have been mentioned as the main precipitating factors for SS. The occurrence of the disease in other members of the family is rare.

This condition is progressive and may cause death, being clinically characterized by skin thickening and problems in visceral organs, including the gastrointestinal tract, lungs, heart, and kidneys. Fibrosis and vascular occlusion occur in all the organs affected by SS. Because of vascular occlusion, the lumen of arteries and capillaries reduces and, as a consequence, isolated pulmonary hypertension takes place, as well as the reduction of capillary network (some vessels are obstructed and others are dilated, forming the telangiectasias), arterial hypertension due to the reduced renal artery lumen, myocardial perfusion disorder, and the Raynaud’s phenomenon.

Patients diagnosed with SS are classified in two subgroups, according to the extension of skin compromise: diffuse SS, in which skin, trunk and limbs, face, and neck are compromised; and limited SS, in which there are no trunk injuries, affecting only distal parts up to the knees and elbows, besides face and neck. The CREST syndrome (calcinosis, Raynaud’s phenomenon, esophageal dysmotility, sclerodactyly, and telangiectasia) is one form of limited SS.

In general, the disease has an insidious onset, and skin manifestations are followed or preceded by the Raynaud’s phenomenon and edematous fingers. Because multiple organs are affected, the disease can limit the relationships of the individuals and make social interactions more difficult, thus compromising the quality of life. A study conducted in 2012 reveals that 21–55% of the patients who have had the disease for approximately 10 years could not resume work due to their physical limitations. Literature has also shown the presence of auditory and vestibular symptoms and changes in autoimmune diseases such as SS. However, studies that investigate these changes in SS are scarce, and the understanding of its physiopathology is still unclear. Considering that hearing enables interaction and social experiences, there is a need to investigate the integrity of this function, to intervene and minimize the losses resulting from auditory and vestibular changes, whenever possible. Besides, no publications on this subject were found in Brazil. The information about auditory and vestibular impairments in patients with SS can contribute to the knowledge of gaps about the problem, thus providing subsidies to professionals who work with these patients, improving quality of life of these patients.

OBJECTIVE

The objective of this study was to conduct a systematic literature review to describe and verify the prevalence of auditory and vestibular changes among individuals with SS, as well as to list the hypotheses formulated to explain such changes.

RESEARCH STRATEGY

For this analysis, a systematic literature review, without meta-analysis, was conducted on studies that investigated auditory and vestibular changes in patients with SS. By checking PubMed, LILACS, Web of Science, SciELO, and SCOPUS, articles published until December 2011 were obtained by a combination of the following keywords in the abstracts: “systemic sclerosis AND balance OR vestibular” and “systemic sclerosis AND hearing OR auditory.” These keywords were selected after DeCS (Descritores em Ciências da Saúde) and MESH (Medical Subject Headings) were consulted. On the basis of articles that dealt with the studied topic, other free terms were also used.

SELECTION CRITERIA

The study included original research articles on auditory and vestibular changes in patients with SS published in Portuguese, Spanish, or English. All the literature reviews, letters, and editorials were excluded from the study. After reading their titles and abstracts, 15 articles were pre-selected. They all studied the matter using a script including location, period of publication, study design/base, and population. Afterwards, the characteristics of investigation and met the inclusion criteria defined for the study. Of the 15 articles, 5 were not available in the BIREME database; therefore, 10 articles were included in this study (Figure 1).

DATA ANALYSIS

For each selected article, the study design was described by using a script including location, period of publication, study design/base, and population. Afterwards, the characteristics and frequency of auditory and vestibular changes identified in individuals with SS were described. Then, the hypotheses described in the selected studies were presented to explain the physiopathology of auditory and vestibular changes in SS.

RESULTS

After consulting the electronic databases, 254 articles were found. However, only 10 were selected for this study. The most common reason for exclusion was the investigation of diagnosis, treatment, physiopathology, and other manifestations of SS that did not include auditory and vestibular changes (39%), followed by studies that reported other pathologies (28%), those in which the type of study did not meet the inclusion (16%) or language criteria (8%), and when the full text was not found (2%).
The earliest identified study was conducted in 1980, and 10 cases of patients diagnosed with SS who developed injury of the cranial nerve were reported\(^9\) (Chart 1). From 2006 onward, more publications on studies conducted in Europe, and, in Brazil, no studies analyzing auditory and vestibular compromise in SS were identified. It can be observed that in the past four decades there were only a few studies with the objective of investigating auditory and vestibular changes among individuals with SS. It is believed that because the disease is rare and its diagnosis is late, studies to investigate auditory and balance in these individuals become compromised. Besides, we should consider that the evolution, symptoms, limitations, and prognosis of the disease draw the attention of professionals who work with these patients towards restoring the functions and skills essential to maintain their lives. However, it is noteworthy that the integrity of hearing and balance enables these patients to socialize and gain autonomy, contributing to their well-being and quality of life.

The most used study design for describing auditory and vestibular changes in individuals with SS was the cross-sectional design, observed in five analyses\(^{10-14}\), followed by four case reports\(^{6,9,15,16}\), and one case-control study\(^{17}\). Because cross-sectional studies reported findings of a single moment, it was difficult to determine a temporal sequence between exposure to a specific factor and the subsequent development of the disease. This type of study is not considered to be the most appropriate or indicated one to investigate hearing and balance among individuals with SS. This is because there is a possibility that at the time of evaluation, auditory and vestibular changes had not manifested. Besides, it is impossible to ensure that SS was present before the development of hearing loss or vestibular changes. Therefore, and considering the progressive aspect of SS, more follow-up studies are required to establish a temporal sequence between the disease and the onset or aggravation of auditory and vestibular changes among individuals with SS.

The largest sample was observed in the study by Amor-Dorado et al.\(^{12}\), which included 42 individuals with SS, and the smallest sample was observed in the study by Maciaszczyk et al.\(^{13}\), including 20 people diagnosed with SS. The reduced number of the samples is probably because of the fact that SS is a rare and progressive disease, with several incapacitating manifestations leading to death.

Amor-Dorado et al.\(^{12,17}\) and Maciaszczyk et al.\(^{13}\) had differences in the study population with regard to the form of disease (limited or diffuse); however, only in the analysis by Amor-Dorado et al.\(^{12}\) results were presented by considering the type of disease. Prevalence of changes in the vestibular system was higher for patients with diffuse SS (Charts 1 and 2). Considering that the level of systemic manifestations is directly related to the form of disease presentation (limited or diffuse), it is hypothesized that auditory and vestibular changes also have these variations, which emphasizes the importance of distinguishing the study population according to the form of disease.

Only the studies by Berrettini et al.\(^{10}\) and Amor-Dorado et al.\(^{17}\) investigated auditory and vestibular involvement in patients with SS. However, six studies aimed at assessing or reporting only auditory changes in these patients\(^{6,9,11,13,15,16}\) and only two\(^{12,14}\) investigated vestibular dysfunctions (Chart 1).

It is possible to observe that symptoms of hearing loss, tinnitus, and vertigo were more frequently reported by individuals with SS. As to the presence of changes, hearing loss was the most common finding in the studies (Chart 2). However, it is important to consider that, of the 10 analyzed studies, 6 observed only hearing changes. Of the eight studies that investigated changes, all identified hearing loss in most individuals with SS, and the prevalence in this finding ranged from 20 to 77%. The type of hearing loss was bilateral sensorineural\(^{6,9,11,13,16,17}\) in most cases, corroborating the physiopathology of the systemic disease. However, mixed\(^{11,15,17}\) and unilateral\(^{13,17}\) hearing loss were also described in few cases.

The configuration and level of hearing loss were little explored in the studies. As this information is very important because it will guide the auditory rehabilitation process among these individuals, it should be included while conducting new studies. Amor-Dorado et al.\(^{17}\) described the flat hearing loss configuration in 13 individuals and high-frequency hearing loss in 14, accounting for...
for 27 patients with sensorineural hearing loss (SNHL). Deroee et al. (16) observed high-frequency hearing loss in one of the ears and severe loss in another. In a study by Maciaszczyk et al. (13), most individuals in the case group presented moderate hearing loss (Chart 2).

Regarding vestibular changes, prevalence ranged from 11 to 63%, and the most common findings were changes in caloric testing, positional nystagmus, abnormal oculocephalic reflex, changes in clinical tests of sensory integration, and benign paroxysmal positional vertigo (6,10,12,17). The prevalence of vestibular changes deserves the attention of professionals assist these patients, because these changes lead to balance changes, which, together with locomotive changes (characteristics of the disease), can be incapacitating and compromise the autonomy of patients with SS.

The analyzed studies have tried to elucidate the frequency and magnitude of auditory and vestibular changes by evaluating the peripheral and central systems. The basic audiological evaluation (tone audiometry, vocal audiometry, and acoustic immittance) was the most used evaluation method to identify these changes in individuals with SS (6,10,11,13,15,17). However, the brainstem auditory-evoked potentials (BAEP) and vestibular evaluation were also used for diagnostic investigation (6,10,13,17). The highest prevalence of peripheral auditory changes can be the reflex of the assessment instrument used in the studies. Therefore, it is important that new studies be conducted with the objective of investigating the presence of auditory changes in the central system, as well as changes related to balance in individuals with SS.

Regarding the biological plausibility of auditory changes in individuals with SS, only one study suggested the possible mechanisms. According to Deroee et al. (16), hearing damage in patients with SS first affects high-frequency hearing, and patients may not be aware of their hearing loss due to the severity of other manifestations of the disease. These authors suggest that the inflammation of small vessels in the cochlear nerve and the reduced blood flow to the cochlea may be the beginning of the inner ear pathogenesis in autoimmune diseases.
The inner ear compromise, especially the cochlea, may be a consequence of a vascular change, once this change is the base of SS physiopathology. The irregular structure of blood vessels associated with the reduced density of capillaries may result in decreased blood flow and, consequently, hypoxia\(^{(16)}\). The cochlea is a highly sensitive organ to blood changes, and the hypoxia resulting from vascular changes of the disease may lead to the death of hair cells\(^{(19)}\). Clinically, these changes are expressed by SNHL, which corroborates the findings of the studies that indicate lower prevalence of this type of hearing loss.

Literature also describes mixed hearing loss in these individuals; however, no biological mechanisms that could explain this implication on SS have been reported. However, studies show the occurrence of conductive hearing loss in other autoimmune diseases such as recidivant polychondritis, thus attributing this change to the expansion of the inflammatory process to the middle ear and auditory tube\(^{(19)}\).

In one of the reviewed articles, the authors\(^{(11)}\) identified the presence of eustachian tube pathology in 3 of the 30 evaluated patients. This change may explain the presence of the conductive component in the hearing loss of patients with ES. As the physiopathology of SS involves not only vascular injuries but also tissue and joint injuries, there is also a possibility of middle ear involvement, because of the mechanical impairment of structures composing the tympanic-ossicle system, leading to changes in sound conduction.

### Chart 2. Symptoms and auditory and vestibular findings in systemic sclerosis

<table>
<thead>
<tr>
<th>Author</th>
<th>Symptoms</th>
<th>Auditory and vestibular changes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Santarelli et al.(^{(10)})</td>
<td>Insidious onset of symptoms, with progressive difficulty to understand the speech</td>
<td>Slightly higher auditory thresholds 1 year after the onset of the disease, and significant aggravation after 2 years; normal tympanometry and absence of bilateral acoustic reflexes; bilateral absence of speech intelligibility; DPOAE at high frequencies, and absence in low frequencies 2 years after the onset of the disease; normal vestibular evaluation; after cochlear implant, 70% of speech recognition</td>
</tr>
<tr>
<td>Teasdall et al.(^{(15)})</td>
<td>Of the 10 patients, 3 presented tinnitus</td>
<td>Bilateral HS in 2 of 10 patients</td>
</tr>
<tr>
<td>Berrettini et al.(^{(10)})</td>
<td>Of the 37 patients, 11 complained of tinnitus; 4 complained of vestibular dysfunctions; and 8 complained of auditory and vestibular dysfunctions</td>
<td>SNHL in 10 and mixed HL in 4 cases; unilateral HL in one case; in patients with mixed HL, acoustic immittance findings indicate tympanosclerosis; two of them presented positional nystagmus; one had unilateral vestibular dysfunction, and one had bilateral deficit after caloric stimulation</td>
</tr>
<tr>
<td>Kastanioudakis et al.(^{(11)})</td>
<td>Not reported</td>
<td>Six of them had SNHL, and one had mixed HL; out of these, one belonged to group A, one to group B, four to group C, and one to group D; 10% of these had bilateral eustachian tube pathology; vocal audiometry compatible with cochlear disorder and tympanometry without changes</td>
</tr>
<tr>
<td>Amor-Dorado et al.(^{(17)})</td>
<td>Not reported</td>
<td>Seven were diagnosed with BPPV; two patients with limited SS had involvement of the posterior semicircular canal; three with limited SS, and two with diffuse SS presented compromised horizontal semicircular canal; changes in sensory integration tests observed in 20 patients</td>
</tr>
<tr>
<td>Maciaszczyk et al.(^{(18)})</td>
<td>Of the 20 patients, 60% reported vertigo, 55% had headaches, 50% had tinnitus, 40% had hyperacusis, 40% had hearing loss, and 30% had auricular plentitude</td>
<td>Eight had SNHL, all were women aged between 42 and 77 years; two had unilateral SNHL; mild for three of them, and moderate for the rest. Five of them presented acoustic reflex, one did not, and two had absence in frequencies of 2 or 4 kHz; BAEP had no changes</td>
</tr>
<tr>
<td>Bassyouni et al.(^{(14)})</td>
<td>Not reported</td>
<td>Vestibular dysfunction in 33% of the sample; no relationship was observed between vestibular compromise and age, duration and other clinical parameters of the disease</td>
</tr>
<tr>
<td>Abou-Taleb and Linthicum(^{(10)})</td>
<td>Progressive auditory loss</td>
<td>Bilateral mixed HL; 68% speech discrimination in the right ear; 78% in the left ear</td>
</tr>
<tr>
<td>Deroee et al.(^{(16)})</td>
<td>Sudden bilateral hearing loss, especially at the left</td>
<td>High-frequency SNHL in the right ear direita and severe HL in the left ear; 100% speech discrimination in the right ear and 0% in the left ear; normal bilateral tympanometry; 1 year after treatment for SS, improved HL and neuropathy, and increasing speech discrimination — to 68% in the left ear</td>
</tr>
<tr>
<td>Amor-Dorado et al.(^{(17)})</td>
<td>Of the 35 patients, 19 (54%) reported HL</td>
<td>HL in 27 patients, being 24 with SNHL (11 asymmetric and 13 symmetric), and 3 with mixed HL; 13 had flat configuration and 14 had high-frequency loss; 2 had unilateral HL; 2 had absent bilateral acoustic reflex, and 11 had unilateral; 10 presented abnormal oculocephalic response, and 9, positional nystagmus; changes in caloric testing and clinical sensory integration tests were found in 31 and 46% of the patients, respectively</td>
</tr>
</tbody>
</table>

**Caption:** HL = hearing loss; SNHL = sensorineural hearing loss; DPOAE = distortion product otoacoustic emissions; BPPV: benign paroxysmal positional vertigo; SS: systemic sclerosis; BAEP = brainstem auditory-evoked potentials
The physiopathology of vestibular changes was not discussed in any of the articles included in this study. However, it is important to investigate how these changes occur, to intervene early, when possible, thus contributing to improvement in the quality of life of individuals who present with dysfunctions in this system, once they can empower locomotion limitations in people with SS.

Auditory and vestibular changes, when added to manifestations resulting from the disease, may lead to even significant damage when it comes to the socialization of these individuals. Therefore, knowing about these changes can indicate the need to include the evaluation of peripheral and central auditory system in the diagnosis protocol, as well as the follow-up of these people, to detect such changes and adopt rehabilitation procedures or medicines that can be applied early and be effective.

CONCLUSION

The results of the analyzed studies indicate auditory and vestibular compromise in patients with SS, and it is possible to observer higher prevalence of bilateral SNHL. The configuration and level of hearing loss were considered in a few studies, and results were inconsistent.

Auditory symptoms may precede the manifestations of the autoimmune disease; therefore, they can be useful in the early diagnosis, when SS is suspected. The early diagnosis and comprehension of the physiopathology of auditory and vestibular changes caused by SS are essential for taking necessary measures for their prevention and rehabilitation.

New investigations should be conducted with the objective of clarifying the relation between the configuration and level of hearing loss with the severity and form of disease presentation (limited or diffuse), contributing to the knowledge of whether or not the progression and form of disease affect the level of auditory and vestibular manifestations.

The analysis of the selected articles for this review indicates the importance of standardizing the measurement of study outcomes, so that it is possible to perform future meta-analyses that will contribute to practices based on evidence. Besides, the need for studies investigating the physiopathology of auditory and vestibular changes among individuals with SS is worth mentioning.

REFERENCES