Primary Sjogren’s syndrome presenting as ptosis and eyelid swelling: A case report.

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Introduction:

Primary Sjogren’s syndrome (pSS) is an idiopathic, autoimmune disorder characterized by both localized and systemic manifestations. The most common clinical presentations include the “sicca” or “dryness” symptoms such as xerostomia and keratoconjunctivitis sicca due to diminished salivary gland and lacrimal gland function, respectively. Patients may also present with nonspecific symptoms such as fatigue and arthralgia, as well as pulmonary, gastrointestinal, neurological and renal involvement. Secondary Sjogren’s syndrome can co-exist with other autoimmune disorders such as rheumatoid arthritis (RA) and systemic lupus erythematosus (SLE).

According to literature, ocular signs and symptoms in pSS range from foreign body sensations, punctate or filamentous keratitis, to overt seborrheic blepharitis and eyelid swelling is reported to be rare. We report an unusual/rare presentation of Sjogren’s syndrome in which the patient developed severe evolving mechanical ptosis and eyelid swelling to the extent that it impaired her ability to lift open her eyes and see, along with a positive anti-acetylcholine receptor antibody test.

Case presentation:

A 27-year-old woman presented to our Rheumatology department with the complaint of periorbital swelling for the past five months. Prior to coming to our clinic, she presented to the Neurology department of one of our allied hospitals, Benazir Bhutto Hospital with a complaint of drooping of eyelids for the last three months where she was presumed to be a case of MG on the basis of positive anti-acetylcholine receptor antibodies. She was given a therapeutic trial of pyridostigmine but she failed to respond to the treatment and was referred to our rheumatology clinic for an autoimmune workup.

The periorbital swelling was gradual, bilateral, and had progressed to the degree that the patient could not lift open her eyelids and had to open them with her fingers to be able to see. Figure 1 shows the extent of her ptosis and eyelid swelling. It was not associated with any pain, redness, discharge, loss of vision, or diplopia. Upon systemic inquiry, she complained of undocumented weight loss, productive cough for the past three months, dry gritty eyes, dry mouth, hair loss, and a photosensitive facial rash. There was no significant family history of disease other than her father who was treated for tuberculosis 10 years back.

On clinical examination, our patient had periorbital puffiness with closed palpebral fissures, there was no redness or discharge, and she had normal extraocular eye movements. There was no assessable lymphadenopathy and none of the salivary glands were palpable. On oral examination, Schirmer test was positive with
less than five mm tear production in five minutes which is suggestive of dry eyes. On respiratory examination, bilateral coarse crepitations were heard. Her neurological examination was unremarkable and muscle strength and endurance in all muscle groups was normal which was in contrast to the symptoms of MG. We could not perform all the provocative maneuvers for ocular MG (sustained upgazed, Herring’s sign, peek sign) because of marked eyelid swelling that impaired lifted the eyelids. There was no fatigable diplopia. The rest of the systemic examination was normal.

Peripheral blood test results were as follows: hemoglobin 14.0 g/dl, total leucocyte count 14000/mm³, neutrophils 70.2%, lymphocytes 25.1%, mixed 5.7%, red blood cells 4.21 million/mm³, platelets 223000/mm³, hematocrit 38.9%, erythrocyte sedimentation rate 44mm/hour, C-reactive protein 4.1 mg/L, Serum biochemical testing results were as follows: serum urea 3.0 mmol/L, serum creatinine 55 umol/L, serum sodium 137 mmol/L, serum potassium 4.6 mmol/L, alanine aminotransferase 15 U/L, alkaline phosphatase 86 U/L, total bilirubin 0.4 mg/dl. The results of her autoimmune workup are displayed in Table 1. Repetitive nerve stimulation showed no significant decremental or incremental response which helped rule out generalized myasthenia gravis but not ocular myasthenia.

We proceeded with a High-Resolution Computed Tomography (HRCT) scan of her chest which revealed bilateral cylindrical bronchiectasis involving all lung lobes, consolidation in the right lower lobe, and mild pleural effusion. PCR was negative for COVID-19, sputum for Gene Xpert, and QUANTI-FERON TB gold test were also negative. Sputum culture revealed heavy growth of Candida and Klebsiella Pneumoniae. Magnetic Resonance Imaging MRI of the brain and orbits revealed bilateral symmetrical homogenous enlargement of lacrimal glands with accompanied symmetrical swelling and edematous changes in bilateral pre-septal and para-septal soft tissues. Figure 2 shows these MRI images prior to commencing therapy.

We considered primary Sjogren syndrome, systemic lupus erythematosus as she gave a history of a photosensitive facial rash, ocular myasthenia with idiopathic orbital inflammation as she gave a history of ptosis and periorbital inflammation, sarcoidosis, ANCA associated vasculitis and IgG4 related orbital pseudotumor as our differential diagnoses. We had ruled out SLE, sarcoidosis, ANCA associated vasculitis on the basis of autoimmune investigations all of which were negative for corresponding antibodies. Myasthenia Gravis was ruled out on the basis of electrodiagnostic studies. As she tested positive for Anti SSA antibodies, we proceeded with a lip biopsy to confirm the diagnosis of primary Sjogren’s syndrome and to rule out IgG4-related disease. The lip biopsy indicated normal-looking mucinous acini with lymphocytic infiltrate. There were four microscopic foci of lymphoid aggregates per 4 mm², with no evidence of plasma cells and fibrosis (hence ruling out IgG4-related disease), suggestive of primary Sjogren syndrome (pSS). Figure 3 shows the histopathological images of her lip biopsy.

A pulmonology consultation was sought and she was started on an antibiotic cover to control infective pulmonary etiology prior to beginning immunosuppressive treatment for pSS. The patient was advised Acetylcysteine, Azithromycin 250mg for three months along with Influenza and Pneumococcal vaccine. After her chest infection subsided, we started her on Hydroxychloroquine 200 mg/day, Azathioprine 2.5 mg/kg/day, and Prednisolone 0.5 mg/kg/day. Cyclophosphamide was deferred because we believed it would worsen her bronchiectasis. The patient was referred for an ophthalmology consultation for her persistent eyelid swelling and the consensus was to control her primary disease process which would ultimately resolve her swelling. As the persistent swelling was hampering her ability to see, she underwent a sling procedure to open her eyelids. Figure 4 shows an image of the patient after her sling procedure. Though the swelling had considerably subsided on therapy, her lacrimal gland were persistently inflamed, we shifted her towards a biological agent Rituximab, 1g IV infusion. She has received two doses and the next dose is scheduled for after 6 months. Figures 5 and 6 show MRI of her brain and orbits after starting treatment.

Discussion

This case highlights a rare presentation of pSS. The patient developed bilateral ptosis along with prominent eyelid swelling which evolved to the extent that she could not lift her eyelids to see. Eyelid swelling has been reported in primary Sjogren’s syndrome but to our knowledge, this is the first case in which the swelling...
has evolved to the extent that impairs the patient’s ability to lift open her eyes. Sjögren syndrome has a multitude of ophthalmological manifestations ranging from chronic conjunctivitis, sterile keratolysis, and non-healing corneal ulcers, but lacrimal gland involvement is reported to be rare.

Due to her initial presentation with ptosis and eyelid swelling along with a positive anti-acetylcholine receptor antibody test, we presumed her to be a case of MG and she was given a therapeutic trial of pyridostigmine to which she failed to respond. This made us reconsider her initial diagnosis of MG and we started her autoimmunity workup. Our case is unique in this aspect as well because MG and Sjögren’s syndrome very rarely co-exist in the same patient and very few cases have been reported in the current literature. There are many diagnostic modalities for MG including Ice pack test, Tensilon test, Anti-acetylcholine receptor antibodies, Repetitive nerve stimulation, and Electromyography, all with varying sensitivity and specificity. Of these, the positive titer of anti-acetylcholine receptor antibodies has the highest specificity (97-99%) for the diagnosis of MG. Our patient had no typical signs or symptoms of MG aside from ptosis but she had a positive titer of anti-acetylcholine receptor antibodies. We proceeded with repetitive nerve stimulation tests which showed no incremental or decremental response hence ruling out generalized MG but not ocular MG. A positive titer for this assay in the absence of typical signs and symptoms, a negative therapeutic trial of pyridostigmine and electrodiagnostic studies ruling out generalized MG could mean that the test result was spuriously positive in our patient. The pathogenesis of false positive antibody test results in the context of autoimmune diseases is very interesting and has been reported in a number of other autoimmune diseases as well. These false positive antibody results can be attributed to the phenomenon of heterophil antibody interference in which some of the patient’s antibodies react with the immunometric sandwich assays and cross-link the assay antibodies yielding a false positive test result. Heterophil antibody interference has been well documented in the literature and there are examples of catastrophic patient outcomes due to this phenomenon such as the HCG scandal in which false positive HCG levels, later attributed to heterophil antibodies, caused unnecessary treatment in a number of women. Heterophile antibodies are common in autoimmune diseases and should be suspected in cases of positive antibody titers in absence of a solid clinical picture of the disease and clinicians need to be made aware of this phenomenon. If such a spurious test result is encountered either on a single occasion or repeatedly, it should be followed by confirmatory testing and taking the clinical context into account. Our case thus highlights the importance of interpreting antibody test results in the context of clinical findings as our patient did not have MG despite a positive anti-acetylcholine receptor antibody test result indicated by a negative therapeutic trial of pyridostigmine and electrodiagnostic studies.

The recent 2016 ACR-EULAR Classification Criteria for primary Sjögren’s Syndrome is most commonly used to establish a diagnosis of pSS. On the basis of focal lymphocytic salalndenitis upon labial salivary gland biopsy, positive Anti-SSA(RO) antibodies, and positive Schirmer test, our patient scored seven out of nine where a score of 4 or more is required for a definite diagnosis of pSS.

Based on the HRCT findings, our patient was also had co-existent bronchiectasis, because of which she could not be started on immunosuppressive therapy for pSS immediately until the pulmonary infection had been resolved. Pulmonary involvement is an extraglandular manifestation of pSS and it can be in the form of both interstitial parenchymal disease and airway disease. Bronchiolitis and bronchiectasis are the common airway lung diseases seen in pSS. The prevalence of bronchiectasis is up to 10% with involvement of the inferior lobes being more common. Cylindrical bronchiectasis is the most common type seen and patients of bronchiectasis have a higher frequency of respiratory infections and pneumonia. Our patient similarly had bilateral cylindrical bronchiectasis involving all of the lung lobes along with a positive sputum culture for Klebsiella pneumoniae thus indicating an infective etiology coexisting with her autoimmune pulmonary manifestation. The pathogenesis of pulmonary involvement in pSS seems to involve epithelial damage due to any environmental factor such as infection or an extension of the primary immune response in salivary glands, followed by epitope spreading, antigen presentation and lymphocyte activation, formation of antibodies and release of cytokines leading to an inflammatory state damaging airways and lung parenchyma. Pulmonary involvement also proved to be a challenge in management as we had to control the infective etiology with antibiotics prior to commencing immunosuppressive therapy to avoid a flare-up of the lung disease.
Conclusion:
Our case highlights a unique ophthalmological manifestation of pSS and provides key insight into managing this autoimmune disease. We must always look at the clinical picture of a disease instead of relying solely on investigations as elucidated in our case where a positive anti-acetylcholine receptor antibody test led to a misleading diagnosis of MG which was ruled out subsequently.

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None.

Conflict of Interest:
The authors declare that there are no conflicts of interest in the publication of this paper.

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References


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