The SSc community – overview of registries and educational opportunities in SSc

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The role of registries in understanding the disease & education opportunities

1. The disease course
2. The Mortality
3. Disease features & Organ manifestations
4. The Gender enigma
5. The challenge of prediction
6. The problem of measuring the outcome
7. Education & opportunities
8. Conclusions
# Worldwide estimates of economic burden of SSc

<table>
<thead>
<tr>
<th>Country (patient number)</th>
<th>Annual direct cost per patient (healthcare, ambulatory, pharmaceutical)</th>
<th>Total annual indirect cost per patient (productivity loss)</th>
<th>Total annual cost per patient (US $) per year</th>
</tr>
</thead>
<tbody>
<tr>
<td>Australia</td>
<td>AUD$11,622 (USD$8,554)</td>
<td>$8,024 (USD$6,178)</td>
<td>$15,127</td>
</tr>
<tr>
<td>Canada</td>
<td>$3,879</td>
<td>$10,341</td>
<td>$14,208</td>
</tr>
<tr>
<td>USA</td>
<td>$17,365</td>
<td>not measured</td>
<td>$17,365</td>
</tr>
<tr>
<td>Spain</td>
<td>$9,372</td>
<td>$14,571</td>
<td>$23,948</td>
</tr>
</tbody>
</table>

**Estimated cost of SSc in Australia per year**

<table>
<thead>
<tr>
<th>Direct</th>
<th>Indirect</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>64 Million</td>
<td>44 Million</td>
<td>108 Million</td>
</tr>
</tbody>
</table>

The course of the disease

Early Intermediate Late

LIMITED SSc

PH, malabsorption

DIFFUSE SSc

Disease duration (years)

Skin thickness

Lung, heart, GI, kidney

Modified from Medsger T & Steen V, Systemic Sclerosis, 1995, p 51, Williams & Wilkins.
What have multicentre registries across the world taught us about the disease features of systemic sclerosis?

Largest meta-cohort analysis in SSc to date, reporting on frequency of disease manifestations and mortality in 17,838 SSc patients from ASIG, CSRG, EUSTAR ad SCORE cohorts
Kaplan-Meier analysis of survival following disease onset in each cohort.
Overall Survival

### Survival

<table>
<thead>
<tr>
<th></th>
<th>1-year</th>
<th>3-year</th>
<th>5-year</th>
<th>8-year</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Inception</strong></td>
<td>99.0%</td>
<td>94.8%</td>
<td>88.9%</td>
<td><strong>81.3%</strong></td>
</tr>
<tr>
<td><strong>Non-Inception</strong></td>
<td>100%</td>
<td>100%</td>
<td>99.8%</td>
<td><strong>98.8%</strong></td>
</tr>
<tr>
<td><strong>Prevalent</strong></td>
<td>99.5%</td>
<td>98.0%</td>
<td>96.7%</td>
<td>94.6%</td>
</tr>
</tbody>
</table>

**23 Years of Life Lost (YLL) in women and 26 YLL in men**
Causes of deaths-SSc related (62.1% of deaths in inception and 55% deaths in prevalent cohort)

<table>
<thead>
<tr>
<th>Organ system</th>
<th>Inception</th>
<th>Prevalent</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Heart and Lung</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PAH</td>
<td>22 (25.3)</td>
<td>88 (36.1)</td>
</tr>
<tr>
<td>ILD</td>
<td>18 (20.7)</td>
<td>53 (21.7)</td>
</tr>
<tr>
<td>PAH and ILD</td>
<td>8 (9.2)</td>
<td>32 (13.1)</td>
</tr>
<tr>
<td><strong>Gut involvement</strong></td>
<td>12 (13.8)</td>
<td>24 (9.8)</td>
</tr>
<tr>
<td><strong>Sepsis</strong></td>
<td>1 (1.1)</td>
<td>4 (1.6)</td>
</tr>
<tr>
<td><strong>Myocardial involvement</strong></td>
<td>13 (14.9)</td>
<td>22 (9.0)</td>
</tr>
<tr>
<td><strong>Renal crisis</strong></td>
<td>12 (13.8)</td>
<td>17 (7.0)</td>
</tr>
<tr>
<td><strong>Pericardial effusion</strong></td>
<td>1 (1.1)</td>
<td>4 (1.6)</td>
</tr>
</tbody>
</table>
To explore the prevalence and clinical associations of elevated systolic pulmonary artery pressure (sPAP), measured by Transthoracic Doppler-echocardiography (TTE) in patients with early systemic sclerosis (SSc).

- A cross-sectional analysis of the prospective EULAR Scleroderma Trial and Research (EUSTAR) database was performed. **SSc patients with <3 years from the first non-Raynaud’s phenomenon (RP) symptom** at baseline EUSTAR visit, were selected. Elevated sPAP was defined as sPAP>40 mmHg on baseline TTE. First visit SSc related variables, including disease subsets, antibodies and visceral involvement, were examined.

- From 1,188 patients, elevated sPAP was found in 17% of patients, both lcSSc and diffuse cutaneous SSc (dcSSc).
- In lcSSc, older age at first non-RP symptom, ACA positivity, joint contractures, restrictive defect and lower DLCO, were independently associated with elevated sPAP.
- In dcSSc, older age at first non-RP symptom, longer time between RP onset and first non-RP symptom, digital ulcers, cardiac blocks, and proteinuria were associated with elevated sPAP.

The prevalence of elevated sPAP on TTE in early SSc patients is considerable. **Association with cardiac, lung and renal involvement suggests that, although some patients might have pulmonary arterial hypertension, others may present pulmonary hypertension secondary to lung or heart involvement.**

Our findings emphasize the need to consider right heart catheterisation in selected early SSc patients with PH suspicion, to clearly determine the cause of PH.
Reducing the burden of SSc: Value of screening for SSc-PAH

<table>
<thead>
<tr>
<th>Survival</th>
<th>Prevalent PAH</th>
<th>Incident PAH</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>3yr</td>
<td>42.7%</td>
<td>94.7%</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Mean time to death (years)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prevalent PAH</td>
<td>2.3 (±2.3)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Incident PAH</td>
<td>4.7 (±2.4)</td>
<td></td>
</tr>
</tbody>
</table>
## Causes of deaths-non-SSc related

<table>
<thead>
<tr>
<th>Organ system</th>
<th>Inception</th>
<th>Prevalent</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n=34</td>
<td>n=148</td>
</tr>
<tr>
<td></td>
<td>n(%)</td>
<td>n(%)</td>
</tr>
<tr>
<td>Malignancy</td>
<td>13(38.2)</td>
<td>55(37.2)</td>
</tr>
<tr>
<td>Sepsis</td>
<td>5(14.7)</td>
<td>14(9.5)</td>
</tr>
<tr>
<td>CVD</td>
<td>4(11.8)</td>
<td>7(4.7)</td>
</tr>
<tr>
<td>IHD</td>
<td>3(8.8)</td>
<td>18(12.2)</td>
</tr>
<tr>
<td>Liver disease</td>
<td>2(5.9)</td>
<td>3(2.0)</td>
</tr>
<tr>
<td>Post operative</td>
<td>1(2.9)</td>
<td>9(6.1)</td>
</tr>
<tr>
<td>complications</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trauma</td>
<td>1(2.9)</td>
<td>8(5.4)</td>
</tr>
<tr>
<td>Sudden death</td>
<td>1(2.9)</td>
<td>4(2.7)</td>
</tr>
<tr>
<td>Renal failure</td>
<td>1(2.9)</td>
<td>1(0.7)</td>
</tr>
<tr>
<td>Asthma/COPD</td>
<td>0(0)</td>
<td>6(4.1)</td>
</tr>
<tr>
<td>Peripheral vascular disease</td>
<td>0(0)</td>
<td>2(1.4)</td>
</tr>
<tr>
<td>Pulmonary embolism</td>
<td>0(0)</td>
<td>1(0.7)</td>
</tr>
<tr>
<td>Arrhythmia</td>
<td>0(0)</td>
<td>1(0.7)</td>
</tr>
<tr>
<td>Drug related</td>
<td>0(0)</td>
<td>1(0.7)</td>
</tr>
<tr>
<td>Other</td>
<td>3(8.8)</td>
<td>18(12.2)</td>
</tr>
</tbody>
</table>
To analyze the characteristics of anti-RNA polymerase III antibodies (anti-RNAP3)- positive patients with systemic sclerosis (SSc) with a focus on the risk of cancer and the characteristics of malignancies, and the aim to provide guidelines about potential cancer screening in these patients.

- Analysis of the EUSTAR database: **4986 patients** with information on their anti-RNAP3 status were included.
- **Case-control study**: additional retrospective data, including malignancy history, were queried in 13 participating EUSTAR centers.
- **158 anti-RNAP3+ cases** were compared with 199 local anti-RNAP3- controls, matched for sex, cutaneous subset, disease duration, and age at SSc onset. (3) A Delphi exercise was performed by 82 experts to reach consensus for cancer screening in anti-RNAP3+ patients.

- In the EUSTAR registry, anti-RNAP3 were associated in multivariable analysis with renal crisis and diffuse cutaneous involvement.
- In the case-control study, anti-RNAP3 were associated with gastric antral vascular ectasia, rapid progression of skin involvement, and malignancies concomitant to SSc onset (OR 7.38, 95% CI 1.61-33.8). When compared with other anti-RNAP3+ patients, those with concomitant malignancies had older age (p < 0.001) and more frequent diffuse cutaneous involvement (p = 0.008). The Delphi exercise highlighted the need for malignancy screening at the time of diagnosis for anti-RNAP3+ patients and tight followup in the following years.

**Anti-RNAP3+ patients with SSc have a high risk of concomitant malignancy. These results have implications for clinical practice and suggest regular screening for cancer in anti-RNAP3+ patients.**

Causes of death- 1072 out of 11193 SSc patients

SSc-related: 617 (57.6%)  
SSc-unrelated: 270 (25.2%)
Causes of death according to SSc-cutaneous subset
Incidence and Risk Factors of Organ Manifestations in the Early Course of Systemic Sclerosis: A Longitudinal EUSTAR Study

Jaeger VK et al PLOS ONE October 5, 2016

11,290 patients in EUSTAR

9,891 patients fulfil the 1980 ACR classification criteria for SSc
→ the entire EUSTAR cohort

1,399 patients did not fulfil the 1980 ACR classification criteria for SSc

695 patients fulfil the 1980 ACR criteria and have a baseline visit recorded within one year of the onset of RP
→ the study population

9,196 patients fulfil the 1980 ACR classification criteria for SSc but had no baseline visit recorded within one year of the onset of RP

Early SSc were identified as those who had a visit within the first year after RP onset. Incident SSc organ manifestations and their risk factors were assessed using Kaplan-Meier methods and Cox regression analysis.
incidence of organ involvement in SSc patients after RP onset
Heart Involvement

Cumulative percentage of patients with organ involvement

Years after onset of RP

(F) p=0.0002

(G) ACA vs. anti-TOPO: p=0.84
ACA vs. anti-RNAP-III: p=0.25
anti-TOPO vs. anti-RNA-III: p=0.19

(H) p=0.16
Scleroderma renal crisis

Cumulative percentage of patients with organ involvement

Years after onset of RP

ACV vs. anti-TOPO: p=0.013
ACV vs. anti-RNAP-III: p<0.0001
anti-TOPO vs. anti-RNA-III: p=0.0003
Higher burden on internal organ involvement in male patients
The gender enigma !!!

Years after onset of RP
9182 patients with SSc were available (1321 men) for the baseline analyses.

In multivariate analysis, male sex was independently associated with:
✓ a higher risk of diffuse cutaneous subtype (OR: 1.68, (1.45 to 1.94); p<0.001),
✓ a higher frequency of digital ulcers (OR: 1.28 (1.11 to 1.47); p<0.001)
✓ pulmonary hypertension (OR: 3.01 (1.47 to 6.20); p<0.003).

• In the longitudinal analysis (n=4499), after a mean follow-up of 4.9 (±2.7) years, male sex was predictive of new onset of pulmonary hypertension (HR: 2.66 (1.32 to 5.36); p=0.006) and heart failure (HR: 2.22 (1.06 to 4.63); p=0.035).

• 908 deaths were recorded, male sex predicted deaths of all origins (HR: 1.48 (1.19 to 1.84); p<0.001), but did not significantly account for SSc-related deaths.

SSc appears as strikingly more severe in men. Our results obtained through the largest worldwide database demonstrate a higher risk of severe cardiovascular involvement in men. These results raise the point of including sex in the management and the decision-making process.
Incidence and predictors of cutaneous manifestations during the early course of systemic sclerosis: a 10-year longitudinal study from the EUSTAR database
Elina G Wirz et al ARD 2016;75:1285–1292

**to analyse the incidence of skin sclerosis and DUs in patients who developed SSc within 1 year after the onset of Raynaud’s phenomenon**

- The median **modified Rodnan skin score (mRSS)** peaked 1 year after RP onset, and was 15 points.
- The 1-year probability to develop a mRSS ≥2 in at least one area of the arms and legs was 69% and 25%, respectively.
- **25%** of patients developed dSSc in the **first year** after RP onset. This probability increased to **36%** during the subsequent **2 years**. Only 6% of patients developed dSSc thereafter.
- The probability to develop **DUs increased to a maximum of 70% at the end of the 10-year observation**.
- The main factors associated with dSSc were the presence of RNApolIII, followed by Topol and male sex.
- The main factor associated with incident DUs was the presence of Topo I autoantibodies.

**Early after RP onset, cutaneous manifestations exhibit rapid kinetics in SSc.**
Time to peak mRSS. The histogram plots the percentage of patients as a function of the time to reach their maximal mRSS from RP onset; The median peak mRSS was

- 15 points (IQR 7–24) overall, 9.5 points (IQR 6–14) in patients with lSSc involvement
- 23 points (IQR 16–29.5) in patients with dSSc.
Prediction of improvement in skin fibrosis in diffuse cutaneous systemic sclerosis: an EUSTAR analysis

AIM: to identify predictors for improvement of skin fibrosis in patients with dcSSc

From the 919 patients included, 218 (24%) improved and 95 (10%) progressed. Eleven candidate predictors for skin improvement were analysed. The final model identified high baseline mRSS and absence of tendon friction rubs as independent predictors of skin improvement.

The baseline mRSS was the strongest predictor of skin improvement, independent of disease duration. An upper threshold between 18 and 25 performed best in enriching for progressors over regressors.

Patients with advanced skin fibrosis at baseline and absence of tendon friction rubs are more likely to regress in the next year than patients with milder skin fibrosis.

Dobrota R et al ARD 2016;75:1743–1748

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Lower mRSS at inclusion enriches for patients with skin fibrosis progression.

(A) Percentage of progressors and regressors per baseline mRSS range. Patients with lower skin score are more likely to progress, whereas those with higher skin scores are much more likely to regress.

(B) Sensitivity for progression and regression depending on different cut-off values for baseline mRSS.

Dobrota R et al ARD 2016;75:1743–1748
Progression of skin fibrosis is associated with FVC decline at follow up.

Survival Probability

Time to FVC decline ≥ 10% (years)

Patients’ number at risk by time

Non-progressor 34.4%  Progressor 53.6%  p<0.001

Wu et al, oral presentation WSC 2018
Progression of skin fibrosis is associated with all-cause death at follow up.

Non-progressor 7.3%  
Progressor 15.4%  
p = 0.003

Time to All-cause Death (years)

Patients’ number at risk by time

Survival Probability

943 854 657 502 397 307 197 103 16

78 72 54 42 35 23 8 4 0

Years
Determine inter/intra-observer reliability of tender and swollen joint counts (TJC, SJC) and Modified Rodnan Skin Score (MRSS) in diffuse systemic sclerosis (dcSSc) and assess content validity of TJC/SJC

✓ Ten rheumatologists completed SJC, TJC, and MRSS on 7 patients. Musculoskeletal ultrasound (MSUS) was performed

• Inter-observer and intra-observer reliability for TJC was 0.97 and 0.99, for SJC was 0.24 and 0.71, and for MRSS was 0.81 and 0.94, respectively. MSUS abnormalities did not correspond with SJC/TJC

excellent inter and intra-observer reliability for MRSS and TJC in dcSSc. However, SJC and TJC did not correspond to MSUS
Conclusions

• Prevention of skin and organ damage in dcSSc - a new clinical trial concept

• Progression of skin fibrosis can be modeled – short disease duration, lower mRSS, arthritis

• Progression of skin fibrosis is associated with ILD worsening and worse survival at follow up
Conclusions

1. The presence of Malignancy is today an issue which needs to be carefully addressed by the physician.
2. In the first 2 years of the disease the skin and GI and DLCo as well are the most affected.
3. The most severely affected gender is male.
4. Overall, the main clinical issues are the Cardiopulmonary involvement, Cancer and Infections.
5. Progression of skin fibrosis can be modeled identifying a short disease duration and a lower mRSS.
6. Skin progression is associated with ILD worsening and worse survival at follow up.

Prevention of skin and organ damage in dcSSc is a new clinical trial concept.
Registries are very useful to understand the disease and show us their particular realities.

Registries from different continents can show different aspects of the disease.

However, sometimes they do not show the same results.

Speculate about Why
There may be several answers to the *Why*...

1. Selection bias depending on the centers
2. Cultural differences
3. Time when patients are entered
4. Environmental differences
5. Genetic differences

Because of the differences we can design better our RCT
Education opportunities on SSc and its complications

- Online courses
- Books
- Courses
- Hands on courses
- Masterclasses
The EULAR On-line Course on Systemic Sclerosis consists of **10 modules** which deal with physiopathology, clinical aspects and management of this complex disease.

MODULE 1: Introduction Module

MODULE 2: Pathogenesis

MODULE 3: Clinical manifestations: skin, peripheral vascular

MODULE 4: Heart involvement

MODULE 5: Pulmonary interstitial/vascular involvement

MODULE 6: Clinical manifestations: GI

MODULE 7: Clinical manifestations: Kidney

MODULE 8: Clinical manifestations: Musculoskeletal, disability

MODULE 9: Management

MODULE 10: Special conditions
WSF Courses & Masterclasses

• **PAH** Firenze 2011
• **Ulcers** 2015-2016
  ✓ Bucharest
  ✓ Split
  ✓ Tel Aviv
Eustar Courses

• Budapest 2005
• Bad Nauheim 2007
• Paris 2009
• Belgrade 2011
• Cluj Napoca 2013
• Katowice 2015
• Split 2017
• Nijmegen 2019