Primary Sjögren’s syndrome is an autoimmune disorder with external exocrine glands dysfunction and multiorgan involvement. The pathogenesis of primary Sjögren’s syndrome is still unclear; however, our knowledge of the involvement of different cells (e.g., B and T cells, macrophages and dendritic cells) and pathways (BAFF/APRIL and interferons) leading to the development of autoimmunity is continually expanding. For clinicians, the most frequent symptoms are dryness of eyes and mouth, but often the patients have musculoskeletal symptoms and systemic manifestations. However, the increased risk of lymphoproliferative disorders in this group of patients, most commonly B-cell marginal zone lymphoma, is particularly important. Recent separation of IgG4-related diseases and attempts to create further diagnostic criteria for pSS testify to the difficulties, and at the same time a large interest, in understanding the disease so as to allow the effective treatment. This article draws attention to the problems faced by the clinician wishing to securely identify pSS by using accurate laboratory biomarkers and useful imaging tools and predict the development of complications associated with this, still not fully understood, autoimmune disease.

Keywords Sjögren’s syndrome · Criteria · New therapies

Introduction

It is known that primary Sjögren’s syndrome (pSS) is considered as an autoimmune disease based on lymphocyte B

hyper-reactivity and polyclonal production of immunoglobulins, and as a consequence, autoantibodies against pSS affected exocrine glands, especially lacrimal glands and salivary glands, but also other external glands such as the pancreas, mucous glands of the gastrointestinal and respiratory tract or bile secretion. In some patients, abnormal H+ secretion in the distal tubules has also been observed and caused distal renal tubular acidosis (type 1 RTA). In connection with the clinical symptoms which will result from the degree of attachment of these glands and epithelial injury, the pSS often uses the term “autoimmune epithelitis” or “autoimmune exocrinopathy” for better imaging of primary initiation sites of inflammation and autoimmunization.

A brief history

First lacrimal and salivary glands enlargement was described in a lecture of a Polish surgeon Jan Mikulicz-Radecki in 1888. In 1925, French dermatologists Henri Gougerot described a few cases of atrophy of the salivary glands with dryness of eyes, mouth and vagina. Later in 1933, Swedish ophthalmologists Henrik Sjögren in his doctoral thesis described keratoconjunctivitis sicca, and his description became a basis for pSS picture.

Epidemiology

It is estimated that primary Sjögren’s syndrome occurs from 0.1 to 3.0 % in general population. The disease is more common for women (female/male ratio 9:1), mainly between the ages of 40–60, with the disease most frequently occurring around 50 years of age.

Pathogenesis of pSS is not clear, nowadays several factors responsible for the development of the disease, such as
genetic factors—genes implicated in B cell or B-cell activation factor (BAFF) known also as B-Lymphocyte Stimulator (BLyS), lymphotoxin α and β and TNF (tumor necrosis factor) are taken into account. Furthermore, it is presumed that genetic predisposition to increase in type I interferon (IFN) may explain the IFN signature and activation of type I IFN signaling in salivary gland and peripheral blood in pSS patients. The HLA-B8, of HLA-Dw3 and HLA-DR3 and DRw52 have also been reported in pSS patients [1, 2]. Another factor responsible for the development of pSS is an infection caused mainly by Epstein–Barr virus (EBV), human T-cell lymphotropic virus type-1 (HTLV-1), Cytomegalovirus (CMV) and Hepatitis C virus (HCV). Also neuroendocrine disturbances with sex hormones and its receptors dependent on hypothalamic–pituitary–adrenal axis (HPA or HTPA axis) interfere with the ratio of estrogen to androgens and affect steroid-dependent cells like epithelial cells and other cells involved in the immune response [3]. In pSS patients, lower basal secretion of ACTH and cortisol has been found. In addition, hypothalamic–pituitary–gonadal (HPG) axis by estrogen deficiency can be responsible for a local autoimmune exocrinopathy [4]. It is assumed that the infection as the trigger (most commonly viral) and other environmental factors caused the disorganisation of epithelial cells. First, due to innate immune response, virus infection is recognized by pattern recognition receptor (PRR) and activates toll-like receptors (TLR) pathway (e.g., TLR 3, 7, 9). After the activation, the innate immune response TLR, cell apoptosis and SS-A RNA complexes stimulate plasmocytic dendritic cells (pDCs) which produce high level of interferons (IFNs) and IFNs as strong stimulators of BAFF production by epithelial cells, monocytes and neutrophils dendritic cells leading to proliferation and differentation of B cells and production of autoantibodies. It is suggested that glandular cell apoptosis triggered by viral infection (EBV, HCV and HIV) leads to progressive damage of glands and their dysfunction with reduced secretion and the appearance of classic clinical symptoms. Damaged epithelial cells release autoantigens, especially Ro/SSA and La/SSB, which create autoimmunity and autoantibodies secretions. The presence of Ro/SS-A antibodies (anti-Ro52 and anti-Ro60) is correlated with longer duration of pSS, bigger destruction of the glands and extraglandular manifestation. Plasmocytoid dendritic cells (pDCs) migrate into the site of damage. pDC is the source of type I INF and initiates the activation of B cell by B-cell activation factor (BAFF) pathway. However, epithelium is infiltrated mainly by CD 4+ limphocytes T subtype and immune response is balanced toward Th1 response and also Th17—with interleukin 17(IL-17) as a main cytokine. Th1 cells produce interferon gamma (IFN γ) which induces plasminogen activator system and together with IL-17 promotes local inflammation. In advanced stages of inflammation, B cells have been detected in salivary glands or other places in exocrine system. Recent study suggests that IL-7 from IL-7+ peripheral blood T cells may contribute to the stimulation of Th1 and Th17 cytokines [5]. It has been recently discovered that newly identified cytokine IL-34 promotes overexpression of CD 14+ monocytes in salivary glands. IL-34 and CSF-1 (colony stimulating factor-1) stimulate survival, proliferation and differentiation of monocytes, macrophages, dendritic cells, Langerhans cells and osteoclasts—cells which have the ability to phagocyte [6]. Studies have shown that also IL-21, cytokine produced by activated CD 4+ T limphocytes and NKT cells play a role in pathogenesis of pSS and correlate with lymphocytes infiltration of salivary gland as well as the presence of autoantibodies and level of gammaglobulins. This cytokine stimulates TH1 and Th17 differentiation and synergistically with BAFF stimulates B-cells differentiation [7]. As in many autoimmune diseases, the increased level of IL-6 has been found in pSS. IL-6 is one more cytokine which enhances lymphocytes B growth and maturation. As well as IL-6, it has a strong influence on persistence of inflammation [8]. The production of IL-6 by monocytes in pSS is strongly stimulated by INF γ. The increased activity of IL-6 may influence the clinical symptoms, and increased IL-6 has been reported in saliva, serum and tears [8]. The break of T cells tolerance by viral infection leads to the activation of self-reactive T cells which by BAFF stimulate activation and maturation of B cells. Extended B-cell activation by BAFF generates self-reactive memory B cells which are an antigen-presenting cells to T cells. The crucial role of B limphocytes in pSS seems obvious—all pathways lead to powerful stimulation of B-cells maturation and to their hyper-reactivity, polyclonal production of gammaglobulins and the emergence of autoantibodies, which are B cell related markers for pSS. At first stage of inflammation, macrophages and T cells are infiltrating salivary glands, but over the time in the glands accumulate memory B cells and their level decreases in peripheral blood, while the level of naïve and transitional B cells increases. Interesting work presented by Bohnhorst et al. [10] showed disturbances in subset of B cells in pSS and differences in proportions between naïve B cells and memory cells in peripheral blood [9, 11]. Baldini and colleagues [12] assessed a prevalence of Vitamin D deficiency in early stage of the disease, considered the impact of hypovitaminosis D in clinical symptoms and disease activity, expressed by the symptoms of dryness and concluded that hypovitaminosis D is relatively frequent in patients with pSS. The investigators speculated that vitamin D may have the effect on Th1 cells and regulatory limphocytes, dendritic cells and CD 8+ T cells and plays a role in pSS pathogenesis. But the fact that vitamin D deficiency may be an independent and associated with reduced supply
and reduced synthesis of this vitamin in relation to the living conditions and nutrition cannot be exclude. In another work, Agmon Levin et al. [13] showed the relationship between reduction of level of vitamin D and the occurrence of peripheral neuropathy and also of non-Hodgkin lymphoma. The pathogenesis of Sjögren’s syndrome schematic is shown in Fig. 1

**Autoantigens and autoantibodies**

The most commonly present autoantibodies in pSS are antibodies against cellular ribonucleoprotein antigens Ro/anti-SS-A and LA/SS-B. These autoantibodies are not specific for pSS, only but their presence is associated with severe glandular manifestations, longer duration of the disease, presence of splenomegaly, lymphadenopathy, vasculitis and high infiltration of salivary glands. Ro52 antibodies were reported in about 80% of pSS patients. In pSS, rheumatoid factor (RF) correlating with early stage of disease and onset of the disease at a younger age is often appearing. RF is present in about 74% of patients with pSS and may also be associated with extraglandular symptoms such as arthritis. In 17% of pSS anti-centromere antibodies (ACA), which are more specific for limited systemic sclerosis, were found. Their presence may signal an overlap syndrome [14].

The prevalence of antinuclear antibodies (ANAs) in pSS reaches 80%, together with RF, in the case of the absence of anti-Ro/La along with histopathological confirmation or confirmation of ocular lesions allows make pSS diagnosis according to the ACR criteria. ANAs are also a predictor of internal organ involvement and the development of lymphoproliferative disorders. In pSS, M3 muscarinic acetylcholine receptor (M3R) is expressed in exocrine glands and its activation is important in the process of exocrine secretion. Sumida and colleagues found that M3R reactive T cells (Th1 and Th17) occur in 40% in peripheral blood of patients with pSS and autoantibodies against M3R may play an important role in the development of autoimmune sialadenitis [15, 16]. In pSS patients, anti-mitochondrial antibodies (AMA), which are characteristic for primary biliary cirrhosis (PBC), can occur. In these patients, changes in the liver are more likely to appear, with periductal lymphocytic infiltrating analogously the changes in salivary glands. Abnormal liver function test (increased AST and/or ALT, AP and/or bilirubin) is common in pSS as extraglandular manifestation of the disease as well as overlapping with another disease as PBC or autoimmune hepatitis [16, 17]. Anti-smooth muscle antibodies (ASMA) are characteristic for autoimmune type 1 hepatitis which can also be found in pSS patients [18]. Also autoantibodies against carbonic anhydrase (anti-CAII) are present in pSS and are associated with distal renal tubular acidosis [19]. The anti-cyclic citrullinated peptide antibodies (anti-CCP), which are highly specific for RA, are not frequent in patient with pSS. The presence of anti-CCP antibodies may be associated with joint erosions and synovitis. Also it cannot be excluded that the patient who has anti-CCP antibodies and rheumatoid factor (RF-IgM) will develop symptoms of RA in the future [20]. In addition, autoantibodies with direct cytoskeletal proteins-α and...
β fodrin and proteasomes have been found in patients with pSS. The current studies showed additional autoantibodies in pSS to salivary gland as follows: protein 1 (SP-1), carbonic anhydrase 6 (CA6) and parotid secretory protein (PSP) [21]. Organ-specific antibodies, such as anti-thyroglobulin (anti-TG) and anti-thyroid peroxidase antibodies (anti-TPO), most characteristic for autoimmune thyroiditis, were also found. D’Arbonneau and colleagues reported that thyroid disease was more frequent in pSS patients in up to 30 % of patients compared with 4 % of the control group [22]. However, the presence of anti-TPO and TG antibodies in pSS may also not be related to the coexistence of Hashimoto’s disease [23].

**Clinical picture of the disease**

The clinical picture of pSS includes symptoms of dryness and extraglandular manifestations with organs involvement. Patients’ complaints of the dryness are the basis for suspicion. The dryness concerns first of all the mouth, eyes and vagina—the latter especially in postmenopausal women. The clinical signs are presented in Table 1. Among laboratory findings, cytopenias (leukopenia, anemia and low platelet count) are the most typical. Elevated erythrocyte sedimentation rate (ESR), hypergammaglobulinemia, ANAs, RF, anti-Ro/SS-A, anti-B/SSB, decreased level of

| Table 1 Clinical features of primary Sjögren’s syndrome |
|--------------------------------------------------------|
| **Dryness**                                            |
| Xerostomia (dry mouth)                                 |
| Diminished secretion of saliva                          |
| Troubles with swallowing                                |
| Dental caries                                            |
| Fungal and bacterial infections                         |
| Xerophthalmia (dry eyes)                               |
| Persistent irritation keratoconjunctivitis sicca        |
| Destruction of cornea                                   |
| Decrease of vision                                      |
| **Extraglandular manifestations**                      |
| General symptoms                                        |
| Fatigue                                                |
| Fever                                                  |
| Weight loss                                            |
| Musculoskeletal features                               |
| Myalgia                                                |
| Arthralgia                                             |
| Arthritis—nonerosive, also rheumatoid arthritis—like myositis |
| Respiratory tract                                      |
| Cough—dryness of trachea or bronchitis                 |
| Intestinal like disease                                |
| Gastrointestinal system                                |
| Dysphagia (dryness of pharynx and esophagus), gastrointestinal reflux |
| Chronic gastritis with atrophy                         |
| Liver involving (rather mild)                          |
| Symptoms of PBC and AIH (autoimmune hepatitis)         |
| Celiac-like disease                                    |
| Urinary tract                                          |
| Distal renal tubular acidosis (RTA type 1)             |
| Nephrocalcinosis (in same cases due to RTA)            |
| Nepritis/glomerulonephritis                            |
| Chronic renal insufficiency                            |
| Vessels and skin                                       |
| Reynaud’s phenomenon (scleroderma like patterns with ACA antibodies) |
| Vasculitis                                             |
| Urticaria, palpable purpura, annular erythema          |
| Neurological manifestations                             |
| Peripheral sensory or motor-sensory polyneuropathy,     |
| Cranial neuropathy,                                    |
| Mononeuritis multiplex                                  |
| Sensonarineural hearing loss                            |
| SM-like syndrome                                       |
| Psychiatric                                            |
| Depression                                             |
| Anxiety                                                |
| Cardiac                                                |
| Pericarditis                                           |
| Pulmonary hypertension                                 |
Complement component C4 and cryoglobulins are more often laboratory abnormalities in patients with pSS.

Classification criteria for pSS and tools useful in the diagnosis

Classification criteria of pSS have evolved from 1960s till new proposed in 2012. Since 2002, clinicians used criteria established and revised by American–European Consensus Group (AECG), which are based on six points: subjective and objectives features of dry eyes and dry mouth, the patient’s need to fulfill at least three subjective or two objective features and the presence of Ro(SS-A) or La(SS-B) antibodies or both, or inflammation in minor salivary glands with focus score (FS) equal or more than 1. FS is defined as 50 inflammatory cells in a 4-mm² glandular section in material from minor salivary gland biopsy (SGB). Newly proposed ACR classification criteria for pSS assumed that in the cases with Ro(SS-A) and La(SS-B) negative, but with symptoms suggestive for pSS and FS >1 in SGB the presence of ANAs titer above/equal to 1:320, and the presence of rheumatoid factor is sufficient to establish a diagnosis [24, 25]. This year Rasmussen et al. [26] compared new ACR criteria and AECG criteria and showed no significant differences and assumed ACR criteria sensitivity to 87.5 (95 % CI 82.9–90.9), specificity 93.4 (95 % CI 86.8–94.00, negative predictive value (NPV) 90.7, positive predictive value (PPV) 91. Among the 268 patients, who have met ACR criteria, 8.9 % did not fulfill AECG criteria, and among 279 patients, who fulfill AECG criteria, 35 were not evaluated using ACR criteria. The differences are due to the fact that not all patients had keratoconjunctivitis sicca or positive serology and histopathological changes did not occur simultaneously. On the other hand, the patients that met only ACR criteria have been negative for Ro/La antibodies, but RF and ANAs positive. Comparative summary AECG criteria and ACR are shown in Table 2.

Diagnosis of pSS according to AECG criteria requires the assessment of salivary glands or salivary gland scintigraphy or sialography which is a specific, but invasive method of imaging. Unstimulated salivary flow and sialometry are other diagnostic tools to assess the function of

### Table 2: Comparison of AECG and ACR proposed 2012 criteria for pSS [24–26]

| The revised AECG criteria 2002 | ACR proposed criteria 2012 |
|-------------------------------|---------------------------|
| **Subjective symptoms**       |                            |
| 1. Ocular symptoms            |                            |
| Persistent dry eyes for more than 3 months? |
| Recurrent sensation of sand or gravel in the eyes? |
| Using tear substitues more than three times a day? |
| 2. Oral symptoms              |                            |
| Feeling of dry mouth for more than 3 months? |
| Recurrent or persistent swollen salivary gland? |
| Frequent drinking to aid swallowing? |
| **Objective symptoms/examinations** |                            |
| 3. Schirmer’s test without anesthesia (<5 mm/5′) | 1. Positive serum autoantibodies to Ro/SS-A or La/SS-B or both or positive rheumatoid factor and ANA ≥ 1:320 |
| Rose Bengal score >4 in van Bijsterveld’s scoring system | 2. Salivary gland biopsy-focus score ≥1 |
| 4. Histopathology-focus score ≥ 1 | 3. Keratoconjunctivitis sicca with ocular staining score in individuals who is not using eye drops for glaucoma, and has not corneal surgery or cosmetic lid surgery in the last 5 years |
| 5. Sialometry—positive results in one of: |                            |
| Unstimulated salivary flow (<1.5 ml in 15 min |                            |
| Sialography showing the presence of sialectasis without destruction of major ducts |
| Scintigraphy with reduced uptake and concentration and/or excretion of tracer |                            |
| 6. Autoantibodies to Ro/SS-A or La/SS-B or both |                            |

**Exclusions**

- Head and neck radiation
- Hepatitis C infection
- AIDS (acquired immune-deficiency syndrome)
- Pre-existing lymphoma
- Sarcoidosis
- GVH (Graft versus Host) disease
- Use of anticholinergic drugs

- Head and neck radiation
- Hepatitis C infection
- Sarcoidosis
- AIDS (acquired immune-deficiency syndrome)
- GVH disease
- IgG4-related diseases
salivary glands [27]. ACR criteria avoid such methods, limiting the estimation of salivary glands for histopathological examination. However, for clinicians, especially in the case of lack of patient’s consent for salivary gland biopsy, it is important to have a simple, available and noninvasive diagnostic tool for the assessment of salivary glands. It appears that salivary gland ultrasonography (SGUS) meets these assumptions. SGUS is relatively cheap, available and can be use not only for making a diagnosis, but also for controlling a disease progression and conversion into lymphoma. In Theander and Thomas study, 105 patients with pSS and 57 individuals in control group were evaluated by SGUS [28]. The authors used a simplified scoring system to evaluate salivary glands, taking into account the parenchymal echogenicity and inhomogeneity. This scoring system has been proposed by Hocevar [29] from normal echogenicity to next steps of hypechoics lesions. For pSS, the score of 2 or 3 is characteristic. This method of scoring seems to be good for more advanced stages of the disease and can predict, together with laboratory findings, such as positive Ro/La antibodies, the presence of RF and ANAs in more severe disease and a higher risk of lymphoma development. Other researchers evaluate glands size, clearness of borders and vascularity, but it is currently believed that only parenchymal homogeneity is appropriate for the diagnosis of changes in pSS [30]. Other methods, such as MR sialography and computed tomography (CT), may have limited availability and are undoubtedly more expensive. The basis of the diagnosis and a strong argument of inflammation and lymphocytes infiltration of salivary glands is histopathological examination. Standardization of techniques of minor salivary glands biopsy makes it possible to reduce the risk of complications and even increase the value of this study. Varela-Centelles et al. [30] suggested the use of S forceps for better positioning of the lower lip than chalazion forceps and to avoid postoperative complications [31]. In histopathological assessment in pSS focal infiltrates, CD 4+ T cells in early stage and more memory B cells in next stages are observed, further aggregation of lymphocytes, organized lymphoid follicle, ectopic germinal center-like structure (GC) and in some cases lymphoma picture. The degree of organization of infiltration correlates with the severity of the disease, large activity of cytokines in situ and by the presence of autoantibodies [32].

**Sjögren’s syndrome and IgG4-related diseases**

The cases of lacrimal and salivary glands being affected by the disease, that manifests itself with the symptoms of dryness and is accompanied by autoimmune pancreatitis and renal disease (IgG4-related kidney disease), have for years been recognized as a subtype of pSS. Japanese clinicians and scientist first demonstrated that such patients show lymphoplasmocytic infiltrations with IgG4 plasma cells. Additionally, diffuse or local organ enlargement, fibrosis and/or hyperplastic lesions with symptoms of organ damage have been observed in this group. The term “IgG4-related diseases” was finally proposed with classification criteria for IgG4 RD established. IgG4 RD affects mainly elderly man (>60 years old) [33]. The serum level of IgG4 is significantly (>135 mg/dl) increased, and a tissue biopsy reveals infiltrations of IgG4 plasma cells, with IgG4/IgG ratio exceeding 40 %. Patients’ plasma tests seronegative for the presence antinuclear antibodies, including SS-A and SS-B. Hypocomplementemia, high serum levels of polyclonal gammaglobulins with increased level of IgG4 and IgE gammaglobulins may also occur. The histopathological assessment shows plasmocyte and lymphocyte infiltrations and may reveal fibrosis. Establishing the diagnosis of IgG4 RD requires first of all the exclusion of malignancy, for which task histopathological examination is vital. Other diseases, chiefly pSS, but also Castelman’s disease, ANCA-associated vasculitis, idiopathic retroperitoneal fibrosis, sarcoidosis should also be considered as an alternative diagnosis and excluded, using diagnostic criteria for each of the listed diseases. Differences between pSS and IgG4 RD are presented in a Table 3.

**HCV infection and primary Sjögren’s syndrome: a topic for discussion**

Patients infected with HCV may have dryness and cryoglobulinemia, but Ro/SS-A and La/SS-B antibodies are rarely present in the HCV infection. The discussion invokes the potential role of HCV infection in the pathogenesis of pSS. Various studies depict the presence of HCV in 14–19% of patients with pSS. Ramos-Casals et al. [34] conducted a study of 137 patients diagnosed with pSS and co-infected with HCV. In 50 % of this group, cryoglobulins were present, and only 23 % had antibodies to Ro/SS-A and/or La/SS-B, 65 % had ANA antibodies. The authors suggest that chronic HCV infection must be considered as a criterion for exclusion pSS mainly by influencing the development and course of pSS and also proposed the term secondary to HCV Sjögren’s syndrome. Also at work, Madaliński et al. [35] who examined 104 patients diagnosed with pSS searched for markers of infection HCV and in 19.2 %—HCV antibodies were found and 4.8 % patients had genetic material of the virus (HCV-RNA); researchers also suggested distinguish disease as different from pSS.

**Risk of lymphoma and development of other cancers**

Patients with pSS have a significantly higher risk of the development of EBV-associated lymphomas. EBV DNA that was found in saliva and B cells. This presence of
epithelial cells in salivary glands suggest active infection (reactivation of EBV). The development of lymphoma has been described in 4–7 % of patients with pSS [36, 37]. This applies particularly to the development of MALT (mucosa-associated lymphoid tissue) lymphomas and also splenic marginal zone lymphomas (SMZL) as well as nodal marginal zone lymphoma (NMZL) [37]. For years, risk factors associated with the development of lymphomas have been analyzed—taking into account both clinical symptoms and potential biomarkers associated with the development of lymphoproliferation, which are presented in the Table 4 [38, 39].

Table 3 Differences between pSS and IgG4 RD

| Primary Sjogren’s syndrome | IgG4-related diseases |
|---------------------------|----------------------|
| Age between 30 and 50     | Age >60              |
| Mainly women              | Mainly men           |
| Salivary glands enlargement may occur | Organ enlargement (local or diffuse) |
| Symptoms of dryness       | Symptoms of dryness may occur |
| Presence of ANA antibodies | Absence of antinuclear antibodies |
| Presence of anti-SS-A, SS-B antibodies | Absence of anti-SS-A SS-B antibodies |
| Polyclonal hypergammaglobulinemia | Polyclonal hypergammaglobulinemia |
| Normal level of IgG4      | Significantly increased level of IgG4 |
| Mainly CD 4+ T lymphocyte infiltrations | Mainly IgG4-positive plasmocytic infiltrations IgG4/IgG ratio >40 % |
| Treatment: symptomatic and systemic (antimalarics, methotrexate, corticosteroids, cyclosporine A, azathioprine, cyclophosphamide) | Treatment: steroids (no response to steroid treatment indicates incorrect diagnosis) |

Table 4 Clinical manifestations and biomarkers associated with lymphoma development in pSS

| Clinical manifestations | Potential biomarkers |
|-------------------------|----------------------|
| Vasculitis              | Cryoglobulins (mixt cryoglobulinemia) |
| Salivary gland enlargement | Low C4 complement component |
| Salivary gland/parotid swelling | Anti-Ro/SS-A antibodies |
| Lymphadenopathy         | Leukopenia |
| Splenomegaly            | Presence of RF expressing B cells |
| Peripheral neuropathy   | *Flt3-L (Fms-like tyrosine kinase3 ligand) |
| Long duration of pSS    | Higher levels of BAFF/BLyS |
| Histopathology: germinal-like structures in minor salivary glands biopsy |

Systemic administration of drugs to enhance salivating includes agonists muscarinic receptors: pilocarpine and cevimeline [41]. Sometimes N-acetylcysteine may be useful. It is important to treat fungal infections of the oral cavity. Vaginal dryness and dysperunia should be treated with lubricants, propionic acid gels, and sometimes, in the case of menopause, vaginal cream with estrogen.

Systemic treatment is based on antimalaric drugs such as hydroxychloroquine and especially in the case of arthritis methotrexate, corticosteroids, cyclosporine A and azathioprine. In severe cases, with life threat involving organs (renal involvement, neurological, pulmonary involvement and vasculitis) cyclophosphamide, infusions of immunoglobulin, plasmapheresis is used. Patients with renal tubular acidosis should receive sodium and/or potassium.

Potential new therapies In view of the demonstrated key role of B cells—their hyperstimulation and overproduction of autoantibodies in pSS—B cells have become a logical target for therapy. Rituximab (RTX), chimeric antibody against anti-CD20, has proven effective in reducing both the destruction of the salivary glands, reducing inflammatory infiltration in salivary glands, which not only shows the effect on the depletion of B cells, but also on T lymphocytes. Also after infusion of rituximab, significant...
decrease of rheumatoid factor Ig-M in serum was observed [42]. RTX treatment had also an influence on reduction of symptoms of dryness, fatigue and artralgias. Efficacy of B-cell depletion in pSS initiated further search for drugs affecting B cells and BAFF/BLyS, currently accepted in SLE treatment. BELLIS open label phase II study showed benefits of belimumab in patients with pSS, but further studies are needed [43, 44]. Epratuzumab is the humanized anti-CD22 monoclonal antibody that causes moderate B-cell depletion, therefore, can be another toll for pSS treatment. However, previous research showed clinical improvement with no influence on level of autoantibodies [45, 46]. The future of pSS treatment may be an IFN-related therapies and influence on IFN-related pathways. Mesenchymal stem cells (MSCs) transplantation, thanks to low immunogenicity and ability of multipotent stem cells to differentiate into various cells, seems to be promising method in pSS. The effectiveness of such treatment has been demonstrated in SLE and systemic sclerosis [47].

Conclusions

Particular challenge is to make the pSS diagnosis in the early stages of the disease. The presence of typical pSS serological markers such as anti-SS-A and anti-SS-B antibodies at an early stage cannot be confirmed, which is taken into account in the newly proposed ACR classification criteria. Discoveries of novel cellular subsets and cytokines involved in pSS pathogenesis may become a new target for future therapies. Both gene transfer and MSC transplantation can find wider application in the pSS treatment.

Conflict of interest None.

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