**THU0586**

**ESTABLISHING THE KEY COMPONENTS OF A EULAR PORTFOLIO FOR TRAINING IN RHEUMATOLOGY: A EULAR SCHOOL OF RHEUMATOLOGY INITIATIVE**


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**Background:** In clinical training, a portfolio is expected to stimulate learning and encourage critical reflection. Some, but not all, European countries use a portfolio in rheumatology training, and their scope varies widely. A EULAR portfolio for Rheumatology trainees could contribute to improve overall training, raise educational standards, foster the setting of common goals and harmonize rheumatology training across countries.

**Objectives:** Develop key components that should be included in a EULAR portfolio of Rheumatology.

**Methods:** A working group (WG) composed of 9 rheumatologists and 1 educationalist was established. A systematic literature review (SLR) was conducted in November 2018, according to the PPRM structure: Population: trainees; Instrument of interest: portfolio; Measurement of properties of interest: content portfolio. A survey was disseminated among the WG group and WG members of the EMerging EULAR NETwork (EMEUNET), inquiring about the content and structure of existing national portfolios. Portfolio materials of selected countries were reviewed. Last, the WG elected the key components of the portfolio.

**Results:** 132/2034 articles were included in the SLR (12 high/1 moderate risk of bias). Information on direct observation of procedural skills (DOPS) (9/13), personal reflections (8/13), learning goals (5/13) and multisource feedback (5/13) were most often included in the portfolio. Twenty-five respondents filled out the survey (response rate ≈ 50%). Reflective writing (n=7), learning goals (n=4) and feedback (n=4) were considered the most useful components of a portfolio. About half indicated that a portfolio was a bureaucratic burden; 4 respondents mentioned lack of feedback by supervisors as a barrier. Portfolio materials of 7 European countries were reviewed. Several portfolios (Germany, Italy, France and Spain) were logbooks, i.e. a record of clinical activities. Other portfolios (UK, Denmark, The Netherlands) also included information on workplace-based assessments, learning goals, and personal reflections. The proposed key components of the portfolio are included in Table 1.

**Table 1. Key components of the EULAR portfolio of Rheumatology.**

<table>
<thead>
<tr>
<th>Key component</th>
<th>Content</th>
</tr>
</thead>
<tbody>
<tr>
<td>Curriculum vitae</td>
<td>Personal record of achievements, experiences, knowledge, and skills</td>
</tr>
<tr>
<td>Personal Development Plan</td>
<td>Learning goals and action plan</td>
</tr>
<tr>
<td>Clinical work</td>
<td>Information on managing patients (e.g. rheumatoid arthritis)</td>
</tr>
<tr>
<td>Professional behaviour</td>
<td>Multisource feedback</td>
</tr>
<tr>
<td>Education</td>
<td>Continuing professional development, list of formal and non-formal learning activities</td>
</tr>
<tr>
<td>Research</td>
<td>List of abstracts, published articles</td>
</tr>
</tbody>
</table>

**Conclusion:** This initiative resulted in the establishment of a list of key components to be included in a EULAR portfolio of Rheumatology. Assessment forms for each key portfolio component are currently being developed. Portfolio implementation, particularly in countries that do not use it yet, may contribute significantly to promote a higher standard of patient care across Europe.

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**Educational cases**

**THU0587**

**TB OR NOT TB? THIS IS THE QUESTION. CASE REPORT OF AN EXTRAPULMONARY TUBERCULAR ARTHRITIS.**

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**Background:** Tuberculous (TB) arthritis consists of 1-3% of all TB cases, whereas TB tenosynovitis & bursitis account for 1%. Primarily it involves large joints but occasionally smaller non-weight-bearing joints. Diagnosis is usually delayed due to lack of awareness, radiographic findings & constitutional or pulmonary involvement.

**Objectives:** We aim to increase rheumatologists awareness to detect possible TB etiology & arthritis & tenosynovitis.

**Methods:** Our case is a 32 years old male complaining of polyarthritis of wrists, MCPs, ankle joints 4 months prior to presentation. Patient was referred as diagnosed rheumatoid patient resistant to treatment based on clinical presentation & laboratory investigation. His lab. was as follows: ESR 76 mm/hr, CRP 56.6 mg/L. RF 18.18 IU/ml, Serum creat 0.8 mg/dl, SGOT 20 SGPT 22, FBS 94, Uric acid 5.4, Hepatitis & HIV negative. CBC showing Hb 14.1 g/dl, TLC 7030/mm³ & platelets 289000/ml. There was no history of genitourinary, gastrointestinal manifestations, oral/genital ulcers, ophthalmological, mucocutaneous, cardiac, pulmonary, hepatic nor renal manifestations. The treatment at time of presentation was Methotrexate 25mg/week IM injection, Leflunomide 20mg/d & low dose steroids, prednisolone 5mg/d. Patient was referred to our department to assess activity, perform musculoskeletal ultrasound on the various involved joints. Hence, expected by referring physician to shift from DMARDs to biologic treatment.

**Results:** MSUS study following eular guidelines showed active synovitis in both radiocarpal & midcarpal joints bilaterally grade II by doppler signal (figure 1). Other active synovitis in multiple MCPs as well as tenosynovitis of Peroneus longus & brevis bilaterally was detected (figure 1). The swelling around the ankle was alarming though the other swollen joints seemed to be consistent with a case of RA in activity. This swelling revealed a well-defined hypoechoic heterogeneous cystic fluid collection with posterior through-transmission (figure 2) & hypechoic hyperemic wall on PD imaging opposite medial malleolus of right fibula. The laboratory investigations prior to shifting patient had to included TB tests, tuberculin test and PCR following the positive result that we found in the skin test. Aspiration was performed from the cystic swelling and sent for clinical pathology analysis. Thick yellowish fluid aspirate on cytology revealed moderately cellular mainly of PMN cells, neutrophils, nuclear debris in proteinaceous background no atypical or malignant cells were found. As regards bacteriology no pus with no growth (both aerobic & anaerobic). These results warranted us to perform a culture for atypical bacteria and revealed growth of mycobacterium tuberculosis. AntiTB therapy was started for 9 months in the form of 2 months of isoniazid (INH) and rifampicin (RIF), pyrazinamide (PZA) and ethambolot (EMB) followed by 7 months of INH and RIF. Excision of the synovial cyst was done on the spot.

**Figure 1.**
Background: Takayasu Arteritis is a chronic, large vessel arteritis that commonly involves the aorta and its major branches, mostly the ascending/descending aorta, subclavian arteries, and carotids [1]. Herein, we report a case of a 23-year-old medically free Indian male who presented to our hospital in acute distress complaining of cough, hemoptysis and shortness of breath for one week as well as intermittent fever and fatigue for five months. He presented with a BP of 140/100 mmHg as well was both systolic and early diastolic murmurs in the mitral and aortic areas, respectively. He also had a paraumbilical bru. Unilateral clubbing in patients with TA occurs as a result of subclavian artery stenosis that leads to tissue ischemia and hypoxia [2-4]. In turn, the bone marrow release megakaryocytes, which enter the systemic circulation when an A-V shunt exists [5]. Platelet-derived growth factor (PDGF) play a central role in the pathogenesis of unilateral clubbing, patients' clinical condition including presence of clubbing improved after initiation of immunosuppressive therapy.

References:

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TAKAYASU'S ARTERITIS PRESENTING WITH UNILATERAL DIGITAL CLUBBING IN A 23 YEAR-OLD MALE

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Background: Takayasu Arteritis is a chronic, large vessel arteritis that commonly involves the aorta and its major branches, mostly the ascending/descending aorta, subclavian arteries, and carotids [1]. Herein, we report a case of a 23-year-old medically free Indian male who presented to our hospital in acute distress complaining of cough, hemoptysis and shortness of breath for one week as well as intermittent fever and fatigue for five months. He presented with a BP of 140/100 mmHg as well was both systolic and early diastolic murmurs in the mitral and aortic areas, respectively. He also had a paraumbilical bru. Unilateral clubbing in patients with TA occurs as a result of subclavian artery stenosis that leads to tissue ischemia and hypoxia [2-4]. In turn, the bone marrow release megakaryocytes, which enter the systemic circulation when an A-V shunt exists [5]. Platelet-derived growth factor (PDGF) play a central role in the pathogenesis of unilateral clubbing, patients' clinical condition including presence of clubbing improved after initiation of immunosuppressive therapy.

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