Heart involvement in systemic sclerosis: an ultrasonic tissue characterisation study

Clodoveo Ferri, Vitantonio Di Bello, Antonella Martini, Davide Giorgi, Franca A A Storino, Massimiliano Bianchi, Alessio Bertini, Marco Paterni, Costantino Giusti, Giampiero Pasero

Abstract

Background—Clinicoepidemiological findings indicate that symptomatic heart involvement in patients with systemic sclerosis (SSc) predicts a very poor prognosis. At necropsy studies, SSc heart involvement without significant coronary lesions is characterised by patchy myocyte necrosis and contraction band necrosis with collagen replacement leading to myocardial fibrosis. There is a discrepancy between the frequency of clinically evident myocardial disease (25%) and autoptical myocardial fibrosis (81%).

Objective—The aim of this study was to detect preclinical myocardial alterations in SSc patients by ultrasonic videodensitometric analysis.

Methods—Thirty five SSc patients (three male, aged 48.6 (11) SD years, range 22–65) with normal ventricular function and 25 age and sex matched healthy controls were studied. All patients had a negative maximal exercise stress; in all cases arterial hypertension, renal involvement, and diabetes were excluded. Echocardiographic images were digitised by a real time videodigitiser (Tomtec Imaging Systems). Quantitative texture analysis was performed on data from the septum and the posterior wall, obtaining mean gray level histogram (MGL) at both end-diastole (d) and end-systole (s). The cyclic variation index (CVI), was calculated according to the formula ((MGLd−MGLs)/MGLd) × 100. Left ventricular mass (LVM), body surface corrected, was calculated according to Penn convention.

Results—Comparable systolic and diastolic blood pressure, LVM, diastolic and systolic function were recorded in both SSc patients and controls. In contrast, in SSc patients the CVI, which is the expression of the intrinsic myocardial structural function, was significantly lower than in controls (septum: −18 (28)% vs 35 (10)%, p<0.0001; and posterior wall: −13 (32)% vs 50 (20)%, p<0.0001). Changes in cyclic echo amplitude, probably related to myocardial fibrosis, were detected in the large majority of SSc patients (88%).

Conclusions—Ultrasonic videodensitometric analysis represents a non-invasive, feasible method that can detect early myocardial changes in SSc patients, which could be related to both fibrosis and microcirculatory abnormalities. Their potential evolution towards ventricular dysfunction and their link with cardiac sudden death, because of severe conduction system or rhythm disturbances, should be further investigated.

Systemic sclerosis (SSc) is a connective tissue disease characterised by fibrosis of multiple target organs including the skin, lung, gastrointestinal tract, kidney, and heart.1 Invasive and non-invasive diagnostic investigations, as well as necropsy studies showed that scleroderma cardiac involvement is one of the most frequent visceral complication that can affect the overall prognosis of the disease.1–8 More interestingly, heart involvement can appear independently from other typical complications of SSc.9 Although the myocardial fibrosis is a frequent pathological finding, cardiac involvement can be clinically silent. In particular, congestive heart failure and myocardial infarction are uncommon, while conduction system and rhythm disturbances were frequently recorded.5,10 Necropsy studies9 11 demonstrated that scleroderma hearts without significant coronary lesions are frequently characterised by patchy myocardial fibrosis in both ventricles, resulting from collagen deposition in the areas of myocyte necrosis. Moreover, the presence of contraction band necrosis suggested that myocardial microcirculation disturbances, the so called “myocardial Raynaud’ phenomenon” are responsible for the myocardial fibrosis, as indirectly supported by in vivo clinical investigations.12–17

Quantitative analysis of two dimensional echocardiographic (2D echo) pattern, or “texture”, represents a novel approach for myocardial tissue characterisation.18–19 This technique was able to identify acute ischaemic and contused myocardium lesions in animal models,18 19 as well as in human heart diseases including amyloid, diabetes, hypertrophic cardiomyopathies, myocarditis, myocardial ischaemia, and more recently hypertensive cardiopathy.20–22 The aim of this study was to detect changes in myocardial structure by means of ultrasonic videodensitometric analysis in a series of SSc patients without clinical evidence of ventricular disfunction or other complications responsible for secondary heart involvement.

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Cutaneous involvement (skin score), 27 presence of SSc; namely, extent of the disease was carefully investigated and reported previously; in particular, all patients underwent a physical examination, chest radiograph (heart size, lung fibrosis, size of main pulmonary arteries), pulmonary function testing (total lung capacity, forced vital capacity, forced expiratory volume in one second), diffusing lung capacity for carbon monoxide (DLco).

Methods

Thirty five SSc patients (three males, aged 48.6 (11) years, mean (SD), range 22–65) and a sex and age matched group of 25 control subjects (two males; mean age 45.6 (10), range 25–63) without known systemic, immunological, and cardiovascular diseases were selected for the study.

All patients met the American College of Rheumatology (formerly, the American Rheumatism Association) 1980 preliminary criteria for the classification of SSc, 31 patients had a limited cutaneous involvement (distal to elbows and knees) and four the diffuse variant. 4

Patients were consecutively recruited at the routine ambulatory control visits according to the following inclusion criteria: age ≤ 65 years; absence of heart failure on clinical examination with normal chest radiographs and left ventricular systolic function at echocardiographic examination; absence of myositis, arterial hypertension, renal involvement, diabetes, and any other systemic diseases. In all cases ECG maximal exercise stress test was negative. All subjects had given their informed consent to enter the study, which was approved by our Institutional Ethical Committee.

At the time of the study, all patients underwent a wide evaluation of the signs and symptoms of SSc; namely, extent of the cutaneous involvement (skin score), 27 presence of telangiectasias, cutaneous calcinosis (radiographs), hypermelanosis, cutaneous ulcers, Raynaud’s phenomenon, arthritis, oesophageal involvement (radiographic hypomotility with or without dysphagia), and nephropathy (proteinuria >300 mg/24 hours and/or serum creatinine >1.2 mg/dl). The presence of lung involvement was carefully investigated, as previously reported; in particular, all patients underwent a physical examination, chest radiograph (heart size, lung fibrosis, size of main pulmonary arteries), pulmonary function testing (total lung capacity, forced vital capacity, forced expiratory volume in one second), diffusing lung capacity for carbon monoxide (DLco).

Nailfold capillary examination was performed using a Leitz microscope and interpreted according to the procedures of Maricq et al. 28 Capillaroscopic changes included diffuse tortuosity, enlarged capillary loops, reduced number of capillaries (total score 4.5).

In all patients the following serological markers were detected using standard techniques: anti-nuclear (ANA) antibodies by indirect immunofluorescence on rat liver at a dilution of 1:20, and on Hep 2 cell line at a dilution of 1:40; anti-centromere antibody (ACA) 39; anti-extractable nuclear antigen (ENA) antibodies, including anti-Scl70, -Sm, -RNP, -SSB, -SSA, -PCNA, -SL and Jo1. 4, 31

Cardiological investigations

In all patients resting and exercise ECG, 24 hour Holter ECG monitoring were performed and evaluated as previously described. 4

Conventional echo Doppler analysis

Conventional echocardiographic studies were performed with a Hewlett-Packard (Andover, Mass, USA) 77020A phased array sector scanner with a 2.5 or 3.5 MHz transducer. Two dimensional images were obtained in the parasternal long axis and short axis views and apical two and four chamber views. Left ventricular diameters (EDD: left ventricular end diastolic diameter), end diastolic septal thickness (Sthd) and posterior wall thickness (PWthd) were also measured, according to the criteria of the American Society of Echocardiography. 32 Left ventricular per cent fractional shortening was calculated as end diastolic diameter minus end systolic diameter divided by end diastolic diameter multiplied by 100. Left ventricular mass was calculated by the Devereux formula (Penn convention) 41 and adjusted for body surface area (LVMi(BS)).

Pulsed Doppler transmitral flow velocity profile was obtained from the apical four chamber view, and the sample volume was positioned just below the mitral valve leaflets. The following parameters were evaluated: peak E (peak transmitral flow velocity in early diastole); peak A (peak transmitral flow velocity in late diastole); E/A ratio. Video recordings of M-mode imaging measurements were analysed off line by an experienced echocardiographer, using the Hewlett-Packard software of analysis. All recordings were analysed on two separate occasions within one week, for intraobserver variability, to assess the reproducibility of these measures, as well as by a blinded investigator for interobserver variability. Both interobserver and intraobserver variability were minimal.

| Table 1 Demographic and clinico-serological data of 35 SSc patients |
|----------------|----------------|
| **Age (y; mean (SD))** | 48.6 (11) |
| **M/F ratio** | 3/32 |
| **Disease duration (y; mean (SD))** | 8.3 (6.0) |
| **Raynaud’s syndrome** | 97% |
| **Calcification** | 26% |
| **Telangiectasias** | 86% |
| **Oesophageal involvement** | 74% |
| **Lung involvement** | 88% |
| **Renal involvement** | 0% |
| **Autoantibodies** | |
| **ANA** | 82% |
| **ACA** | 40% |
| **anti-Scl70** | 37% |

ANa = anti-nuclear; ACA = anti-centromere antibodies.
Quantitative textural analysis
(videodensitometry)
For a reproducible sampling of textural ultrasonic parameters, the gain settings and the gain compensation profiles were adjusted to obtain an apparently uniform myocardial brightness. The gray scale transfer function was adjusted to be linear for the entire video signal range, and no reject, no enhancement of dynamic range were used (18–19) with a 25–30 dB amplification at a depth of 18 cm. The optimal echocardiographic images were directly transferred from the echocardiograph to a calibrated video digitisation system. Images were converted into 256 × 256 pixels of 256 gray levels each (0 = black, 255 = white) with eight bits of intensity range, by a commercially real time videodigitiser (Tomtec Imaging Systems, Boulder, Colorado, USA). One cardiac cycle (R-R wave) was automatically divided in 12 frames, independently by heart rate, and the images corresponding to the end diastolic and end systolic phases, all in left parasternal long axis view, were selected with an optimal visualisation of the interventricular septum and of the left ventricular posterior wall. Using an interactive computer program the region of interest, always the same size (32 × 42 pixels), was placed at mid-septum and mid-posterior wall level, both in end systole and end diastole, including only the myocardium and excluding the endocardial and epicardial specular echoes to avoid areas of echo dropouts and obvious artefacts. For each region of interest, a histogram of the echocardiographic gray level distribution was generated, plotting the gray level distribution on the abscissa and the frequency of the occurrence on the ordinate. The mean gray level in each cavity region (background signal) was subtracted from the absolute mean gray level obtained at myocardial level obtaining both end systolic mean gray level (MGLes) and end diastolic mean gray level (MGLed). Cyclic variation index of gray level amplitude was also calculated according to the formula: \( \text{CVI} = \frac{\text{MGLed} - \text{MGLes}}{\text{MGLed} \times 100} \). All measurements were obtained from the average of five consecutive cardiac cycles. Coefficient of variation regarding MGL, both at end systolic and end diastolic values, ranged from 8% to 10% for interobserver, and from 9% to 12% for intraobserver variability.

**Table 2** Conventional echo Doppler parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Systemic sclerosis (n=35)</th>
<th>Controls (n=25)</th>
<th>Analysis of variance</th>
</tr>
</thead>
<tbody>
<tr>
<td>EDD (mm)</td>
<td>44.5 (4.1)</td>
<td>45.7 (3.6)</td>
<td>0.21</td>
</tr>
<tr>
<td>FS (%)</td>
<td>39.2 (6.4)</td>
<td>40.7 (6.2)</td>
<td>0.31</td>
</tr>
<tr>
<td>LA (mm)</td>
<td>32.4 (5.9)</td>
<td>31.3 (4.0)</td>
<td>0.39</td>
</tr>
<tr>
<td>Sthd (mm)</td>
<td>9.3 (1.1)</td>
<td>8.2 (1.1)</td>
<td>0.02</td>
</tr>
<tr>
<td>Pwthd (mm)</td>
<td>8.9 (1.2)</td>
<td>8.0 (1.2)</td>
<td>0.06</td>
</tr>
<tr>
<td>LVMi (BS) (g/m²)</td>
<td>83.7 (20.2)</td>
<td>81.5 (19.8)</td>
<td>0.19</td>
</tr>
<tr>
<td>Peak E (m/sec)</td>
<td>0.69 (0.22)</td>
<td>0.80 (0.21)</td>
<td>0.07</td>
</tr>
<tr>
<td>Peak A (m/sec)</td>
<td>0.70 (0.12)</td>
<td>0.57 (0.13)</td>
<td>0.007</td>
</tr>
<tr>
<td>E/A ratio</td>
<td>1.03 (0.42)</td>
<td>1.44 (0.28)</td>
<td>0.001</td>
</tr>
</tbody>
</table>

MGLed = Mean gray level (background corrected) in end diastole; MGLes = mean gray level (background corrected) in end systole; CVI = cyclic variation index.

**Table 3** Ultrasonic textural data (first order analysis)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Systemic sclerosis mean (SD)</th>
<th>Controls mean (SD)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>MGLed (septum)</td>
<td>65.8 (24.2)</td>
<td>89.9 (13.5)</td>
<td>0.001</td>
</tr>
<tr>
<td>MGLes (septum)</td>
<td>77.2 (18.4)</td>
<td>50.0 (14.7)</td>
<td>0.001</td>
</tr>
<tr>
<td>CVI (%) (septum)</td>
<td>−23.7 (30.3)</td>
<td>35.9 (20.9)</td>
<td>0.001</td>
</tr>
<tr>
<td>MGLed (p. wall)</td>
<td>78.1 (27.6)</td>
<td>92.3 (15.8)</td>
<td>0.01</td>
</tr>
<tr>
<td>MGLes (p. wall)</td>
<td>88.5 (28.7)</td>
<td>44.9 (18.4)</td>
<td>0.005</td>
</tr>
<tr>
<td>CVI (%) (p. wall)</td>
<td>−20.5 (32.7)</td>
<td>50.9 (20.6)</td>
<td>0.0001</td>
</tr>
</tbody>
</table>

MGLed = Mean gray level (background corrected) in end diastole; MGLes = mean gray level (background corrected) in end systole; CVI = cyclic variation index.

**Results**
Table 1 shows the clinicopneumological and serological findings of 35 SSc patients. Among clinical manifestations lung involvement, Raynaud’s phenomenon, and telangiectasias were the most frequently found. In all patients the presence of one or more serum autoantibodies was recorded.

Ambulatory 24 hour Holter ECG monitoring showed the presence of significant arrhythmias (premature ventricular complexes ≥ 200/24 hours) in four of 35 SSc patients. In our group of SSc patients no conduction disturbances were detected during ECG Holter monitoring. The four patients with significant premature ventricular complexes, showed a clear cut abnormal ultrasonic textural pattern.

On the whole, the main values of echocardiographic parameters were comparable in both SSc patients and controls (table 2). In particular, mean values of fractional shortening, which represents a global index of systolic function, were comparable in both groups; similarly, left ventricular mass (body surface indexed) did not show significant differences between patients and controls.

Of the Doppler parameters, evaluating left ventricular diastolic filling, late peak flow (peak A) velocity was significantly higher in SSc patients than controls (p<0.007, table 2), while early peak flow (peak E) was reduced, but not significantly, if compared with normal controls. The E/A ratio was significantly lower in SSc
patients than in normal subjects (p<0.001). When individual patients’ analysis was performed, E/A ratio was under the normal range (95% confidence intervals) in only five of 35 (15%) of patients (fig 1).

Table 3 shows the videodensitometric parameters, in particular the mean gray level of the septum and of the posterior wall, both at end systole and end diastole. The mean values of CVI for both septum and posterior wall were significantly lower in SSc patients than controls (table 3, and fig 2; p<0.0001). In SSc patients the same parameter was abnormally reduced (< 95% confidence intervals) in 80% of cases (28 of 35) for the septum and in 88% (31 of 35) for posterior wall, respectively.

Figure 3 compares the digitised 2D echo images of the left ventricle and the diagrams of echo intensity, in a selected region of the septum during one cardiac cycle, in one normal subject and one SSc patient. The mean gray values were totally changed in the patient compared with the normal subject; in particular, in the SSc patient the echo intensity values, during the cardiac cycle, showed a paradoxical behaviour.

No significant correlations were found between CVI and septum or posterior wall thickness, LVMi, FS%, and diastolic Doppler transmitral flow parameters (peak E, peak A, E/A ratio).

Finally, no significant correlations were detected between epidemiological (age, sex, disease duration), clinical (extent of cutaneous involvement, nailfold capillaroscopy changes, skin calcinosis, lung involvement) and serological features (anti-centromere, anti-Scl70 antibodies) and textural videodensitometric findings. Interestingly, during the time period after this study one patient died because of severe cardiopulmonary involvement. This patient was one of those with more pronounced videodensitometric changes (CVI septum: −102.7%).

![Figure 2](http://ard.bmj.com/)

Figure 2  Cyclic variation index (CVI) at mid-septum and at mid-left ventricular posterior wall level for both systemic sclerosis (SSc) and controls. −2SD: 2 standard deviations under the mean value of control subjects.

![Figure 3](http://ard.bmj.com/)

Figure 3  Comparison between digitised 2D echo images of the left ventricle (left side) and the diagrams of echo intensity variations (right side), in a selected region of the septum during one cardiac cycle, in one normal subject (A) and one SSc patient (B). Arrow 1 indicates the interventricular septum: the patient shows a clear cut increase in echo intensity suggesting the presence of fibrosis; arrow 2 indicates the left ventricular cavity for background subtraction. In the SSc patient the diagram of echo intensity variations shows a paradoxical behaviour compared with the normal subject.
Discussion
Heart involvement in SSC patients is often clinically occult and its detection depends, at least in part, on the sensitivity of the diagnostic methods used. It has been estimated that scleroderma heart disease is clinically evident in only 20–25% of cases; in these people the prognosis becomes severe and a 70% of mortality at five years occurs. The discrepancy between the percentage of symptomatic heart involvement and necropsy scleroderma myocardial fibrosis (30%–81%) should be improved by the introduction of more sensitive, possibly non-invasive, diagnostic tools.

This study demonstrated that myocardial tissue changes were detectable in the large majority of SSC patients (88%) by means of ultrasonic videodensitometric analysis. In particular, the CVL, which is the expression of the intrinsic myocardial structural function was significantly lower, for both the septum and posterior wall, in scleroderma patients than in age matched controls. These findings can be the expression of preclinical heart abnormalities; in fact, all SSC patients had a normal left ventricular systolic function as well as a normal left ventricular mass. Coronary artery disease and other disorders responsible for myocardial damage—that is, arterial hypertension, diabetes, and other cardiomyopathies—have also been excluded in the patients in this series. Moreover, left ventricular diastolic function was seldom impaired; actually, a change in E/A ratio was detected in only 15% of patients; this is particularly true as our age and sex matched control group showed a mean (SD) value of E/A ratio comparable with that reported by other authors. The diastolic change observed in SSC patients mainly regards the relaxation phase, resulting in an increase of atrial contribution to filling (peak A) and a decrease of early diastolic filling (peak E). Left ventricular diastolic function depends on the complex interaction of ventricular isovolumic relaxation, diastolic filling, and compliance. Abnormalities of the Doppler transmitral profile have been described in a large number of cardiac diseases; unfortunately, there is a wide overlap of individual data points between patients and controls, which prevents the routine evaluation of diastolic function in individual patient by Doppler echocardiography. The rarity of diastolic changes and much more the absence of systolic left ventricular dysfunction in our SSC patients emphasise the clinical relevance of myocardial tissue changes detected by videodensitometry.

Several clinical studies focusing on the heart involvement in SSC patients have been performed by means of traditional echo Doppler, which showed various abnormalities—that is, right ventricular enlargements, concentric left ventricular hypertrophy with asymmetrical septal hypertrophy, impaired shortening fraction or hyperdynamic systolic function, diastolic functional abnormalities, left atrial enlargement, pericardial effusion. On the whole, echocardiographic left ventricular systodiatolic myocardial dysfunction have been detected in a limited percentage of patients, ranging from 11% to 32%. In contrast, high incidence (71%–100%) of thallium perfusion abnormalities was detected in SSC patients. The coronary angiography usually shows normal epicardial coronary arteries, while coronary reserve was often impaired; these findings suggested a prevalence of microcirculatory disturbances. The scintigraphic abnormalities are strikingly associated with pathological lesions, even in the absence of left ventricular dysfunction. Similarly, the high incidence of videodensitometric abnormalities observed in this study could reflect the presence of myocardial structural changes, detectable at a preclinical level.

The myocardial involvement in SSC has been reported in up to 81% of SSC patients at necropsy. The main pathological findings were represented by extensive focal myocardial fibrous replacements that are prevalent in perivascular areas and contraction band necrosis, a pattern similar to that observed after reperfusion following transient coronary occlusion. Other necropsy findings, namely focal infiltrates of round cells and thickening of smaller coronary vessels, were less frequently represented. However, the exact mechanism leading to the myocardial fibrosis in SSC remains unknown. The paucity of structural vascular disease in the presence of contraction band necrosis could be explained by different pathogenetic hypotheses: namely, functional abnormalities such as small vessel vasospasm, the so called “intramyocardial Raynaud’s phenomenon”, or structural intramyocardial microcirculatory changes. The presence of exercise and cold induced thallium perfusion abnormalities and abnormal coronary vasodilator reserve, together with the above pathological findings suggested that scleroderma myocardial changes are the consequence of various cofactors. Among these myocyte replacement by collagen and coronary vasospasm seem to be the main pathogenetic factors of the myocardial fibrosis. The possible contribution of small artery obliterative disease and the damage of coronary capillary is suggested by in vivo physiological studies such as thallium perfusion changes and impaired coronary reserve; however, preliminary necropsy studies did not confirm this hypothesis.

The high frequency of myocardial texture changes detected in SSC patients is particularly significant if compared with that recorded in other disorders, in which comparable changes, but to a lesser degree, were found. In normal subjects a cardiac cycle dependent variation in ultrasound signals within the myocardium was detectable: maximum values occurred at end diastole, minimal values at end systole. These cyclic changes could be related to intrinsic contractile function, distinct from myocardial wall thickening, although the physiological basis of this index is not fully understood and it is almost certainly multifactorial. For the first time we applied the videodensitometric approach to study the health of SSC patients.
The significantly lower values of CVI are essentially because of the abnormal increase of the mean gray level at the end systole, than in control the reduction of CVI could mainly be related to scleroderma myocardial fibrosis, which causes an increase in acoustic impedance. This finding is mirrored by the high percentage of patients with myocardial fibrosis, the hallmark of scleroderma heart involvement, detected in necropsy studies.\(^9\) More interestingly, the results of this study further support the demonstration of myocardial fibrosis at necropsy and contraction band necrosis even in patients without clinical evidence of myocardial dysfunction.\(^9\) This latter condition was indirectly confirmed in our patients by the lack of correlation between myocardial changes and patients’ age and disease duration as well as other prognostically relevant clinicorheological manifestations—that is, skin sclerosis extent, capillaroscopic changes, lung involvement, and serum anti-Scl-70 antibody. In the light of this, it could be supposed that primary myocardial involvement in SSC patients represents a precocious event, characterised clinically by infrequent evolution to overt heart failure and more frequently by conduction system abnormalities, arrhythmias, and spontaneous cardiac death. More severe, life threatening arrhythmias.\(^2,4,9\)

In conclusion, videodensitometry represents a non-invasive, feasible ultrasound tissue characterisation method, able to early detect myocardial changes in SSC patients. The physiological and clinicorheological relevance of ultrasonic myocardial abnormalities in scleroderma patients needs to be further evaluated by prospective clinicopathological studies. In this respect, the patient with rather marked videodensitometric changes who died during the follow up suggests a potential prognostic relevance of such myocardial changes. Moreover, the analysis of scleroderma heart disease, mainly at a preclinical level, can give us new insights in the important field of primary cardiomyopathies.

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