Longterm Effects of Endothelin Receptor Antagonism on Microvascular Damage Evaluated by Nailfold Capillaroscopic Analysis in Systemic Sclerosis

MAURIZIO CUTOLO, GIUSEPPE ZAMPOGNA, LAURA VREMIS, VANESSA SMITH, CARMEN PIZZORNI, and ALBERTO SULLI

ABSTRACT. Objective. Systemic sclerosis (SSc) is characterized by microvascular injury, fibrosis, and hypoxia of involved tissues. The vasoactive peptide endothelin-1 (ET-1) seems to be implicated in these events. Using nailfold videocapillaroscopy (NVC), we evaluated longterm effects of ET-1 antagonist treatment on nailfold microvascular damage in patients with SSc, over a 3-year followup period.

Methods. Thirty patients with SSc (mean age 64 ± 5 yrs, mean disease duration 8 ± 1 yrs) were recruited during their programmed standard treatment protocols. At baseline (T0), 15 patients with SSc (mean age 63 ± 15 yrs, mean disease duration 7 ± 3 yrs), already receiving cyclic intravenous infusion of iloprost (5 continuous days, average 80 µg/day, every 3 mo), continued the treatment for a further 3 years (ILO group). The remaining 15 patients with SSc (mean age 68 ± 13 yrs, mean disease duration 8 ± 4 yrs), although they continued the same cyclic intravenous iloprost treatment as the previous group, also received bosentan 125 mg twice a day for 3 years (ILO+BOS group). Qualitative analysis (scleroderma patterns) and semiquantitative scoring of the microvascular damage were performed by validated routine NVC methods.

Results. During followup, a statistically significant increase of capillary number was observed in the ILO+BOS group (p < 0.02), with a significant and progressive increase of angiogenesis (p < 0.01). In contrast, the ILO group showed a statistically significant decrease of capillary number (p < 0.05). After 3 years the number of capillaries was significantly higher in the ILO+BOS group than in the ILO group (p < 0.05). The score for giant capillaries decreased significantly in both groups of patients with SSc (p < 0.05).

Conclusion. In this open study, longterm treatment with ET-1 receptor antagonist in combination with iloprost was found to interfere with progression of nailfold microvascular damage in patients with SSc, as assessed by NVC over a 3-year followup period. (First Release Nov 1 2012; J Rheumatol 2013;40:40–5; doi:10.3899/jrheum.120416)

Key Indexing Terms:
NAILFOLD VIDEOCAPILLAROSCOPY   SYSTEMIC SCLEROSIS   MICROANGIOPATHY   ENDOTHELIN-1   BOSENTAN

Systemic sclerosis (SSc) is characterized mainly by microvascular damage, fibrosis, and hypoxia of involved tissues causing progressive organ failure1. SSc microangiopathy is a dynamic and sequential process, progressing from early enlargement of capillaries (giant capillaries) to capillary loss, tissue ischemia, and finally tissue fibrosis and reactive neoangiogenesis (capillary ramifications)2.

Currently, the structural capillary changes are investigated by nailfold videocapillaroscopy (NVC) and 3 different and progressive patterns of nailfold microvascular damage have been described in SSc, i.e., the early, active, and late scleroderma patterns3.

Briefly, NVC allows the investigator to carry out early diagnosis of SSc, and to characterize, quantify, and monitor the progression of nailfold microvascular damage4. Therefore, SSc microangiopathy was found to be predictive for progression of primary Raynaud’s phenomenon (RP) to SSc and related clinical complications such as digital ulcers5,6,7.

There are several therapeutic approaches to improve RP or to reduce the appearance of new digital ulcers8. Iloprost is a prostacyclin analog that blocks platelet aggregation and adhesion, dilates arterioles and venules, activates fibrinolysis, and reduces the release of oxygen-reactive species,
reducing the effects of RP9. Endothelin-1 (ET-1), a 21-amino acid peptide, is a potent vasoconstrictor, mainly produced by endothelial cells, that is found in high concentrations in the skin as well as in lung, kidney, and plasma of patients with SSc10,11. Bosentan, a dual ET-1 receptor antagonist, has been licensed to treat pulmonary hypertension and to prevent the onset of new digital ulcers in patients with SSc and history of digital ulcers8.

This investigation was planned to evaluate by NVC the longterm effects (3 years) of bosentan treatment on progression of nailfold microvascular damage in patients with SSc who were already receiving cyclic treatment with intravenous iloprost.

**MATERIALS AND METHODS**

*Patients.* Thirty patients (mean age 64 ± 5 years) with SSc (mean disease duration 8 ± 1 years) according to the American College of Rheumatology criteria for SSc12 or the LeRoy criteria for the classification of early SSc13, and who were attending the Academic Unit of Clinical Rheumatology at the University of Genoa during 2007-2008, were followed during their standard treatment protocols. All patients provided informed consent.

A complete medical history and comprehensive laboratory examination results were available for all patients, including results of a full medical assessment. All patients were regularly taking aspirin (100–150 mg/day) and buflomedil (300–600 mg/day), a vasoactive drug mainly inhibiting adrenoreceptors and platelet aggregation. In addition, all patients were treated with intravenous iloprost in cycles of 5 days, by continuous 24-h intravenous infusion (median 80 µg/day) every 3 months, with temporary withdrawal of buflomedil.

At study baseline (T0), 15 patients (mean age 63 ± 15 years, mean disease duration 7 ± 3 years) already receiving cyclic intravenous iloprost for an average of 5 ± 1 years continued such treatment for 3 additional years (ILO group). The remaining 15 patients (mean age 68 ± 13 years, mean disease duration 8 ± 4 years), although they continued the same cyclic intravenous iloprost treatment as the previous group (for an average time of 7 ± 3 years), also received bosentan, according to the general recommendations, since they experienced the appearance of new digital ulcers (ILO+BOS group). Bosentan was administered at a dosage of 62.5 mg twice a day orally for 1 month; after liver function assessment, the dosage was increased to 125 mg twice a day. All patients with SSc were followed for at least 3 years and were assessed yearly by NVC (i.e., T0, T1, T2, T3). Liver function and other serum measurements were monitored every 4–6 months. Patients' demographic data are reported in Table 1.

**Nailfold videocapillaroscopy.** NVC was performed yearly in each patient using an optical probe videocapillaroscope equipped with a 200× contact lens and connected to image analysis software (Videocap, DS MediGroup). The same operator (CP), blinded to patients' clinical data, performed all NVC examinations and scored all images. Each patient was inside the building for a minimum of 15 min before the nailfold was examined and the room temperature was 22°–23°C. Nailfolds of second, third, fourth, and fifth fingers of both hands (sparing the thumbs) were examined in each patient, after a drop of immersion oil was placed on the nailfold bed to improve image resolution14. The first line of capillaries at the nailfold was evaluated, and the following capillaroscopic variables were considered, based on our previous studies: number of capillaries, presence of giant capillaries, microhemorrhages, capillary ramifications (reactive angiogenesis), and disorganization2,3,4.

A semiquantitative and validated rating scale to score each capillary abnormality was adopted, in accord with previous studies (0–3 score for each variable: 0 = no changes, 1 = less than 33% capillary alterations/reduction, 2 = 33%–66% of capillary alterations/reduction, 3 = greater than 66% capillary alterations/reduction, per linear mm)5,16.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>ILO Group (15 patients)</th>
<th>ILO+BOS Group (15 patients)</th>
</tr>
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<tbody>
<tr>
<td>Mean age, yrs</td>
<td>63 ± 15</td>
<td>68 ± 13</td>
</tr>
<tr>
<td>Mean SSc duration, yrs</td>
<td>7 ± 3</td>
<td>8 ± 4</td>
</tr>
<tr>
<td>Sex (male/female)</td>
<td>1/14</td>
<td>2/13</td>
</tr>
<tr>
<td>LSSc, n</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>LeSSc, n</td>
<td>9</td>
<td>7</td>
</tr>
<tr>
<td>DeSSc, n</td>
<td>3</td>
<td>8</td>
</tr>
<tr>
<td>ANA speckled pattern, %</td>
<td>13</td>
<td>27</td>
</tr>
<tr>
<td>ANA homogeneous, %</td>
<td>13</td>
<td>40</td>
</tr>
<tr>
<td>ANA nucleolar, %</td>
<td>7</td>
<td>7</td>
</tr>
<tr>
<td>ANA centromeric, %</td>
<td>67</td>
<td>27</td>
</tr>
<tr>
<td>Sc70-positive, %</td>
<td>20</td>
<td>27</td>
</tr>
<tr>
<td>New digital ulcers, n</td>
<td>T0 0/15</td>
<td>15/15*</td>
</tr>
<tr>
<td>T3</td>
<td>2/15</td>
<td>0/15</td>
</tr>
</tbody>
</table>

* Patients were enrolled in the ILO+BOS group upon the appearance of new digital ulcers. LSSc: limited systemic sclerosis; LeSSc: limited cutaneous systemic sclerosis; DeSSc: diffuse cutaneous systemic sclerosis; ILO: iloprost; BOS: bosentan; ANA: antinuclear antibodies.

In relation to nailfold capillary abnormalities, patients were distributed within the appropriate NVC pattern of microangiopathy (i.e., early, active, and late scleroderma patterns), as described2,3. The “microangiopathy evolution score” (MES; the sum of 3 scores: loss of capillaries, disorganization of the microvascular array, and capillary ramifications) was also adopted to globally assess the progression of vascular damage, as reported15,17.

**RESULTS**

**Baseline NVC findings and pattern transition.** No statistically significant difference was observed between the 2 groups of patients concerning the scores for different NVC measures at the baseline visit (Table 2).

At baseline (T0), in the ILO group the late NVC pattern was present in 5 of 15 patients (33%), the active pattern in 7 (47%), and the early pattern in 3 (20%), while in the ILO+BOS group, 11 of 15 patients (73%) showed the late NVC pattern and 4 (26%) the active pattern of microangiopathy.

**Progression of nailfold microvascular damage.** At the end of the followup (T3), in the ILO group 1 patient with the early and 3 patients with the active pattern at baseline shifted to the late microangiopathy pattern. Conversely, in the ILO+BOS group, 1 patient shifted from the late to the active pattern of microangiopathy.

Statistically significant modifications of the scores for the capillaroscopic measures were observed in the 2 groups of patients with SSc, as described below.

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Followup are reported in Table 2. The microvascular damage, along with a greater amount of microhemorrhages, and capillary ramifications) during the ILO group. These results suggest a reduced progression of reactive angiogenesis.

Combined treatment with iloprost and bosentan. A statistically significant progressive increase of capillary number with a significant and progressive increase of capillary was observed in the ILO+BOS group of patients, together with a significant and progressive increase of capillary ramifications (Figure 1). Interestingly, the increase of capillary ramifications was greater than that observed in the ILO group. These results suggest a reduced progression of the microvascular damage, along with a greater amount of reactive angiogenesis.

The detailed variations of scores for the single NVC morphological markers (capillary loss, giant capillaries, microhemorrhages, and capillary ramifications) during the followup are reported in Table 2.

Of note, after 3 years (T3), the number of capillaries was significantly higher in the ILO+BOS group than in the ILO group and the difference was statistically significant (Figure 1). Giant capillaries decreased significantly from T1 to T3 (Figure 1).

No relevant side effects leading to withdrawal of treatment and no new digital ulcers were observed in the ILO+BOS patient group at the end of the followup. Minor side effects such as nausea and mild increases of transaminase were managed by temporary reduction of dosages.

DISCUSSION

Our study reports the longterm effects of ET-1 receptor antagonism on nailfold microvascular damage changes, as assessed by NVC analysis in patients with SSc who had digital ulcers. In particular, NVC revealed that treatment with the combination of bosentan with iloprost significantly increased the number of capillaries after 2 years of treatment; in contrast, a progressive and significant loss of capillaries, as expected in the natural followup of the disease, was observed under iloprost treatment alone. These data are intriguing, but should be interpreted with caution because this was a nonrandomized, unblinded, uncontrolled trial, with 2 groups of patients with dissimilar baseline characteristics.

Dynamic modifications of the nailfold microvascular bed during the course of SSc have been demonstrated by a 20-year longitudinal study, and progressive transition through NVC patterns has been linked to the progression of clinical complications in SSc. However, the therapeutic influence on transition of the NVC pattern has rarely been evaluated in SSc, excluding the effects of immunosuppressive treatments (i.e., cyclosporine, cyclophosphamide) that showed a regression from active/late compared to the early/active NVC patterns, respectively.

Besides being a potent vasoconstrictor, ET-1 is a direct downstream target of transforming growth factor-β1 (TGF-β1) and it behaves as a profibrotic cytokine, stimu-

<table>
<thead>
<tr>
<th>Time, yrs</th>
<th>T0</th>
<th>T1</th>
<th>T2</th>
<th>T3</th>
<th>p (T0 vs T3)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ILO group</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Loss of capillaries</td>
<td>1.71 ± 0.70</td>
<td>1.86 ± 0.52</td>
<td>1.93 ± 0.46</td>
<td>2.00 ± 0.38</td>
<td>0.05</td>
</tr>
<tr>
<td>Giant capillaries</td>
<td>1.14 ± 0.83</td>
<td>1.29 ± 0.59</td>
<td>1.14 ± 0.64</td>
<td>0.93 ± 0.59</td>
<td>0.05</td>
</tr>
<tr>
<td>Microhemorrhages</td>
<td>0.86 ± 0.52</td>
<td>0.93 ± 0.46</td>
<td>0.93 ± 0.46</td>
<td>0.86 ± 0.52</td>
<td>NS</td>
</tr>
<tr>
<td>Ramifications</td>
<td>1.50 ± 0.91</td>
<td>1.64 ± 0.72</td>
<td>1.79 ± 0.56</td>
<td>1.71 ± 0.88</td>
<td>0.05</td>
</tr>
<tr>
<td>ILO+BOS group</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Loss of capillaries</td>
<td>1.79 ± 0.89</td>
<td>1.79 ± 0.89</td>
<td>1.71 ± 0.88</td>
<td>1.43 ± 0.65</td>
<td>0.02</td>
</tr>
<tr>
<td>Giant capillaries</td>
<td>0.93 ± 0.73</td>
<td>1.00 ± 0.78</td>
<td>1.00 ± 0.78</td>
<td>0.71 ± 0.83</td>
<td>NS</td>
</tr>
<tr>
<td>Microhemorrhages</td>
<td>0.57 ± 0.65</td>
<td>0.50 ± 0.65</td>
<td>0.43 ± 0.65</td>
<td>0.36 ± 0.63</td>
<td>NS</td>
</tr>
<tr>
<td>Ramifications</td>
<td>1.21 ± 0.80</td>
<td>1.43 ± 0.51</td>
<td>1.79 ± 0.70</td>
<td>2.07 ± 0.83</td>
<td>0.003</td>
</tr>
</tbody>
</table>

ILO: iloprost; BOS: bosentan; T0: baseline; T1: 1 year; T2: 2 years; T3: 3 years; NS: nonsignificant.
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The significant increase of normal capillaries observed during longterm ET-1 receptor antagonism in SSc patients with digital ulcers, as assessed by NVC, suggests important interference exerted on the TGF-β1/ET-1 axis, particularly through the mechanisms involved in the endothelial-to-mesenchymal cell transition (EndMT)\textsuperscript{27}. The importance of the EndMT as one of the leading processes in fibrosis, including a role for ET-1, has recently been supported, because diabetes-induced cardiac fibrosis seems to be associated with the emergence of fibroblasts from endothelial cells, and this transition process is stimulated by ET-1\textsuperscript{28}.

Thus, the results of our study might suggest that ET-1 receptor antagonism exerted over the long term in patients with SSc may interfere with the mechanisms inducing progressive microvascular damage such as capillary loss.
and consequent tissue hypoxia and fibrosis. In addition, the larger extent of angiogenesis observed in SSc patients treated with bosentan might be explained as an attempt to revascularize the tissues under hypoxia. It is suggested that elevated angiogenic serum markers, as reported in patients with SSc, may appear as a compensatory response to maintain vascular function with the formation of new vessels, which might lead to enhanced capillary formation.29

However, angiogenesis resulting from defective vascularization and tissue hypoxia is insufficient in SSc, despite elevated levels of proangiogenic factors (i.e., vascular endothelial growth factor and circulating endothelial progenitor cells).30,31 Therefore, the reasons that a significant increase of angiogenesis together with increased capillary formation was observed by NVC in SSc patients treated with bosentan are not clear at present, but deserve further investigation.

Nevertheless, the reversal of progressive loss of capillaries by ET-1 receptor blocker treatment needs to be interpreted with caution. First, our study was observational, and not a randomized controlled trial; second, because the ET-1 receptor blocker therapy was added upon clinical indication (the appearance of digital ulcers), baseline characteristics of our 2 study groups were different, which may bias our results.

Interestingly, no new digital ulcers were detected in the patients with SSc during longterm treatment with ET-1 receptor antagonist combined with iloprost, in contrast to findings in other studies, in which the treatment with bosentan alone was associated with a reduction of only 30% in the number of new digital ulcers in the short term (24 weeks).32 Therefore, the intensive longterm combination treatment with intravenous iloprost (5-day cycles, with 24-h continuous intravenous infusion, every 3 months) seems to further enhance the effects of the ET-1 receptor antagonism compared to the monotherapy. It is known that ET-1 receptor antagonism combined with vasoactive drugs (i.e., iloprost or sildenafil) induces significant improvement in pulmonary arterial hypertension and reduces the appearance of new digital ulcers in patients with SSc.33,34 From the capillaroscopic viewpoint, in SSc patients with the more severe (late) NVC pattern at baseline (i.e., most patients in the ILO+BOS group), based only on the natural history of the disease, one should wait for further worsening of the microangiopathy during followup; but on the contrary, an increase of capillary number was observed precisely in the ILO+BOS group compared to the ILO group, showing the latter had less severe nailfold microangiopathy at baseline. This observation underlines the possibility of a real treatment-related effect of ET-1 receptor antagonism on nailfold microvascular damage. Nevertheless, for some clinical complications in SSc (pulmonary arterial hypertension and digital ulcers), progressive capillary loss, as detected by NVC, seems to have predictive value, and specific predictive indexes have been proposed.6,7,35,36

Longterm treatment of patients with SSc who have digital ulcers with the ET-1 receptor antagonist in combination with iloprost seems to interfere with the progression of nailfold microvascular damage, as assessed by NVC.

REFERENCES


