The European network for care of children with Paediatric Rheumatic Diseases: care across borders.

P. Dolezalova¹, J. Anton², T. Avcin³, M. Beresford⁴, P. A. Brogan⁵, T. Constantin⁶, Y. Egert⁷, I. Foeldvari⁸, H.E. Foster⁹, V. Hentgen¹⁰, I. Kone-Paut¹¹, J. Kummerle-Deschner¹², P. Lahdenne¹³, B. Magnusson¹⁴, A. Martini¹⁵, L. McCann⁴, K. Minden¹⁶, S. Ozen¹⁷, C. Schoemaker¹⁸, P. Quartier¹⁹, A. Ravelli¹⁵, I. Rumba²⁰, N. Ruperto¹⁵, S. Vastert¹⁸, C. Wouters²¹, F. Zulian²², and N.M. Wulffraat¹⁸ for the SHARE consortium

¹: Charles University in Prague and General University Hospital, Department of Paediatrics and Adolescent Medicine, Praha, Czech Republic
²: Hospital Sant Joan de Déu, University of Barcelona, Department of Pediatric Rheumatology, Barcelona, Spain
³: University Children’s Hospital, University Medical Centre Ljubljana, Department of Allergology, Rheumatology and Clinical Immunology, Ljubljana, Slovenia
⁴: Alder Hey Children's NHS Foundation Trust, Paediatric Rheumatology, Liverpool, United Kingdom
⁵: University College London Great Ormond St Institute of Child Health. London, United Kingdom
⁶: Semmelweis University, Unit of Paediatric Rheumatology, Dept of Pediatrics, Budapest, Hungary
⁷: European network childhood arthritis patient organisation, Israel
⁸: Hamburg Centre for Pediatric and Adolescent Rheumatology, Hamburg, Germany
⁹: Newcastle Hospitals NHS Foundation Trust, Great North Children’s Hospital, Newcastle Upon Tyne, United Kingdom
¹⁰: A. Mignot University Hospital Versailles – CEREMAI, Pediatric Rheumatology, Le Chesnay (Paris), France
¹¹: Le Kremlin-Bicêtre University Hospital, Paris-Sud University, Paris, France
¹²: University Children’s Hospital Tuebingen, Centre for Rheumatology / Autoimmune diseases, Tuebingen, Germany
13: Children's Hospital, Helsinki University Central Hospital, Pediatric Rheumatology, Helsinki, Finland
14: Karolinska University Hospital, Pediatric Rheumatology Unit, Karolinska Hospital, Stockholm, Sweden
15: Giannini Gaslini Institute, Department of Pediatrics, Genoa, Italy
16: Charite University Hospital Berlin, Dpt. of Dermatology and Allergy, Berlin, Germany
17: Hacettepe University Children's Hospital, Department of Pediatric Nephrology and Rheumatology, Ankara, Turkey
18: Wilhelmina Kinderziekenhuis, Department of Pediatric Immunology and Rheumatology, University medical center Utrecht, Utrecht, Netherlands
19: Paris-Descartes University, Necker Children’s Hospital, Paris, France
20: University of Latvia, Faculty of Medicine, Riga, Latvia
21: University Hospital Gasthuisberg, Department of Pediatrics, Division of Pediatric Rheumatology, Leuven, Belgium
22: Clinica Pediatrica I, Unità di Reumatologia Pediatrica, Padova, Italy
Abstract

Objectives: To provide an overview of the paediatric rheumatology (PR) care in Europe to inform future specialist service provision.

Methods: An online survey was developed and presented to national coordinating centres of the Pediatric Rheumatology International Trials Organisation (PRINTO) representing a single EU member state (35 centres; Country survey); and to 288 individual PR centres (Centre and Disease Surveys) as a part of the EU project SHARE. The survey contained country and centre specific components covering organisation of PR care, composition of teams, education, healthcare and research facilities, and assessment of needs. The national coordinating centres completed both centre as well as country questions.

Results: Response rates were 83% for Country and 57% for Centre surveys. Data from both surveys were itemised to organisational, quality of care and educational aspects. Across the EU, only one paediatric rheumatologist is available per million population, located in one of the 288 centres with specialised PR care. In all EU member states, there is overall good access to specialist care and to approved medications, although off-label medication availability is worse in Eastern European countries. Full financial coverage is provided for most prescribed medications. PR education is widely available for physicians but is insufficient for allied health professionals. Participation in clinical trials is generally high. Among important gaps identified, lack of widely accepted clinical guidelines/recommendations; and insufficient adolescent transition management planning were highlighted.

Conclusions: This study provides a comprehensive analysis of specialist PR service provision across Europe. Seen from the perspective of health care providers, there are no major differences between EU member states. Rarity, chronicity and complexity of diseases form a major challenge to paediatric rheumatology care. Therefore, strengthening subspecialty networks (Paediatric Rheumatology European Society, PReS, PRINTO, and SHARE) and the recently created European Reference Networks (ERN) will facilitate provision an dissemination of standards of care and treatment recommendations to further improve patient-centred healthcare across Europe.

Introduction

The care and research for rare and complex disorders need strong collaborative networks, widely
applicable treatment recommendations and standardized follow-up measures. We present an example of such a collaboration in the area of Paediatric Rheumatic Diseases (PRD), a heterogeneous group of rare inflammatory conditions affecting children. The EU project “Single Hub and Access point for paediatric Rheumatology in Europe” (SHARE, grant number No. 20111202) was set up to provide an overview of the current organisation of paediatric rheumatology (PR) with the aim to harmonise the care for children with rheumatic diseases in Europe (1). By doing so it promotes one of the main EU goals: to deliver equal health care across all EU member states. In this report we present results of a detailed survey from 35 European countries (online appendix), to provide an overview of paediatric rheumatology (PR) care in Europe to inform future specialist service provision, and identify currently unmet needs or deficiencies in specialist PR service provision.

Methods
The SHARE consortium included PR representatives from 15 European countries and 2 patient organisations (CS and YE). Participants had been actively involved in educational and scientific activities of the Paediatric Rheumatology European Society (PReS) and Pediatric Rheumatology International Trials Organisation (PRINTO). (2-7). The survey methodology and strategy were agreed at an initial consensus meeting. Three key components were identified and formulated into surveys probing Country, Centre and Disease-specific issues (the latter spanning all the main groups of PRD identified by consensus). The PRINTO provided a platform for online data capture. Access to the country survey was provided to one national PRINTO co-ordinator per country (n=35), including Israel and Turkey. The latter two were included because of their long-term participation in PReS and PRINTO activities, with significant historic contribution to PR provision of care and research. The centre and disease surveys were presented to all PRINTO centres (n=288) from these 35 countries.

Characteristics of surveys
The country survey included 10 questions on the healthcare system, organisation of PR care, and access to care; 4 questions on evidence based and qualified care; and 10 questions on education and employment issues (Supplemental Table 1). The Centre survey covered assessment of needs (21 questions), evidence based and qualified care (24 questions), preventive care and adequate
treatment of co-morbidities (6 questions), transitional care and transferral to adult care (6 questions), and education and employment issues (3 questions; Supplemental Table 2). In 4 disease-specific surveys, 6 questions covered areas of the characteristics and numbers of patients followed in the centre, timing of clinic appointments, use of standardised disease measures, availability of drugs, and patient information strategies.

Data analysis
Descriptive data were reported as medians (1st-3rd quartile) for continuous variables; or as absolute numbers and percentages for categorical variables. Percentages were presented for both Eastern and Western European respondents only if they differed by more than 10%.

Results

From 35 countries approached, 29 representatives replied: 17/18 from Western European Countries (WEC), 12/17 from Eastern European Countries (EEC), with total response rate 83%. For this analysis, Turkey was grouped with EEC, and Israel with WEC. The Centre survey was returned by 165/288 respondents from 33 countries (125 WEC, 40 EEC; response rate 57%), ranging between 1 and 21 per country (Figure 1). Upon analysis, results of both surveys were itemised to organisational, quality of care and educational aspects.

Organisation
General characteristics
In 25/29 (86%) countries full coverage is provided for most prescribed medications (94% WEC, 75% EEC). Limited coverage of prescribed medication was reported by two EEC only. Inpatient and outpatient medical care is fully covered in 62.5% (70.6% WEC, 54.5% EEC) and 42.9% of countries, respectively.

Organisation and access to care
The median number of paediatric rheumatologists per 1 million population was 1.1 (IQR 0.6-1.8 for WEC, 0.5-1.3 for EEC). PR professional organisations were identified in 96.4% of countries, and combined physicians and allied health professionals (AHP) in 43% of countries. In 82% of
countries such societies organize regular meetings, provide educational materials or provide a platform for communication. In over 60% of countries, more than half of patients were managed by paediatric rheumatologists in tertiary centres (53% WEC, 73% EEC). In 64% and 46% of countries a significant minority (< 25%) of patients were managed by other specialists.

National referral guidelines/clinical recommendations were available in 54% of countries (59% WEC, 45% EEC) as reported by national representatives. Nearly 50% of responding centres were based in paediatric hospitals (Table 1). From those, 66% were tertiary centres in university hospitals. More than half of these units acted as paediatric rheumatology national referral centres.

The catchment population for PR services varied significantly according to the size and population of the country and type of centre, ranging from 300 000 in Norway to 80 million inhabitants in Germany. Nearly 20% of patients travelled more than 150 km to their hospital to obtain access to PR care.

The staff, bed and patient numbers at individual centres are shown in Tables 2 and 3. The average waiting time for new referrals was below 4 weeks in 47.7% of centres, and was 4-8 weeks in 21.9%. A new referral was allocated 30-45 minutes clinical time in 54% centres. Follow-up visits were 20-30 minutes in 59% of centres. Combined clinics with other paediatric specialists were available in 78% of centres.

**Quality of care**

**Availability of treatments**

National guidelines/recommendations for PRD were available in 54% of countries (59% WEC, 45% EEC). Licensed drugs for approved indications were readily available (within weeks) in 96%, without limitation in 43% (53% WEC, 27% EEC), and limited by budget in 29% of countries (24% WEC, 36% EEC).

Disease modifying anti-rheumatic drugs (DMARDs) including subcutaneous methotrexate were readily available in 97% of centres (100% WEC, 89% EEC). Licensed biologics could be prescribed by any qualified paediatric rheumatologist in 82% of countries (88% WEC, 73% EEC).
Limitation of prescription at designated centres was reported by 43% of countries (35% WEC, 55% EEC). Biologics for off-label indications were generally available in 79% (94% WEC, 58% EEC) and could be prescribed without limitations in 21% (35% WEC, 0% EEC), and with budget limitations in 18% of countries. From 31% of all individual centres where off-label biologics were not readily available (21% WEC, 63% EEC), over 80% reported regulatory issues as the main barrier to prescription. The use of biologics was recorded in a registry in 96% of countries: 29% national, 21% international, and 29% both types.

*Treatment monitoring, prevention*

Standardised drug monitoring was provided to patients on biologics as well as DMARDs in over 90% of all centres. For patients on long-term corticosteroids, standardised drug monitoring included regular blood pressure measurement, ophthalmology assessment, and bone densitometry in 90% of centres (88% WEC, 97% EEC). Compliance to medication was routinely assessed in 95% of centres. Treatment outcomes of patient cohorts were regularly reviewed in 72% of centres (68% WEC, 86% EEC).

*Transitional care and transferral to adult care*

Specific issues such as alcohol, drugs, sexual health and transition to adult rheumatology were addressed with adolescents in 88% of centres. Preparation for the transition to adult services started after the age of 16 years in 69% of centres (65% WEC, 79% EEC). Management plans in transfer groups were standardized between paediatric and adult care in 50% of centres only (53% WEC, 40% EEC). Potential difficulties regarding transition to adult care were addressed in the teams of 69% of centres (75% WEC, 49% EEC). Dedicated/transition clinics for adolescent patients were provided by 54% of centres (58% WEC, 37% EEC).

*Clinical research*

In over 90% of centres patients had an opportunity to participate in clinical studies. Laboratory-based scientific research was performed in 41%; and investigator-initiated clinical research in 70% of centres (76% WEC, 51% EEC). Sponsored drug trials were reported in 54% of centres. The median number of patients enrolled over the past 5 years per centre were: 100 (IQR 30-200) in
cohort studies; and 10 (IQR 5-20) in drug trials, similar for WEC and EEC centres. At least part of the team was trained in Good Clinical Practice in 79% of centres.

**Education**

Paediatric rheumatology was officially recognised as a subspecialty in 48.5% of countries. A syllabus describing the paediatric rheumatic disorders was endorsed by the European Academy of Pediatrics (EAP) and PRES. (8) A defined training scheme for paediatric rheumatology was available in 46% of countries. Sufficient resources were available for the specialist training in 68% of countries (59% WEC, 82% EEC). Insufficient English language proficiency was identified as an educational burden mainly for AHPs in 5 countries (3 EEC).

Medical schools in 60% of countries provided under-graduate teaching in PR. A training programme in PR for primary care physicians/general paediatricians was available in 14% of countries, while a specific training for nurse practitioners in PR was missing in 96%.

*Continuing Medical Education (CME)*

Over 80% of centres reported more than 20 CME hours per year being devoted to PR by their team members. A national medical registration of specific post-graduate/continuing education for physicians existed in 54% of countries (41% WEC, 73% EEC). A similar system for AHP was available only in 18% of countries. A need for more paediatric rheumatologists was reported by 72%, while a process for manpower planning was available in only 18% of countries (12% WEC, 27% EEC). The majority of countries (93%) reported an improvement in the situation for PR at the time of this survey as compared to 10 years ago.

**Disease-specific issues**

Numbers of patients with individual PRD seen by individual centres are shown in Table 3. The PRINTO website for families (www.pediatric-rheumatology.printo.it) was the most common patient information resource used by more than 60% of centres for all groups of PRD. Patient-reported outcome tools were available for the assessment of physical functioning (Childhood Health Assessment Questionnaire, CHAQ), quality of life and school attendance. These were
widely used for all PRD in 37-66% of all centres. Where available, disease-specific tools were applied by a variable proportion of centres, with JIA assessments being used most widely. There were no significant differences between WEC and EEC centres in the use of any assessment tools.

**Discussion**

This is the first pan-European report on the current state-of-art of PR care published to date. In general, PR differs from many other paediatric subspecialties and from adult rheumatology because virtually all PRD are rare diseases. Therefore, direct comparison with other medical specialties, generally caring for both common and rare diseases, is challenging. A comprehensive report from 2011 provided an overview of PR workforce requirements from a US perspective, and identified 4 main educational and economic barriers to service development (9-11). The Author has synthesized published data into specific policy goals offering solutions to each of the barriers. The main attention has been paid to more general aspects of education, organisation, coverage and availability of care. The more recent review describes various approaches of the paediatric rheumatology learning network established in 2011 to improve health outcomes of PRD in USA and Canada(12).

The European PR network has been expanding for more than 20 years. This study aimed to investigate the current organisation and delivery of specialist PR care across Europe and allied countries. Response rates were generally high. A remarkable general finding was the homogeneity of outcomes across replies from 29 countries, despite their significant economic and cultural differences. The PRINTO National Coordinating Centres from the bigger EU member states all replied, as did no less than 165 other paediatric health care providers. That said, the survey was available in English only which may have limited response rate in some countries.

Children with rheumatic conditions should be treated by paediatric rheumatologists. This is in keeping with the replies of 155 centres who indicated that only a minority of children with PRD were treated by other specialists. In the USA almost 25% of surveyed adult rheumatologists care for children with JIA (13). In Europe most of the paediatric rheumatology care is delivered in tertiary care academic hospitals, especially in EEC (8).
Availability of national referral guidelines and clinical recommendations for PRD was surprisingly low both in WEC and EEC. This important unmet need is currently being addressed by another work package of the SHARE project, with several sets of recommendations and clinical guidelines now published for various PRD (14-16), as several others pending publication. In 15 countries there were differing replies from individual centres within one country to the question on the existence of an "official system of national referral centres". This may reflect a discrepancy between the clinical practice and official healthcare system rules, or a lack of communication within these country. We also detected limited attention to active patient participation in PR care and research: input from patients and caregivers was collected only in 66% of centres. Management plans for patient transition from paediatric to adult specialist care were lacking in approximately half of the centres. This is a clear area for improvement that should be endorsed by PReS, and patient advocacy groups in order to better foster patient involvement.

Access to specialist PR clinical services and to standard medications in Europe was good, with nearly 70% of new patients being seen within 8 weeks of referral. Most prescribed drugs were fully reimbursed in the majority of countries. The availability of expensive biologics for approved indications was surprisingly high, both in the West and the East. Nevertheless, access to their off-label use was restricted, especially in EEC. Unsurprisingly, over 80% of centres who reported this problem indicated regulatory issues as the main barrier.

The use of various disease outcome parameters was reported by 37-66% of centres. Standardised assessment of outcomes is considered one of the main quality improvement methods and has been incorporated among overarching recommendations for the management of PRD (12, 17). Disease and treatment registries also belong to clinical research as well as care improvement instruments (18). Accordingly, the majority of patients treated with DMARDs and/or biologics are followed in pharmacovigilance registries. An overall high level of clinical study participation was also encouraging, reflecting the engagement of paediatric rheumatologists and patients alike with rare PRD research.

Although resources for continuing education are higher in WEC, centre replies indicated that national registration of specific post-graduate education exists in only 41% of WEC, compared to
73% EEC. This finding was relatively independent of the subspecialty recognition status reported by 50% of both WEC and EEC. To this end, a European Syllabus for Training in Paediatric Rheumatology has been prepared by the PReS Education and Training Committee, and approved by the European Academy of Paediatrics. Also, members of the recently established European Reference Networks (ERN) must document extensive training and registration for their staff (https://ec.europa.eu/health/rare_diseases/european_reference_networks_en). Lack of specialist postgraduate education for allied health professionals is another important unmet need revealed by this study. More detailed analyses of education in PR is a subject of a separate report.

In agreement with reports from other continents (19-21) insufficient numbers of paediatric rheumatologists have been reported by more than 70% of both EEC and WEC, suggesting that the median number of 1.1 PR specialist physicians per 1 million inhabitants is insufficient. Differences regarding the definition of "paediatric rheumatologist" may account for this discrepancy. For instance, paediatric rheumatologists from smaller countries and centres might devote only 50% of their clinical hours to this subspecialty service, with the remainder devoted to general paediatric services. The number of specialists thus might not correspond to the number of full-time paediatric rheumatology posts. The suboptimal organisation of care is also reflected by the lack of systematic manpower planning in the subspecialty healthcare in about 80% of countries.

Our results are limited by the fact that we were not able to verify survey replies by cross-checking individual hospital returns with national healthcare figures, and by the aforementioned limitation that the surveys were only available in English which may have limited responses from some centres. There may be inherent biases, as in all survey studies. For instance, respondents used to higher standards of care could have evaluated more negatively their current resources, whereas responders used to lower standards may have evaluated their services more positively, thus narrowing any potential gaps between East and West. However we do not have any hard evidence to support this notion. Another potential source of bias relates to geographical distribution of centres, which was denser in the West (125 centres) compared to the East (40 centres). This could have biased the results due to inequity of the number of larger high-quality centres in these two geographic locations. Despite that, a strength of our study is the high number of replies from almost all EU countries, and the fact that at the time of final analysis there were no missing answers from
any of the surveys from the centres that replied. Thus to the best of our ability we believe that the data provide an accurate picture of PR services in the EU and allied countries surveyed.

In conclusion, our survey provides the first comprehensive analysis of current PR specialist services in Europe. We have identified its strengths as well as weaknesses at multiple healthcare levels. Rarity, chronicity and complexity of PRD pose particular challenges for paediatric rheumatology care world-wide. Therefore, understanding what the current state of service provision across the EU is essential in order to further develop optimal care for PRD. Importantly, strengthening existing networks to allow better knowledge transfer regarding best practice, development and dissemination of evidence-based clinical guidelines to benchmark care, and ongoing efforts to facilitate international collaborative research for PRD are key to further improve specialist service provision in paediatric rheumatology across the EU and beyond. Of note, the recent formation of European Reference Networks (ERN) on rare diseases within the EU project has been an important starting point for healthcare improvement in this area.

Acknowledgement:
SHARE, grant number No. 2011 1202
all the country representatives who replied (national coordinators) Printo to provide the list please
Table 1: Type and size of hospitals/units where paediatric rheumatology centres are based

<table>
<thead>
<tr>
<th>Centre origin</th>
<th>WEC</th>
<th>EEC</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>125</td>
<td>40</td>
<td>165</td>
</tr>
<tr>
<td>Children's hospital</td>
<td>54 (43.2%)</td>
<td>23 (57.5%)</td>
<td>77 (46.7%)</td>
</tr>
<tr>
<td>General hospital</td>
<td>35 (28.0%)</td>
<td>4 (10.0%)</td>
<td>39 (23.6%)</td>
</tr>
<tr>
<td>Both</td>
<td>29 (23.2%)</td>
<td>9 (22.5%)</td>
<td>38 (23.0%)</td>
</tr>
<tr>
<td>Other (specify)</td>
<td>7 (5.6%)</td>
<td>4 (10.0%)</td>
<td>11 (6.7%)</td>
</tr>
<tr>
<td>Number of all paediatric inpatient beds</td>
<td>110 (40 - 175)</td>
<td>112 (60 - 233)</td>
<td>111 (40 - 195)</td>
</tr>
<tr>
<td>Number of paediatric rheumatology inpatient beds</td>
<td>3 (1 - 5)</td>
<td>8 (5 - 13)</td>
<td>4 (2 - 8)</td>
</tr>
<tr>
<td>Number of all paediatric outpatients per week</td>
<td>350 (150 - 775)</td>
<td>400 (135 - 1033)</td>
<td>400 (150.0 - 889)</td>
</tr>
<tr>
<td>Number of paediatric rheumatology outpatients per week</td>
<td>30 (15 - 50)</td>
<td>48 (25 - 85)</td>
<td>30 (20 - 50)</td>
</tr>
</tbody>
</table>

Legend: WEC Western European Countries  EEC Eastern European Countries

Table 2: Paediatric Rheumatology Centre characteristics

<table>
<thead>
<tr>
<th></th>
<th>Median Number</th>
<th>1st-3rd quartile</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paediatric Rheumatologists</td>
<td>3</td>
<td>2 - 4</td>
</tr>
<tr>
<td>Fellows/Trainees</td>
<td>1</td>
<td>1 - 2</td>
</tr>
<tr>
<td>Nurses</td>
<td>2</td>
<td>1 - 4</td>
</tr>
<tr>
<td>Other AHP</td>
<td>2</td>
<td>1 - 3</td>
</tr>
<tr>
<td>Paediatric Beds total</td>
<td>111</td>
<td>40 - 211</td>
</tr>
<tr>
<td>Paediatric Rheumatology beds</td>
<td>4</td>
<td>2 - 8</td>
</tr>
<tr>
<td>Paediatric Outpatients total/week</td>
<td>425</td>
<td>150 - 897</td>
</tr>
<tr>
<td>Paediatric Rheumatology Outpatients/week</td>
<td>30</td>
<td>20 - 52</td>
</tr>
</tbody>
</table>

Legend: AHP Allied Health Professionals
Table 3: Patient population characteristics in individual centres: Median numbers of reviewed and new patients in one year per centre

<table>
<thead>
<tr>
<th></th>
<th>125 WES</th>
<th>40 EEC</th>
<th>165 Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total / New</td>
<td>Total / new</td>
<td>Total / new</td>
</tr>
<tr>
<td>JIA</td>
<td>165 / 20</td>
<td>181 / 30</td>
<td>168 / 25</td>
</tr>
<tr>
<td>SLE</td>
<td>8 / 2</td>
<td>10 / 3</td>
<td>9 / 2</td>
</tr>
<tr>
<td>APS</td>
<td>1 / 0</td>
<td>2 / 1</td>
<td>1 / 0</td>
</tr>
<tr>
<td>VAS</td>
<td>4 / 1</td>
<td>6 / 2</td>
<td>5 / 1</td>
</tr>
<tr>
<td>JIIM</td>
<td>5 / 1</td>
<td>5 / 2</td>
<td>5 / 1</td>
</tr>
<tr>
<td>ISS</td>
<td>0 / 0</td>
<td>1 / 1</td>
<td>1 / 0</td>
</tr>
<tr>
<td>AID</td>
<td>15 / 3</td>
<td>12 / 5</td>
<td>14 / 4</td>
</tr>
</tbody>
</table>

Legend: WEC Western European Countries  EEC Eastern European Countries JIA Juvenile Idiopathic Arthritis  JIIM Juvenile Idiopathic Inflammatory Myopathies  JS Juvenile Scleroderma  SLE Systemic Lupus Erythematosus  APS Antiphospholipid syndrome  VAS Vasculitides  AID Autoinflammatory Diseases
Figure 1: Overview of countries participating in the survey. The numbers indicate the ratio of the total number of sites in the country, and the number of sites that actually participated

Supplemental material

Table 1: Country survey

Table 2: Centre survey

References