The level of antibodies to pneumococcal capsular polysaccharide was determined using the EIA PCP IgG kit (TestLine Clinical Diagnostics s.r.o., Czech Republic) before vaccination, 1, 3, and 12 months after vaccination.

Results: The dynamics of the concentration of antibodies to pneumococcal capsular polysaccharide in patients with SpA is presented in the Table 1.

Table 1. Concentration of pneumococcal antibodies, U/ml, Me [25; 75 percentile]

<table>
<thead>
<tr>
<th>Visit</th>
<th>Concentration of pneumococcal antibodies, U/ml, Me [25; 75 percentile]</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 visit (initial)</td>
<td>160.1 [73.5; 245.7]</td>
</tr>
<tr>
<td>2 visit (after 1 month)</td>
<td>214.5 [103.2; 255.0]</td>
</tr>
<tr>
<td>3 visit (after 3 months)</td>
<td>175.0 [120.1; 260.1]</td>
</tr>
</tbody>
</table>

*p=0.01  **p=0.005

At 1, 3 and 12 months after vaccination, the concentration of antibodies to pneumococcal capsular polysaccharide was significantly higher compared to the baseline values. In 81% of patients, vaccination tolerance was good. Reactions at the injection site (pain, swelling and hyperemia of the skin up to 2 cm in diameter), resolved independently after 1-5 days, were observed in 6 patients. In 2 patients, a severe local reaction was registered in the form of pain in the arm, infiltration and hyperemia of the skin up to 8 and 15 cm in diameter, respectively, accompanied by low-grade fever in one patient for 2 days, and febrile fever in the other for 3 days. In both cases, these symptoms were completely stopped after administration of paracetamol and antihistamines. Exacerbation of SpA and the emergence of new autoimmune disorders were not detected. During the follow-up period, no patients developed lower respiratory tract infections. Patients suffering from frequent sinusitis and otitis reported the absence of these infections after vaccination.

Conclusion: The obtained data indicate satisfactory immunogenicity and good tolerability of PPV-23 in patients with SpA. Further studies are needed to better assess the immunogenicity and safety of vaccine, as well as to study of the influence of anti-rheumatic therapy on the effectiveness of immunization.

Disclosure of Interests: None declared.

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COMORBID INFECTIONS IN PATIENTS WITH SPONDYLOARTHRITIS.

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Background: Data on the frequency and structure of comorbid infections (CI) in spondyloarthritis (SpA) are few and contradictory. Objectives: The aim of the study was to study the frequency and structure of CI in the inpatient population of SpA patients in the course of a one-month retrospective study.

Methods: The study included 208 patients with SpA (121 men, 87 women, mean age 39.1±12.2 years) who were hospitalized at the V.A. Nasonova Research Institute of Rheumatology. Anti-cytoplasmic antibodies were diagnosed in 133 patients, psoriatic arthritis - in 57, spondyloarthritis associated with Crohn's disease - in 1, undifferentiated spondyloarthritis - in 17. The majority of respondents had higher education (60.6%). None of the patients consumed alcohol on a daily basis, 124 patients never smoked. The Charlson comorbidity index, equal to 0, had 98 respondents, 1 - 51, 2 - 27, 3 - 15, 4 - 10, 5 or more - 7. Most patients (n=168) received nonsteroidal anti-inflammatory drugs (NSAIDs), as well as glucocorticoids-GC (average duration of administration 239.5±65.8 months), methotrexate-MT (32.4±46.2), sulfasalazine (21.0±32.1), lefunomide (24.0±46.6), biological drugs - TNF-α inhibitors (21.5±23.3), interferon (9.0±5.2), IL-17 (10.0±9.3). Patients were interviewed by a research doctor with the completion of a unified questionnaire, additional data were obtained from medical documentation.

Results: Leading in the structure of CI in patients with SpA were respiratory tract infections: acute nasopharyngitis (more often 3 times a year), sinusitis, acute bronchitis, pneumonia and herpes-viral infections, in particular herpes zoster. 29.8% of patients reported a more severe course of CI against the background of SpA (12 of them did not receive immunosuppressive drugs). Temporary discontinuation of therapy due to the development of CI occurred in 26.4% of patients. At the same time, in 5 patients treated with GC (including in combination with MT, n=3), the development of furunculosis was the reason for changing the treatment regimen. In one patient, MT therapy was discontinued due to the frequent development of purulent tonsillitis. Exacerbation of SpA after CI was diagnosed in 84 patients (70 of them received immunosuppressive therapy).

Conclusion: The data obtained indicate the important of the problem of CI in SpA. Further studies are needed to determine the large samples of patients in order to find significant risk factors for CI, study their relationship with clinical characteristics and influence on the course of SpA.

Disclosure of Interests: None declared.

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EFFECTIVENESS OF SCREENING IN PATIENTS WITH RHEUMATOID DISEASE ON BIOLOGICAL THERAPY AND RISK OF ACTIVE TUBERCULOSIS

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Background: Treatment with biologic therapy has been associated with a high risk of reactivation of latent tuberculosis (TB). Preventive strategies for tuberculosis remain a crucial step before initiating biologics in rheumatoid disease. Treatment with biologic therapy has been associated with high risk of reactivation of latent tuberculosis (TB). Prevention strategies remain a crucial step before initiating biologics.

Objectives: We aimed to assess the effectiveness of TB screening before the initiation of biologics and the risk of occurrence of active TB among patients with rheumatic diseases on biologic therapies.

The study aimed to access the effectiveness of TB screening recommendations before the initiation of biological therapy and identify the incidence of active TB among these patients.

Methods: We performed a hospital-based retrospective cohort study among rheumatoid disease patients on biological therapy in two centers in Jeddah between January 2005 to December 2019. Medical files were retrospectively reviewed for demographics data, baseline screening for TB, use of prophylaxis, information on DMARDs and biological therapies, and outcomes results were collected.

Results: A total of 365 patients were included over a period of 14 years. Two hundred ninety-two (80%) had Rheumatoid arthritis (RA), 13% psoriatic arthritis (PSA), 9% spondyloarthritis (SPA), 2% SLE, and 4% others. The mean age was 47.5±4 (14.2), 311 (85%) were females with a mean duration of disease 8.45 years (± 6.19). Hundred sixty-nine (49.3%) patients were on steroids. Anti-TNFs were prescribed in 213 (58.4%) patients, Non Anti-TNFs 124 (36.6%) patients, and Jak inhibitors 18 (5%) patients. TB screening was done to all patients except 3 patients (data missing) before commencing biologics. Forty-four (12.1%) patients had latent TB at baseline and all received chemoprophylaxis with isoniazid before starting biologics. Four patients with active TB were identified (one with Behcet’s disease and three with RA). One patient had a reactivation of latent TB and 3 patients developed de novo TB. Three out of four had an infection in the first 6 months of treatment (one on infliximab and two on rituximab) and one case after 1 year of stopping adalimumab. Two cases had pulmonary TB and two others with extrapulmonary TB (pericarditis and brain abscess each). All four patients active TB were treated with standard ant TB medications. Three had complete resolution of their TB and one died.

Conclusion: Baseline screening has been effectively carried out in our cohort as per recommendations. Physician should be vigilant not only for reactivation of latent TB but occurrence of de novo TB in patients on biological therapy.

REFERENCES:
BACKGROUND: The development of biologics for the treatment of systemic rheumatic diseases increased the risk of infections. The management of this complication deserves particular attention since it remains a major cause of morbidity and mortality.

OBJECTIVES: The aim of our study was to determine infection frequency under biological treatment and consequences on the therapeutic management.

METHODS: Patients included in the Biological National Registry (BINAR) from 2016 to 2020. Data related to the disease, biological agents, and infections occurring under biologic disease-modifying antirheumatic drugs (bDMARDs) were collected.

RESULTS: The study included 298 patients with a mean age of 49.2 years (18-79). 175 patients with rheumatoid arthritis and 123 with spondyloarthopathies (Axial Spondyloarthopathies=48, Enteropatic Arthritis=41, Psoriatic Arthritis=34). Anti-Tumor necrosis factor-alpha (Anti-TNF) agents were the most prescribed bDMARDs in 87.9% (n=263) of patients: Infliximab 20.4% (n=61), Etanercept 21.3% (n=69), Adalimumab 24.6% (n=74) and Certolizumab (n=79). No patients were treated with Golimumab. Tocilizumab and Rituximab were prescribed respectively in 10.4% (n=31) and 5% (n=15) of patients. Infections occurred in patients (3.1%) with a total of 13 infectious episodes 12 bacterial and a viral one. The site of infections was: respiratory (38%), urinary (15%), cutaneous (23%). OLT (8%), infective endocarditis (8%), and other (8%). The infectious agent was identified in only 3 patients. The outcomes were favorable in most cases except in one patient where there was a definitive interruption of bDMARDs. The patient was hospitalized for sepsis complicating a cutaneous infection with favorable outcomes under antibiotics within a week. The biological agent with higher risk of infections was Tocilizumab (p = 0.056), unlike Rituximab (p = 0.483) and Anti-TNF (p = 0.082).

Conclusion: Our results confirm that bDMARDs are predisposing to infections, data from BINAR showed that most infections were trivial with no serious outcomes. Therefore, infections should be assessed in patients under bDMARDs for an early therapeutic intervention.

Disclosure of Interests: None declared.

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POS1154
CHARACTERISTICS OF SURGICALLY TREATED TUBERCULOUS AND BRUCELLAR SPONDYLODYSICTIS
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BACKGROUND: Although rare in most developed countries, tuberculous and brucellar spondylodiscis are still frequent in the Mediterranean countries. Management of spondylodiscitis may require a surgical intervention in cases of spinal cord or root compression, bone destructions and deformities, unsuccessful medical treatment or large abscesses.

OBJECTIVES: To study characteristics and outcome of surgically treated tuberculous and brucellar spondylodiscis.

METHODS: Medical records of patients who were admitted into the physical therapy and rehabilitation ward immediately post surgery of the spine were retrospectively studied. The study included records from January 2010 to December 2020.

RESULTS: Twenty patients were diagnosed with tuberculous or brucellar spondylodiscitis for which they underwent surgery. The mean age of diagnosis was 49.7± 11.9 years. The sex ratio was 1:1. The diagnosis was delayed in 60% of the cases due to atypical presentations. Fatigue was reported in 45% of the patients, back pain in 60% of the cases, radicular pain in 15% of the cases, fever in 20% of the patients and abnormal gait in 60% of the cases. Tuberculosis was identified in 75% of cases and brucellar infection in 25% of the cases. On biopsy, elevated inflammatory markers were noted in 60% of the cases. Intradermal reaction for tuberculosis was positive in 9 patients while cultures allowed the diagnosis of tuberculosis in 1 patient. Wright serology test confirmed the diagnosis of brucellar infection in all 5 cases. In the remaing ones, vetalbral biopsy or intraoperative samples allowed the diagnosis of the tuberculous spondylodiscitis. Cervical spine involvement was noted in 5% of cases, lumbar spine in 25% and dorsal spine in 30% of the cases. All patients received antibiotics and spinal immobilization. Surgery was indicated in 78.6% of cases for spinal cord decompression and in 21.4% of the cases for voluminous abscesses. Anterior approach was used in 71% of the cases and posterior approach in 92.9% of the cases. 23.1% of the patients benfited concomitantly with spinal fusion surgery. Post operative success was immediately confirmed by recovery of normal gait in 15.4% of the patients. Complications after surgery were reported in 35.7% of the cases. They consisted in spinal instability in 60% of the cases, abcesses in 20% of the cases and the emergence of secondary articular localizations in 20% of the cases. 28.6% of the patients underwent a second surgery mainly to stabilize the spine. Before surgery, walking was impossible for 65% of the patients and 12 months after surgery only 30% of the patients couldn’t walk (p<0.001). 15.8% of the patients developed later on pressure ulcers and 45% of the patients were prescribed appropriate wheelchairs.

Conclusion: Spinal surgery in tuberculous or brucellar spondylodiscitis may be indicated if medical treatment is insufficient and especially if neurological complications occur. Surgery may be invasive with various complications but sometimes it could be the only option to treat a neurological deficit.

DISCLOSURE:

Disclosure of Interests: None declared.

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POS1155
INFECTIOUS RISK DURING BIOLOGIC THERAPY FOR INFLAMMATORY RHEUMATIC DISEASES: DATA FROM THE TUNISIAN BINAR REGISTRY
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Disclosure of Interests: None declared.

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