Safety and efficacy of recombinant γ interferon in the treatment of systemic sclerosis

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Abstract

Objective—To evaluate the safety and efficacy of recombinant γ interferon (rIFNy) in the treatment of patients with systemic sclerosis.

Methods—Sixteen patients with systemic sclerosis were treated with r-IFNγ, 60 μg m⁻² (low dose, n=10) and 150 μg m⁻² (high dose, n=6), three times weekly in an open phase I/II trial of eight months duration. The patients were stratified in low and high dosage according to the severity and the extent of scleroderma; the two groups were comparable.

Results—The treatment was well tolerated. The most common side effects, almost certainly related to r-IFNγ, were fever, chills, dizziness, headache, and severe flu-like syndrome with decreasing intensity with the time of treatment. Severe aphthous stomatitis (n=1), ventricular tachycardia (n=1), severe oesophageal ulcers due to gastro-oesophageal reflux (n=1), disease exacerbation alone with frank arthritis and slight pericardial effusion (n=1), and inability to conform to the requirements of the study (n=1) were the reasons for discontinuing treatment. Side effects and degree of response were evident during the first five months of treatment. A significant decrease in mean skin thickness score was observed and was higher in the high dose group. Reactive oxygen species of peripheral neutrophils and soluble interleukin-2 receptor serum concentrations were higher than those of normal individuals at study entry and decreased in parallel with clinical improvement.

Conclusions—Treatment of systemic sclerosis patients with r-IFNγ was relatively safe and well tolerated for doses as high as 150 μg m⁻² three times weekly. Side effects and degree of response can be seen during the first months of therapy and can be used as predictors of ultimate toxicity or response. The drug seems to be effective in treating cutaneous scleroderma.

Methods

Sixteen patients (15 female, one male) were enrolled in a phase I/II clinical study, (table 1). The patients had clinical evidence of sclerodematous skin involvement (oedematous or hidebound) proximal to wrist, ankle, and face. Patients 6, 13, 14, and 15 had skin thickening restricted to sites distal to the elbow and knee, but also involving the face and neck, and these patients were considered to have limited cutaneous systemic sclerosis (ISSc). The remaining patients had skin thickening centrally to the elbows and knees and nine of them (Nos 1, 2, 5, 7, 8, 9, 10, 12, and 16) had also trunk involvement; these patients were considered to have diffuse cutaneous systemic sclerosis (dSSc), as shown in table 1. The sclerodematous changes occurred during a period of 1 to 46 months before treatment. The patients were able and willing to conform to the requirements of the study and gave a written consent. Thirteen patients were followed up in the rheumatology outpatient clinic, Department of Internal Medicine, University of Ioannina, and three patients at the rheumatology outpatient clinic, Department of Pathophysiology, Laikon Hospital, National University of Athens. The following exclusion criteria were used for the selection of the patients:
(1) Scleroderma secondary to exposure to L-tryptophan, vinyl chloride, or bleomycin.

(2) Significant internal organ damage according to the following criteria: (a) significant renal disease defined by serum creatinine > 265 μmol l⁻¹, or renal crisis defined by a rise in diastolic blood pressure above 110 mm Hg during the final week of observation, associated with haematuria or proteinuria; microangiopathic haemolytic anaemia, retinal haemorrhages, papilloedema, or an increase in creatinine levels over 70% of baseline; (b) significant lung involvement defined by forced vital capacity (PCV) < 50% and/or diffusion capacity of carbon dioxide (DLCO) < 40% of predicted; (c) significant heart involvement defined by left ventricular ejection fraction (LVEF) < 40% of predicted, or by arrhythmias requiring medical treatment; (d) significant gut involvement defined by oesophageal ulcers, diarrhoea severe enough to require antibiotic treatment, or abdominal pain with air fluid levels on abdominal upright radiogram in the absence of identifiable causes other than systemic sclerosis.

(3) Drug treatment that might independently influence outcome of the trial, such as: prednisolone > 10 mg per day, angiotensin converting enzyme inhibitors, proton-pump inhibitors, or therapy with D-penicillamine ≥ 500 mg per day, methotrexate, or cyclosporin within the preceding three months.

End points suggesting discontinuation of the treatment were: doubling of serum creatinine, renal crisis, newly developed hypertension requiring angiotensin converting enzyme inhibitor therapy, FVC < 50% of predicted, DLCO to alveolar volume ratio (DLCO/VA) < 40% of predicted, LVEF < 40% of predicted, arrhythmias requiring medical treatment, multiple air fluid levels in the upright abdominal radiogram in the absence of other causes, oesophageal ulcers due to gastro-oesophageal reflux, and diarrhoea severe enough to require antibiotic treatment. Finally, doubling of skin thickness score for more than three months indicated serious disease exacerbation constituting an end point for skin involvement and suggested discontinuation of the treatment.

DOSE AND ADMINISTRATION
All patients received three weekly intramuscular injections of r-IFNγ (specific activity 3 × 10⁷ units mg⁻¹, supplied by Boehringer-Ingelheim-Hellas SA). The r-IFNγ preparations were supplied, ready for use, in bottles each containing 100 μg of r-IFNγ. The r-IFNγ injections were given in a medical outpatient facility. Ten patients received 60 μg m⁻² of r-IFNγ per injection (low dose) and six patients received 150 μg m⁻² of r-IFNγ (high dose) per injection.

PATIENT STRATIFICATION
The low dose group was comparable with the high dose group in terms of initial mean skin thickness score [15.3 (SD 9.4) v 18.7 (9.3), P = 0.37], age [45.4 (15.6) v 40.3 (16.4) years, P = 0.55], disease duration [15.5 (12.7) v 22.3 (16.5) months, P = 0.66], DLCO [73 (21) v 67 (24) % predicted, P = 0.87], LVEF [70 (10) v 70 (9) % predicted, P = 0.87], Po₂ [11.3 (0.73) v 12.2 (1.06) kPa, P = 0.12], and presence of gastro-oesophageal reflux (6/10 v 3/6 patients positive). Treatment with r-IFNγ was scheduled for eight months.

PREVIOUS AND CONCOMITANT TREATMENT
Patients 1, 2, 9, 12, and 16 had received a previous drug treatment: D-penicillamine (patients 1 and 12) or methotrexate. Treatment was discontinued because of side effects (patient 2) or no response (the remaining patients). Five patients from the low dose group and two patients from the high dose group were taking 20 mg omeprazole daily for gastro-oesophageal reflux. Two additional patients, one from each group, received omeprazole during the study after institution of r-IFNγ treatment because of gastro-oesophageal reflux. Three patients from the low dose group and two patients from the high dose group were receiving 7.5–10 mg prednisolone at study entry, and one patient (patient 12) also received prednisolone 10 mg daily after institution of rIFNγ treatment. Two patients from the low dose group and two from the high dose group were taking acetylsalicylic acid 100 mg per day.

STUDY EVALUATION
A history was taken and physical examination performed at study entry, at two week intervals for the first two months, and each month thereafter. One physician was responsible for selecting patients, keeping their records, and managing the patients’ problems related either to their disease or to their treatment. Three other doctors, unaware of the treatment, participated in the study as follows. A cardiologist evaluated the heart by echocardiogram for cardiac effusion, valve abnormalities, and measurement of left ventricular ejection fraction.
fraction at baseline, and after four, six, and eight months of treatment. A pulmonary physician evaluated lung function by clinical examination. This involved assessing and grading dyspnoea using a well accepted dyspnoea scoring system27 at each patient visit; spirometric evaluation at baseline and at four and eight months of treatment; and measuring arterial oxygen pressure with the patient at rest, at baseline and at four and eight months of treatment. A chest x ray was taken at baseline and at the end of the study, and interstitial lung disease was assessed as previously described.27

One physician evaluated kidney function and the skin thickness score at each patient visit. Skin involvement was assessed at each patient visit by measuring a skin thickness score, according to a modification of the system described by Kahaleh et al, as follows: the body was divided into 15 zones—head and neck, anterior chest, anterior abdomen, posterior chest, buttocks, and five bilateral sites: upper arm, forearm and dorsum of hands, thigh, lower leg, and dorsum of the feet. Patients with scleroderma limited to the fingers and face were not included in the study; therefore measurement of sclerodactyly in the fingers was not included in the evaluation of the total skin score. The gradings were: 0, normal skin; 1, thickened skin; 2, thickened and unable to pinch; 3, thickened and unable to move. Individual zone scores were summed to obtain a total score. Therefore the range of possible skin thickness scores was from 2 (patient with mild skin involvement above the wrists) to 45 (patient with trunk involvement). In addition, at each patient visit, the oral aperture was measured as the distance from the upper to the lower denture borders on full, active mouth opening. All the patients enrolled in the study were followed up on the same day as other patients with systemic sclerosis who did not participate in the study. Complete blood cell count, platelets, erythrocyte sedimentation rate, prothrombin time, serum electrolytes, serum glutamic oxaloacetic transaminase (SGOT), serum glutamic pyruvic transaminase (SGPT), alkaline phosphatases; lactic dehydrogenase, albumin, serum creatinine, cholesterol, triglycerides, and urinalysis were performed at each visit.

AUTOANTIBODIES IN THE SERA OF PATIENTS UNDER STUDY

Antinuclear antibodies (ANA) were detected by indirect immunofluorescence using Hep-2 cells as substrate. In addition, antibodies to extractable cellular antigens were evaluated in all the patients using both counterimmunoelectrophoresis and western blotting.24 The sera of all the patients were positive for ANA in a titre ranging from 1/320 to 1/1280 with a fine speckled pattern, with the exception of the sera from patients 15 and 16 which gave a pattern compatible with the presence of anti-centromere antibodies (ACA). Counterimmunoelectrophoresis revealed that the sera of patients 1, 4, 5, 9, and 12 were positive for anti-topoisomerase-I antibodies (anti-topoI). The presence of anti-topo-I and ACA was reconfirmed with western blotting. Antibodies to U1RNP, SmRo/SSA, and La/SSB were not detected in the sera under study.

MEASUREMENTS OF REACTIVE OXYGEN SPECIES OF PERIPHERAL NEUTROPHILS

At baseline and after three months of treatment with r-IFNy, peripheral neutrophils of the patients were isolated from 25 ml of blood by ficoll-hypaque density gradient centrifugation as previously described25 and 1.5 × 10⁶ cells in 1 ml of Roswell Park Memorial Institute (RPMI 1640) medium were activated by 200 μg ml⁻¹ phorbol myristate acetate (PMA) in the presence of 80 μM cytochrome C; the reduction of cytochrome C was measured by a Shimatzu spectrophotometer as previously described.25 Reduction of cytochrome C was used as a measure of the production of reactive oxygen species (ROS) by the neutrophils as previously described.25 The patients were free of r-IFNy on the day of blood sampling as well as on the previous day, and in each experiment one or two patient samples were tested along with two samples of age and sex matched normal individuals.

SOLUBLE INTERLEUKIN-2 RECEPTOR IN THE SERA OF PATIENTS

The soluble interleukin-2 receptor (SIL-2R) was determined by the CELLFREE interleukin-2 receptor test kit (T Cell Diagnostics) according to the manufacturers’ instructions: 50 μl of standard kit controls and sera tested in duplicates from patients and from 50 normal individuals were pipetted into a microtitre plate and precoated with murine monoclonal antibody to human IL-2R along with 50 μl of a second murine monoclonal antibody to human IL-2R conjugated with horseradish peroxidase. The plate was incubated for 3 h on a rotator at room temperature, washed thoroughly, and 100 μl of a chromogen solution were pipetted into the wells. After 30 min incubation at room temperature the reaction was stopped and the absorbance was read at 490 nm. The samples values were determined from a standard curve.

STATISTICAL ANALYSIS

A paired difference t test on each patient’s skin score, oral aperture, LVEF % predicted, DLCO % predicted, P0₂ at baseline versus four, six, and eight months of treatment was performed to evaluate the effects of treatment on skin, heart and lung involvement. One-way analysis of variance (ANOVA) was performed in order: (a) to evaluate the effects of the treatment on the ability of the patients’ peripheral neutrophils to produce reactive oxygen species in response to PMA, at baseline versus three months of therapy, and compare this with the ability of peripheral neutrophils from age and sex matched normal individuals to produce reactive oxygen species; and (b) to compare the SIL-2R sera levels before and after r-IFNy treatment with the SIL2R sera levels of normal individuals. Regression analysis was performed as a tool: (a) to identify a relation between oral aperture and skin thickness score, and (b) to identify a relation
between skin thickness score, oral aperture, LVEF % predicted, and DLCO % predicted with the duration of treatment. A Mann-Whitney U test was performed to compare the low with the high dose group in terms of age, disease duration, DLCO % predicted, Po2 (kPa), LVEF % predicted, and skin thickness score. The Spearman rank correlation coefficient was used to test the hypothesis that there is a trend in pulmonary and heart function as the skin score and/or oral aperture change.

**Results**

**SKIN THICKNESS SCORE**

As shown in Table 1, the patients were subdivided into those receiving low dose of IFNy (60 μg m^-2^) and those receiving high dose (150 μg m^-2^). Four patients (patients 2, 5, 7, and 12) withdrew from the study after treatment for 18, 17, 19, and 16 weeks respectively because of severe oesophageal ulcers, exacerbation of the skin involvement, personal reasons (one patient moved to another country), and severe aphthous stomatitis respectively. One additional patient (No 16) discontinued treatment after the third injection of r-IFNy because of ventricular tachycardia.

Individual skin thickness scores at study entry and after eight months of treatment are shown in Table 1. The mean skin thickness score decreased from an average of 17 at study entry to 7 at the end of treatment in the 11 patients who completed the study, with a mean percentage change in skin thickness score of −60.5%. The mean skin thickness score also decreased in the patients who discontinued treatment, from 15.5 at baseline to 14.8, with a mean percentage change in skin thickness score of −23.5%. Statistical analysis using a paired difference t test on each patient's skin score at baseline versus four months of treatment indicated a statistically significant decrease in the mean skin thickness score for 15 patients who completed four months of treatment (P < 0.001). In addition, for the 11 patients who completed the study the mean skin thickness score also decreased significantly at baseline versus eight months of treatment (P < 0.009). However, the decrease of mean skin thickness score from the fourth to the eighth month of treatment was not significant (P = 0.6).

In general the mean skin thickness score was inversely related to the duration of treatment as shown by regression analysis (r = 0.95, P = 0.00052, fig 1) The distance from the upper to the lower denture borders on full active oral aperture was measured at each visit and increased by the duration of treatment (r = 0.95, P < 0.0005, fig 2). Furthermore, there was a significant inverse correlation between the skin thickness score and the oral aperture (r = 0.92, P < 0.00052). An effort was undertaken to compare the effects of therapy in patients receiving high dose (150 μg m^-2^; n = 6) and low dose (60 μg m^-2^; n = 9) r-IFNy. Patients receiving high dose rIFNy experienced a greater reduction of skin thickness score than patients receiving low dose r-IFNy.

**PULMONARY FUNCTION BEFORE AND AFTER r-IFNy TREATMENT**

Patients 3 and 9 had grade 4 dyspnoea score at study entry, which remained unchanged throughout the study. The remaining patients had a grade 0 dyspnoea score at study entry and of these, patient No 4 developed grade 4 dyspnoea score and patient No 13 grade 1 dyspnoea score at the end of the study. Patients 4, 9, 13, and 15 had a grade 1 interstitial pattern on chest x-ray at study entry, which remained unchanged at the end of the study, while patients 3 and 14 had a grade 2 interstitial pattern which remained also unchanged. Patient No 7 had a grade 4 interstitial pattern at study entry and a grade 2 interstitial pattern at the end of the study. The Po2 values were correlated with the LVEF % predicted values measured by echocardiography (P = 0.028).

Statistical analysis using a paired difference t test on each patient’s forced expiratory volume in one second (FEV1), forced vital capacity (FVC), and DLCO values at baseline versus four, six, and eight months of treatment showed that the changes in these indices over the course of the treatment period were not significant compared to the baseline values. Differences related to dose of r-IFNy were not observed.

**CARDIOLOGICAL EVALUATION BEFORE AND AFTER r-IFNy TREATMENT**

One patient had ventricular premature beats that did not require therapeutic intervention before the institution of r-IFNy treatment. The first injections of r-IFNy were therefore given in an inpatient setting with close monitoring by 24 h Holter; this showed ventricular tachycardia after the third injection of r-IFNy.
Recombinant \( \gamma \) interferon in systemic sclerosis

The treatment was therefore discontinued and the electrocardiographic findings reverted to normal.

Ultrasound examination of the heart revealed a slight pericardial effusion in patient No 11 at study entry. Follow up ultrasound revealed no change in the effusion at the end of the study.

Patient No 5, who discontinued treatment after 17 weeks, presented with a slight pericardial effusion one month earlier, together with the exacerbation of skin thickness. The pericardial effusion in patient No 5 was asymptomatic; the treatment was discontinued because of exacerbation of skin involvement and arthritis. The fluctuations of mean LVEF % predicted values during the study did not reach statistical significance compared to the baseline levels. Differences related to dose of r-IFNy were not observed.

GASTROINTESTINAL INVOLVEMENT IN RESPONSE TO r-IFNy TREATMENT

Nine out of the 15 patients (Nos 2, 3, 4, 9, 10, 11, 12, 14, and 15) experienced severe oesophageal burning due to gastro-oesophageal reflux at study entry and seven of these (Nos 2, 3, 4, 9, 10, 14, and 15) were taking 20 mg of omeprazole daily (alone or with cisapride). Although patients 1 and 8 had a considerable skin involvement response to r-IFNy, they experienced oesophageal burning due to gastro-oesophageal reflux after starting r-IFNy treatment. In addition, patient No 2 developed severe oesophageal ulcers which led to the discontinuation of r-IFNy treatment.

SIDE EFFECTS

Many patients had difficulty tolerating r-IFNy. Fever was the most common side effect (this occurred in 14 patients—87.5%). It was well tolerated by most patients and was minimised by taking 500 mg paracetamol 2 h after the r-IFNy injection. In four of the 14 patients who experienced fever, the symptom resolved two to four months after study entry. In the remaining patients, fever persisted as a side effect, occurring after every r-IFNy injection until the end of the study, but to a lesser degree after two to four months of treatment.

In 10 of the 14 patients who experienced fever, chills also occurred, accompanied by myalgia and fatigue. A severe flu-like syndrome, however, characterised by fever, myalgia, fatigue, chills, and nasal obstruction, occurred in two patients. The intensity of this reaction diminished as treatment progressed and was further reduced by paracetamol.

One male patient experienced severe aphthous stomatitis accompanied by a rise in the ANA titre from 1/320 to 1/1280 with a fine speckled pattern, but no rise anti-dsDNA antibodies. The treatment was discontinued. Prednisolone 10 mg daily was prescribed, together with nystatin by mouth, and the symptom resolved. After one month, injections of r-IFNy were repeated and the same symptom occurred again after the third injection. The patient did not experience arthritis, rash, or renal involvement while his scleroderma was in remission. Thus the severe aphthous stomatitis in this patient should be considered an adverse reaction almost certainly related to r-IFNy.

One female patient (No 16, table 2) with premature ventricular beats before the institution of r-IFNy treatment experienced ventricular tachycardia detected by 24 h Holter monitoring after the injection of r-IFNy and the drug was discontinued.

One patient (No 3) with advanced pulmonary disease at study entry experienced arthralgia after the first injection of r-IFNy. Despite some improvement in skin thickness, her pulmonary function did not improve. Although arthralgia could be considered to be a symptom related to exacerbation of systemic sclerosis, the improvement of skin involvement in this patient indicates that arthralgia occurring shortly after r-IFNy injection was probably an adverse reaction to r-IFNy.

One patient (No 5) developed frank arthritis in wrists, metacarpal and proximal interphalangeal joints, knees, and ankles, as well as doubling of skin thickness score which persisted for more than three months. The same patient developed blurred vision transiently and a non-symptomatic pericardial effusion. Her ANA titre rose from 1/320-fine speckled pattern to 1/1280 with the same pattern, while anti-dsDNA antibodies were not found and the complement levels remained within normal limits.

One patient developed pruritus, hypertrichosis in the shoulders, upper arms and face, and hoarseness. Ultrasonographic examination of the ovaries did not reveal a tumour and computerised tomography of the abdomen did not reveal an adrenal tumour. Plasma testosterone and 17-hydroxyprogesterone levels were within normal limits.

Another patient also developed hoarseness.

REACTIVE OXYGEN SPECIES PRODUCTION FROM PERIPHERAL NEUTROPHILS BEFORE AND AFTER r-IFNy TREATMENT

Neutrophils were isolated from peripheral blood of patients and incubated for 30 min with PMA in the presence of cytochrome C as described in Methods. The production of reactive oxygen species was assessed by measuring...
Table 2  Adverse reactions during eight months of treatment with recombinant interferon-γ

<table>
<thead>
<tr>
<th>Adverse reaction</th>
<th>No of patients (including those who withdrew from the study)</th>
<th>Patients who experienced the adverse reaction (patient No)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fever*</td>
<td>14 1 2 3 4 5</td>
<td>6 7 9 10 11 12 14 15 16</td>
</tr>
<tr>
<td>Chills*</td>
<td>10 1 2 3 4 5</td>
<td>6 7 9 10 11 12 14 15 16</td>
</tr>
<tr>
<td>Headache*</td>
<td>6 1 1 4 5</td>
<td>6 7 9 10 11 12 13</td>
</tr>
<tr>
<td>Severe flu-like symptoms*</td>
<td>2</td>
<td>7 10</td>
</tr>
<tr>
<td>Hoarseness</td>
<td>2</td>
<td>6 8</td>
</tr>
<tr>
<td>Dizziness†</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>Blurred vision†</td>
<td>1</td>
<td>12</td>
</tr>
<tr>
<td>Sever aphthous</td>
<td>1</td>
<td>12</td>
</tr>
<tr>
<td>Somaticatitis§</td>
<td>1</td>
<td>12</td>
</tr>
<tr>
<td>Ventricular tachycardia§</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>Arthralgia†</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>Arthritis†</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>Oesophageal ulcer§</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Itching§</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Hypertrichosis§</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Disease</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>Pericardial effusion§</td>
<td>1</td>
<td>5</td>
</tr>
</tbody>
</table>

Relation to study medication: (∗): almost certain, (†): probable, (§): possible, (§§): reason for withdrawal.

the rate of reduction of cytochrome C according to standard protocols. The mean values (SD) of ROS production rates (nmol per 1.5 million cells min⁻¹) from neutrophils of patients (n = 11) were higher than in normal individuals (n = 12) [8.2 (3.1) v 5.7 (2.1), P < 0.03], while after three months of treatment the mean values decreased significantly [5.1 (2.1) v 8.2 (3.1), P < 0.01] and were comparable to those of normal individuals (fig 3). As shown in fig 3, reactive oxygen species decreased in all the patients except two—one with a mild increase after r-IFNγ treatment and another with a considerable increase. The latter patient (No 5) also experienced arthritis, gastro-oesophageal reflux, and deterioration of scleroderma in the skin. Differences related to dose of r-IFNγ were not observed.

SOLUBLE INTERLEUKIN 2 RECEPTOR CONCENTRATIONS BEFORE AND AFTER r-IFNγ TREATMENT

The SIL-2R serum concentrations were measured at study entry and at the end of the study. Sera from 50 normal individuals were used as controls. Patients 4 and 9 were negative to SIL-2R at study entry. The remaining patients were positive, with values ranging from 500 to 2280 U ml⁻¹ (median 850 U ml⁻¹). Sera from controls gave SIL-2R concentrations ranging from 0 to 168 U ml⁻¹ (median 116 U ml⁻¹). At the end of the study the serum SIL-2R in patients 4 and 9 remained unchanged. Sera from patients 5 and 13 increased and those of the remaining patients decreased. Serum SIL2R after the treatment ranged from 0 to 1550 U ml⁻¹ with a median 350 U ml⁻¹. Comparison between samples on the basis of physical signs indicated that the decrease in SIL-2R after treatment was of marginal significance (P = 0.043). Patients at study entry also had statistically higher SIL-2R concentrations than controls (P < 0.01). Differences related to dose of r-IFNγ were not observed.

Discussion

Our study describes a phase I/II open clinical trial on the toxicity, efficacy, and biological effects of r-IFNγ in the treatment of systemic sclerosis. Several phase I/II trials have been undertaken with the same purpose and have given encouraging results. A significant decrease in skin thickness score was observed at four and eight months of treatment. The decrease in skin thickness correlated with the dose and the duration of treatment with r-IFNγ. A decrease in skin thickness has also been observed in previous studies but no association with the dose and the duration of treatment was seen. Despite the fact that a lower dose of r-IFNγ was used in the current study than in a previous similar study, we observed a greater response in terms of skin...
thickness score. This phenomenon is probably attributed to the fact that the patients in the previous study had more advanced disease in terms of the extent and severity of skin and organ involvement. The ideal therapeutic dose of r-IFNγ remains controversial. In fact doses as low as 50 μg per injection three times weekly and as high as 500 μg m⁻² three times weekly have been tested. The specific activity of rIFNγ preparations used in several studies were not the same; therefore comparisons between the doses should be done with caution. Given that many patients have difficulty in tolerating a dose as high as 500 μg m⁻² per injection, we think that a dose of 150 μg m⁻² or 200 μg m⁻² should be tested in future phase III trials. A second important finding of the current study was that internal organ involvement, in contrast to skin involvement, did not change during the study. Similar findings have also been described by others. Several explanations could be offered for this. (1) The patients who participated in our study had no severe internal organ involvement; therefore the disease process remained unchanged because of endogenous factors and not because of our intervention. (2) The mechanisms of internal organ damage in systemic sclerosis are dissimilar to the mechanisms responsible for skin thickening. (3) Higher doses of rIFNγ may be more effective in treating internal organ involvement. (4) Different organs respond in a different time course. However, a sclerodermal renal crisis has been observed in a patient undergoing treatment with r-IFNγ. This again could be a result of the natural course of disease progression and not due to the intervention. Changes in DLCO and LVEF did not reach statistical significance during the study, as also observed by others, and in contrast to a significant increase in DLCO observed in one study.

Nearly one quarter of our patients met criteria for discontinuation of the drug. Side effects were categorised as almost certain, probably, and possibly related to rIFNγ. Fever and chills, headache, and severe flu-like symptoms were the most common symptoms and were also certainly related to r-IFNγ. Nearly 90% of patients experienced fever, but the intensity of this reaction was ameliorated by paracetamol. Fever could be the result of activation of macrophages by IFNγ, with consequent production of interleukin-1 and hypothalamic stimulation. Alternatively, IFNγ could suppress TH2 lymphocytes which secrete interleukin-10, a potent inhibitor of macrophage activation; therefore, macrophages are activated from products derived from tissue injury in scleroderma, in the absence of downregulatory effects of interleukin-10. Fever, chills, severe flu-like symptoms, and headache were not reasons for discontinuing the treatment. Our patients complained of adverse reactions during the first five months of treatment and they described the greatest response to the therapy during the first two months. The rapid decrease of skin thickness score can be attributed to the decrease of subcutaneous oedema which is not an uncommon manifestation of scleroderma. Other studies have also shown similar results. Arthralgia but not arthritis has been linked with r-IFNγ treatment and occurred in one of our patients who had only a marginal response to r-IFNγ. However, the frank arthritis that occurred in this case cannot be considered a definite side effect of rIFNγ; it may simply have been a manifestation of disease exacerbation.

Exacerbation of systemic sclerosis may be an inevitable result of the disease process itself and not due to our intervention. An effort was made to search for the effects of rIFNγ on biological variables reflecting the presumed underlying mechanism of the drug. We concentrated on two indices known to be associated with the extent and the severity of the disease—the rate of production of reactive oxygen species from peripheral neutrophils and the serum concentrations of SII-2R² in patients before and after r-IFNγ treatment. Enhanced oxidative metabolic activity of neutrophils from patients with systemic sclerosis has been reported¹³ ¹⁴ ²⁷ ²⁸ and increased concentrations of substances such as malondialdehyde, which are produced by oxidative agents, have also been observed in the plasma of patients with systemic sclerosis. Furthermore, an increased susceptibility to oxidation of low density lipoproteins isolated from patients with systemic sclerosis has been described and attributed to high ROS concentrations and to low levels of antioxidants, especially ascorbic acid, in systemic sclerosis patients. Reactive oxygen species can cause several abnormalities related to systemic sclerosis, such as endothelial cell damage, enhanced platelet activation, increased frequency of chromosomal breaks, loss of functional proteinase inhibitors, and oxidation of low density lipoproteins (LDL). Oxidised LDL activate T cells, increase the release of interleukin-1β, induce the expression of genes which encode adhesion molecules in endothelial cells, and increase the expression of platelet derived growth factor from smooth muscle cells. It is therefore plausible that activated neutrophils may represent a pathophysiological mechanism in systemic sclerosis. Neutrophils do not always generate reactive oxygen species. Under the influence of a variety of factors, these cells may increase their capacity of producing ROS after stimulation, a phenomenon called "priming". IFNγ in particular has been regarded generally as an inducer of ROS secretion by neutrophils and this finding encouraged its use in patients with chronic granulomatous disease, with considerable success. However, even in chronic granulomatous disease the clinical response was not in accordance with the changes in superoxide generation from neutrophils after rIFNγ treatment. Moreover, these changes were insignificant. Therefore, knowledge of the previous functional status of neutrophils and their microenvironment is essential in predicting their response to r-IFNγ. Further research is necessary to clarify the exact role of IFNγ on neutrophils in systemic sclerosis patients. We simply wished here to use...
the reduced generation of reactive oxygen species by neutrophils of systemic sclerosis patients as an indicator of some biological action of the drug which might alter the disease process, since large amounts of ROS are found in systemic sclerosis patients compared to normal individuals,13 is in

High levels of SIL-2R were also found at study entry in the sera of all the patients except two. The SIL-2R levels tended to decrease in all patients. The exceptions were patients NO 1 and No 2 who had disease exacerbation, and patient No 4 who had a good response.

In conclusion, our report offers data which suggest that r-IFNγ is relatively safe in the treatment of systemic sclerosis, at least in the dose of 150 μg m⁻² three times weekly and is effective for skin involvement. A decrease in skin thickness score and the expression of side effects from the first five months of treatment could be good indicators for response, non-response, or a tendency to develop side effects.

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