Long-term clinical stabilization of scleroderma patients treated with a chronic and intensive IV iloprost regimen

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Abstract Intravenous iloprost is a first-line option for the treatment of scleroderma-related digital vasculopathy, and some studies have suggested its favourable role on disease progression. The aim of our study is to evaluate the disease progression, specifically in terms of cardiopulmonary function, in a group of consecutive patients chronically treated with intravenous iloprost. Our retrospective study enrolled 68 scleroderma patients (68 F, 54.4 ± 12.3 years) treated with iloprost for 7.1 ± 2.9 years, with a schedule of 5–6 consecutive daily infusions per month (6 h/day, 0.5–2.0 ng/kg/min). In all patients, modified Rodnan skin score (4.7 ± 5.3 vs. 3.7 ± 5.3, p < 0.0001), systolic pulmonary arterial pressure (sPAP) (30.9 ± 6.4 vs. 24.0 ± 3.2 mmHg, p < 0.0001), tricuspid annular plane systolic excursion (22.1 ± 2.4 vs. 23.8 ± 3.5 mm, p = 0.0001), pro-brain natriuretic peptide (97.2 ± 69.3 vs. 65.8 ± 31.7 pg/ml, p = 0.0005) showed statistically significant improvement from baseline. In the subgroup of patients with baseline sPAP ≥36 mmHg (n = 17), a significant sPAP reduction was observed (from 39.5 ± 3.8 to 25.1 ± 4.5 mmHg, p < 0.0001) after 7.6 ± 2.5 years of follow-up. The number of patients with digital ulcers (DUs) at follow-up was reduced from baseline (42.6 vs. 11.8%, p < 0.001), and none of the free-DU patients at baseline presented DUs at follow-up. An intensive and chronic regimen of IV iloprost administration seems to stabilize and potentially improve the long-term development of disease in SSc patients, as suggested by stabilization or significant improvement of cardiopulmonary parameters and vasculopathy.

Keywords Raynaud’s phenomenon · Scleroderma · Iloprost · Systolic pulmonary arterial pressure · Digital ulcers

Introduction

Scleroderma (systemic sclerosis or SSc) is a severe, chronic disease characterized by small vessel vasculopathy, autoantibodies production, and fibroblast dysfunction leading to an excessive deposition of collagen in the skin and internal organs [1, 2].

Severe Raynaud’s phenomenon (RP) is the early onset symptom in most SSc patients and may precede other clinical manifestations of the disease by many years [3]. The clinical course of the disease often involves the cardiovascular and respiratory systems, which can be severely damaged by SSc. The heart can be directly or indirectly involved, with myocardial damage, or with the involvement of other organs, especially kidneys and lungs, respectively [4]. For the respiratory system, SSc can affect lung parenchyma and pulmonary blood vessels, leading to interstitial lung disease (ILD), which may progress to pulmonary arterial hypertension (PAH). The presence of a cardio-pulmonary involvement generally leads to a poor prognosis for the patient [2]. Patients with significant internal organ involvement remain often asymptomatic until the late stages of systemic sclerosis; therefore, routine monitoring...
for the underlying disease and an intensive medical treatment are essential after the first diagnosis. Despite recent advances in the disease management, systemic sclerosis remains a treatable but not curable disease [2].

In the present paper, we report our experience with intravenous iloprost, a stable prostacyclin analogue indicated for the treatment of secondary RP, which may have a favourable effect on the disease progression. Iloprost shows vasodilating, anti-platelet, cytoprotective, and immunomodulating properties, with a long-lasting beneficial effect on the microcirculation [5]. The European League Against Rheumatism (EULAR) guidelines recommend iloprost as a first-line drug for the treatment of SSc-related digital vasculopathy in order to reduce the frequency and severity of SSc-RP attacks and to heal active digital ulcers (DUs) in patients with SSc, representing the gold standard in the treatment of active ulcers [6]. Moreover, recent studies have described a favourable disease course in SSc patients regularly treated with iloprost, with a low occurrence of the most severe vascular complications such as PAH, renal crisis, and digital necrosis [7–12].

The aim of the present study is to evaluate the disease progression, specifically in terms of cardiopulmonary function, in a group of consecutive SSc patients treated with iloprost, at the Unit of Rheumatology of Catania’s hospital, Italy.

**Methods**

Our retrospective study enrolled 68 SSc patients (68 F, 54.4 ± 12.3 years) treated with iloprost for 7.1 ± 2.9 years, with a schedule of 5–6 consecutive daily infusions per month (6 h/day, 0.5–2.0 ng/kg/min). Patients generally received six iloprost infusions per month, with possible suspensions or reductions during the warmest month of the year, for the treatment of secondary RP. The study was conducted under the Declaration of Helsinki and the current ethical standards. Assessed parameters included the following: modified Rodnan skin score (mRss), systolic pulmonary arterial pressure (sPAP), tricuspid annular plane systolic excursion (TAPSE), diffusing capacity of the lung for carbon monoxide (DLCO), forced vital capacity (FVC), alveolar volume (AV), DLCO/AV, pro-brain natriuretic peptide (pBNP), NYHA class, and presence/absence of digital ulcers.

Descriptive statistics was performed by calculating mean and standard deviation (SD) for continuous variables, and percentage for discrete variables. Statistical comparisons of post-baseline versus baseline data were made using Student’s t test for paired data when continuous variables were analysed, and by applying the McNemar test, when categorical variables were considered. Statistical analyses were performed through standard procedures using SAS statistical program, version 9.2 (SAS Institute, Cary, NC, USA).

**Results**

The characteristics of the study population are described in Table 1. Patients were followed up for an average of 7.1 years, varying from 1 to 15 years, during which they were regularly treated with intravenous iloprost. The age of onset of Raynaud’s phenomenon corresponds in most cases to the onset of the disease and the diagnosis of scleroderma. The evaluations refer to baseline data—before the beginning of treatment with iloprost—and to the last available follow-up data for each patient.

The results show a stabilization of the cardiopulmonary function, since these parameters remained unchanged or were significantly improved (Table 2).

A significant sPAP reduction from 39.5 ± 3.8 to 25.1 ± 4.5 mmHg (p < 0.0001) was also observed in the
Discussion

The present results show a disease stabilization in SSc patients during a long-term follow-up, as indicated by the improvement or non-worsening of the assessed parameters.

This patient’s cohort showed a low mRSS at baseline with a slight but significant reduction during the follow-up. This finding might be relevant, since previous studies showed a skin score worsening related to the disease progression in terms of patient survival [13] or occurrence of scleroderma renal crisis [14].

Our results indicate a stabilization of the cardiopulmonary disease, in a very long-term follow-up, possibly suggesting a favourable effect of the treatment against the development of pulmonary arterial hypertension. These conclusions are first supported by NYHA class non-progression and a significant reduction of systolic pulmonary arterial pressure and brain natriuretic peptide levels.

In particular, the importance of sPAP has been recently focused by the EUSTAR working group, since baseline values ≥36 mmHg were significantly associated with an increased risk of death up to 3-year follow-up [HR 1.44 (1.06, 1.96) vs. baseline sPAP < 36 mmHg], regardless of the presence of pulmonary hypertension assessed by right heart catheterization [15]. In our study, average baseline sPAP of the entire population was 30.9 ± 6.4 and 24.0 ± 3.2 mmHg after 7.1 ± 2.9 years of follow-up. Interestingly, in the subgroup of patients with baseline sPAP ≥36 mmHg (n = 17), a significant reduction of the sPAP was observed (from 39.5 ± 3.8 to 25.1 ± 4.5 mmHg) after an average follow-up of 7.6 ± 2.5 years, and no deaths were recorded.

In the absence of control groups that did not receive iloprost, we compared our data with those reported in recent literature. A recent study by D’Alto et al. [16] reported a significant worsening of sPAP (from 26.1 ± 6.0 to 28.8 ± 6.3) after a 3-year follow-up in a group of 74 consecutive SSc patients, in which IV iloprost was administered in 19% of cases. In the EUSTAR cohorts, where the treatment with IV iloprost is reported in 15.6% of cases, overall survival at 5 years is generally comprised between 80 and 90% [15, 18, 19]. Therefore, our results can be considered encouraging.

Table 2 Changes in skin score and cardiopulmonary function over time

|                                | All patients (n = 68) | Baseline | Follow-up | p    |
|--------------------------------|----------------------|----------|-----------|------|
| mRSS ± SD                      | 4.7 ± 5.3            | 3.7 ± 5.3|           | <0.0001|
| sPAP ± SD, mmHg                | 30.9 ± 6.4           | 24.0 ± 3.2|           | <0.0001|
| NYHA class ± SD               | 1.0 ± 0.0            | 1.0 ± 0.4|           | 1.0   |
| TAPSE ± SD, mm                | 22.1 ± 2.4           | 23.8 ± 3.5|           | 0.0001|
| FVC ± SD, % predicted          | 107.1 ± 14.5         | 101.2 ± 21.3|       | 0.0581|
| DLCO ± SD, % predicted         | 83.7 ± 13.5          | 81.4 ± 14.3|           | 0.4121|
| AV ± SD, % predicted           | 91.1 ± 13.0          | 91.3 ± 15.3|           | 0.9855|
| DLCO/AV ± SD, % predicted      | 88.5 ± 13.5          | 91.2 ± 14.0|           | 0.0575|
| pBNP ± SD, pg/ml              | 97.2 ± 69.3          | 65.8 ± 31.7|           | 0.0005|

AV alveolar volume, DLCO diffusing capacity of the lung for carbon monoxide, FVC forced vital capacity, mRSS modified Rodnan skin score, pBNP pro-brain natriuretic peptide, SD standard deviation, sPAP systolic pulmonary arterial pressure, TAPSE tricuspid annular plane systolic excursion

The subgroup of patients with baseline sPAP ≥36 mmHg (n = 17), after an average follow-up of 7.6 ± 2.5 years.

In all patients, mRSS (4.7 ± 5.3 vs. 3.7 ± 5.3, p < 0.0001), tricuspid annular plane systolic excursion (22.1 ± 2.4 vs. 23.8 ± 3.5 mm, p = 0.0001), pro-brain natriuretic peptide (97.2 ± 69.3 vs. 65.8 ± 31.7 pg/ml, p = 0.0005) showed a statistically significant improvement from baseline. DLCO, AV, DLCO/VA, FVC, and NYHA class remained unchanged.

Table 3 Presence of digital ulcers in the study population

| Presence of digital ulcers | Baseline (%) | Follow-up (%) |
|----------------------------|--------------|--------------|
| All patients (n = 68, follow-up 7.1 ± 2.9 years) | 42.6 | 11.8 |
| Patients according to treatment | | |
| Iloprost (n = 35, follow-up 5.8 ± 2.7 years) | 5.7 | 0.0 |
| Iloprost + bosentan (n = 33, follow-up 8.3 ± 2.5 years) | 81.8 | 24.2 |
We also observed a significant change in brain natriuretic peptide levels, which remained below the upper normal limit of 125 pg/ml according to our clinical practice. BNP is released from myocardium in response to wall stress and induces vasodilatation and natriuresis, and high BNP levels are found in patients with cardiac infarction, congestive heart failure, and pulmonary hypertension. High BNP concentration was found to proportionally increase with the degree of right ventricular dysfunction, and further increases are associated with mortality in established severe primary PAH [20]. Thus, BNP levels represent an important diagnostic marker of early pulmonary artery hypertension in SSc patients [20].

Mean TAPSE values remained >20 mm during the whole study period, with significant improvement from 22.1 ± 2.4 to 23.8 ± 3.5 mm. According to the current guidelines for the diagnosis and treatment of pulmonary hypertension, TAPSE has a well-established importance for assessing disease severity, stability, and prognosis in PAH patients, with a cut-off value > 20 mm indicating a satisfactory patient status [21]. The importance of this parameter is related to the multiple components of right ventricular function, especially with right ventricular ejection fraction (RVEF), in patients with SSc-PAH. In SSc patients, TAPSE values lower than 19.6 mm suggest a RVEF <40%; the lower is the value, the higher is the frequency of hospitalization [22]. Raw mortality data were also found to be significantly associated with TAPSE in SSc patients: values ≤17 mm prognosticated a nearly fourfold increased risk of death compared to patients with TAPSE > 17 mm. Moreover, when calculated as a continuous variable, a decrease of 1 mm in TAPSE was associated with a 15% increased risk of death [23].

We also observed a stabilization of interstitial lung disease markers, such as DLCO, FVC, VA, and DLCO/VA. This is important for both patient’s functionality and prognosis.

Also, our results have a relevant impact on patient’s quality of life since DUs represent a frequent and severe source of pain and disability in SSc. The administration of an intensive IV iloprost regimen, in combination with bosentan when indicated, improved pre-existing ulcer healing and prevented the occurrence of new DUs, confirming the importance of these two therapeutic options in the management of SSc vasculopathy [6]. Moreover, these data provide useful information for the chronic administration of IV iloprost, since both EULAR [6] and manufacturer’s [24] recommendations do not provide satisfactory instructions for the therapy repetition over time, probably because all main randomized controlled trials have been carried out with a single infusion cycle and a short-term follow-up [25, 26]. Thus, since the administration of a single 5-day IV iloprost infusion cycle was already known to improve RP and heal DUs, the present results indicate that its monthly repetition maintains a favourable healing effect, and reduce the occurrence of new DUs over time.

The present study has some limitations mainly due to the trial design. Being a retrospective analysis of a patient’s database, a control group was not provided, and the number of patients was relatively limited. Thus, our results need to be confirmed in further prospective and possibly controlled trials.

Conclusions

Scleroderma remains a treatable but not curable disease, characterized by a poor prognosis due to the occurrence of cardiopulmonary complications. Therefore, the long-term disease stabilization represents an important therapeutic goal. Intravenous iloprost may play a role in promoting a favourable disease course during a long-term follow-up. Our results confirm that monthly iloprost infusions for six consecutive days might exert a beneficial healing effect and reduce the occurrence of new DUs as well against cardiopulmonary disease development or worsening in SSc patients.

Compliance with ethical standards

Conflict of interest Alberto Farina is an employee of Italfarmaco S.p.A., and the other authors report no conflict of interest.

Ethical approval All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki Declaration and its later amendments or comparable ethical standards.

Informed consent Informed consent was obtained from all individual participants included in the study.

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References

1. Gabrielli A, Avvedimento EV, Krieg T (2009) Scleroderma. N Engl J Med 360:1989–2003
2. Hinchcliff M, Varga J (2008) Systemic sclerosis/scleroderma: a treatable multisystem disease. Am Fam Physician 78:961–968
3. Herrick AL (2000) Vascular function in systemic sclerosis. Curr Opin Rheumatol 12:527–533
4. Champion HC (2008) The heart in scleroderma. Rheum Dis Clin North Am 34:181–190
5. Scorza R, Caronni M, Mascagni B, Berruti V, Bazzi S, Micallef E, Arpaia G, Sardina M, Origgio L, Vanoli M (2001) Effects of long-term cyclic iloprost therapy in systemic sclerosis with Raynaud’s phenomenon. A randomized, controlled study. Clin Exp Rheumatol 19:503–508
6. Kowal-Bielecka O, Landewé R, Avouac J, Chwieski S, Martin RW, Collier D, Weinstein A, Hurwitz EL, Mayes M, White B, Wigley F, Jimenez S, Mayes M, Cleland PJ, Weiner SR, Porter J, Ellman M, Wise C, Kaufman LS (1994) Intravenous iloprost treatment of Raynaud’s phenomenon. A multicenter, placebo-controlled, double-blind study. Clin Exp Immunol 95:44–50
7. Caramaschi P, Volpe A, Tinazzi I, Barbarea LM, Carletto A, Biasi D (2006) Does cyclically iloprost infusion prevent severe isolated pulmonary hypertension in systemic sclerosis? Preliminary results. Rheumatol Int 27:203–205
8. Caramaschi P, Martinelli N, Volpe A, Piropan S, Tinazzi I, Patuzzo G, Mahamid H, Barbarea LM, Biasi D (2009) A score of risk factors associated with ischemic digital ulcers in patients affected by systemic sclerosis treated with iloprost. Clin Rheumatol 28:807–813
9. Caramaschi P, Dalla Gassa A, Prati D, Barausse G, Tinazzi I, Ravagnani V, Contente S, Biasi D (2012) Severe vascular complications in patients affected by systemic sclerosis cyclically treated with iloprost. Rheumatol Int 32:1933–1938
10. Caravita S, Wu SC, Secchi MB, Dadone V, Bencini C, Pierini P, Rosato E, Carreira PE, Riccieri V, Sarraco M, Denton CP, Riemekasten G, Valentini S, Caramaschi P, Martinelli N, Volpe A, Bazzi S, Micallef E, Arpaia G, Sardina M, Origgio L, Vanoli M (2001) Improvement in skin thickness as a prognostic factor of death in the systemic sclerosis sclerodema trial. Arthritis Rheum 46:2983–2989
11. Steen VD, Medsger TA Jr, Petri M, Taylor NJ, Sierakowski S, Allanore Y, Czirjak L, Hachulla E, Riemekasten G, Farge D, Müller-Ladner U, Matucci-Cerinic M, Földvari I, Furst DE, Müller-Ladner U, Seibold J, Silver RM, Ullman S, Zaiman AL, Girgis RE, Hassoun PM (2011) Tricuspid annular plane systolic excursion. Echocardiography 24:118–125
12. DeMarco PJ, Weisman MH, Seibold JR, Furst DE, Wong WK, Hurwitz EL, Mayes M, White B, Wigley F, Barr W, Moreland L, Medsger TA Jr, Stein V, Martin RW, Collier D, Weinstein A, Lally E, Varga J, Weiner SR, Andrews B, Abeles M, Caramaschi P, Volpe A, Piropan S, Tinazzi I, Patuzzo G, Mahamid H, Barbarea LM, Biasi D (2002) Predictors and outcomes of scleroderma renal crisis: the high-dose versus low-dose n-penicillamine in early diffuse systemic sclerosis trial. Arthritis Rheum 46:2983–2989
13. Hachulla E, Clerson P, Airò P, Cuomo G, Allanoire Y, Caramaschi P, Rosato E, Carreira PE, Riccieri V, Sarraco M, Denton CP, Riemekasten G, Pozzi MR, Zini S, Mihai CM, Ullman S, Distler O, Rednic S, Smith V, Walker UA, Matucci-Cerinic M, Müller-Ladner U, Launay D (2015) Value of systolic pulmonary arterial pressure as a prognostic factor of death in the systemic sclerosis EUSTAR population. Rheumatology 54:1262–1269
14. D’Alto M, Cuomo G, Romeo E, Argiento P, Iudici M, Vettori S, Giovanna Russo M, Calbrò R, Valentini G (2014) Tissue Doppler imaging in systemic sclerosis: a 3-year longitudinal study. Semin Arthritis Rheum 43:673–680
15. Meier FM, Frommer KW, Dinser R, Walker UA, Czirjak L, Denton CP, Allanoire Y, Distler O, Riemekasten G, Valentini G, Müller-Ladner U (2012) Update on the profile of the EULAR Scleroderma Trials and Research group database. Ann Rheum Dis 71:1335–1360
16. Fransen J, Popa-Diaconu D, Hesselstrand R, Carreira P, Valentini G, Beretta L, Airo P, Inanc M, Ulman S, Balbir-Gurman A, Sierakowski S, Allanoire Y, Czirjak L, Riccieri V, Giacomelli R, Gabrielli A, Riemekasten G, Matucci-Cerinic M, Farge D, Hunzelmann N, Van den Hoogen FH, Vonk MC (2011) Clinical prediction of 5-year survival in systemic sclerosis: validation of a simple prognostic model in EUSTAR centres. Ann Rheum Dis 70:1788–1792
17. Mihai C, Landewé R, van der Heijde D, Walker UA, Constantin PL, Gherge AM, Ionescu R, Rednic S, Allanoire Y, Avouac J, Czirjak L, Hachulla E, Riemekasten G, Cozzi F, Airò P, Cutolo M, Mueller-Ladner U, Matucci-Cerinic M (2016) Digital ulcers predict a worse disease course in patients with systemic sclerosis. Ann Rheum Dis 75:681–686
18. Allanoire Y, Borderie D, Meune C, Cabanes L, Weber S, Ekindjian OG, Kahan A (2003) N-terminal pro-brain natriuretic peptide as a diagnostic marker of early pulmonary artery hypertension in patients with systemic sclerosis and effects of calcium-channel blockers. Arthritis Rheum 48:3503–3508
19. Galí N, Hoepfer MM, Humbert M, Torbicki A, Vachiery JL, Barbera JA, Beghetti M, Corris P, Gaine S, Gibbs JS, González-Sanchez MA, Jondeau G, Klepetko W, Optiz C, Peacock A, Rubin L, Zellweger M, Simonneau G, ESC Committee for Practice Guidelines (CPG) (2009) Guidelines for the diagnosis and treatment of pulmonary hypertension: the Task Force for the Diagnosis and Treatment of Pulmonary Hypertension of the European Society of Cardiology (ESC) and the European Respiratory Society (ERS), endorsed by the International Society of Heart and Lung Transplantation (ISHLT). Eur Heart J 30:2493–2537
20. Lee CY, Chang SM, Hsiao SH, Tseng JC, Lin SK, Liu CP (2007) Right heart function and scleroderma: insights from tricuspid annular plane systolic excursion. Echocardiography 24:118–125
21. Mathai SC, Sibley CT, Forfia PR, Mudd JO, Fisher MR, Tedford RJ, Lechtzin N, Boyce D, Hummers LK, Housten T, Zaiman AL, Girgis RE, Hassoun PM (2011) Tricuspid annular plane systolic excursion is a robust outcome measure in systemic sclerosis-associated pulmonary arterial hypertension. J Rheumatol 38:2410–2418
22. Italfarmaco S.p.A. Endoprost (2016) Iloprost trometamol solution for infusion [Package Insert]. Italfarmaco S.p.A, Milano
23. Wigley FM, Seibold JR, Wise RA, McCloskey DA, Dole WP (1994) Intravenous iloprost infusion in connective tissue disease. Eur J Intern Med 22:518–521
24. Mathai SC, Sibley CT, Forfia PR, Mudd JO, Fisher MR, Tedford RJ, Lechtzin N, Boyce D, Hummers LK, Housten T, Zaiman AL, Girgis RE, Hassoun PM (2011) Tricuspid annular plane systolic excursion. Echocardiography 24:118–125
25. Wigley FM, Seibold JR, Wise RA, McCloskey DA, Dole WP (1994) Intravenous iloprost infusion in connective tissue disease. Eur J Intern Med 22:518–521
26. Mathai SC, Sibley CT, Forfia PR, Mudd JO, Fisher MR, Tedford RJ, Lechtzin N, Boyce D, Hummers LK, Housten T, Zaiman AL, Girgis RE, Hassoun PM (2011) Tricuspid annular plane systolic excursion. Echocardiography 24:118–125
27. Mathai SC, Sibley CT, Forfia PR, Mudd JO, Fisher MR, Tedford RJ, Lechtzin N, Boyce D, Hummers LK, Housten T, Zaiman AL, Girgis RE, Hassoun PM (2011) Tricuspid annular plane systolic excursion is a robust outcome measure in systemic sclerosis-associated pulmonary arterial hypertension. J Rheumatol 38:2410–2418
28. Italfarmaco S.p.A. Endoprost (2016) Iloprost trometamol solution for infusion [Package Insert]. Italfarmaco S.p.A, Milano
29. Wigley FM, Seibold JR, Wise RA, McCloskey DA, Dole WP (1992) Intravenous iloprost treatment of Raynaud’s phenomenon and ischemic ulcers secondary to systemic sclerosis. J Rheumatol 19:1407–1414
30. Wigley FM, Seibold JR, Wise RA, McCloskey DA, Dole WP (1992) Intravenous iloprost treatment of Raynaud’s phenomenon and ischemic ulcers secondary to systemic sclerosis. J Rheumatol 19:1407–1414