TITLE:
Long-term outcome and prospective validation of NIH response criteria in 39 patients receiving imatinib for steroid-refractory chronic GVHD

RUNNING HEAD:
CGVHD RESPONSE TO IMATINIB BASED ON NIH CRITERIA.

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KEY POINTS:

1. Efficacy of Imatinib in steroid-refractory chronic GVHD was prospectively compared across 3 different response systems, with high agreement

2. Validity of quantitative-based assessment of response with NIH criteria was confirmed by its prognostic impact on long-term survival

ABSTRACT:

Forty adults aged 28–73 years were entered into a prospective trial of imatinib for the treatment of steroid-refractory chronic graft-versus-host disease (SR-cGVHD). After 6 months, intention-to-treat (ITT) analysis of 39 patients who received the drug, regardless of the duration of treatment, revealed 14 partial responses (PR), 4 minor responses (MR) with relevant steroid sparing (46%) according to Couriel criteria, and 20 ≥PR (51.3%), as per the NIH criteria and NIH severity score changes. The best responses were seen in the lungs, guts, and skin (35%, 50%, and 32%, respectively). After a median follow-up of 40 (range, 12–54) months, 28 patients were alive, with 3-year overall survival (OS) and event-free survival of 72% and 46%, respectively. The 3-year OS was 94% for patients responding at 6 months and 58% for non-responders according to NIH response, suggesting that these criteria represent a reliable tool for predicting OS after second-line treatment. Monitoring of anti-platelet-derived growth factor receptor (PDGF-R) antibodies showed a significant decrease in PDGF-R stimulatory activity in 7 responders, while it remained high in 4 non-responders. In conclusion, this study confirms the efficacy of imatinib against SR-cGVHD and suggests that the response at 6 months significantly predicts long-term survival.

This study is registered at https://eudract.ema.europa.eu/, EUDRACT number: 2009-012927-27.
INTRODUCTION

Steroid-refractory chronic graft-versus-host disease (SR-cGVHD) represents an unmet challenge. The National Institutes of Health (NIH) consensus on chronic graft versus-host disease (cGVHD) has defined SR-cGVHD as follows: progression despite treatment with prednisone 1 mg/kg/day for ≥2 weeks or no improvement after 4-8 weeks on prednisone 0.5 mg/kg/day; or inability to taper below 0.5 mg/kg/day of prednisone [1].

The NIH consensus has also defined the global cGVHD severity score as mild, moderate and severe, the latter having the worst outcome [2-3]. Skin sclerosis is associated with poor functional status [4-5], poor quality of life and the need for prolonged immunosuppressive therapy [6].

Extracorporeal photopheresis (ECP) has shown a significant effect on skin SR-cGVHD, with a consistent steroid-sparing effect; however, there is no evidence of improved outcome [7]. In this setting, pentostatin has achieved promising results, albeit at the price of increased fungal infections [8]. Imatinib showed promising responses in 2 small cohorts of patients with SR-cGVHD, without major toxicities [9-10]; other studies have shown less favourable results [11-13]. Imatinib is a potent dual inhibitor of both transforming growth factor-beta and platelet-derived growth factor receptor (PDGF-R) pathways [14]; these 2 cytokines are both involved in the fibrogenic and inflammatory processes of several fibrotic diseases [15] as suggested by a murine model of cGVHD [16] and by recent data on cGVHD with fibrotic features (ScGVHD) and systemic scleroderma (SS) [17-18]. Moreover imatinib inhibits T-cell proliferation [19] as suggested by clinical improvement in patients with chronic myelogenous leukemia and concomitant autoimmune diseases receiving imatinib treatment [20].

Patients with ScGVHD have stimulating anti-PDGF-R antibodies as in SS, suggesting a similar pathogenetic mechanism [17-18]. However, the role of these antibodies is still debatable, and in a recent study of 15 cGVHD patients receiving imatinib, the presence of anti-PDGF-R antibodies did not correlate with outcome [12].

A major limitation of results from different trials on SR-cGVHD is the lack of standardized response criteria. Despite a considerable effort made by the NIH consensus [21] to create a uniform response evaluation, this is often based on physician assessment, with considerable variability in measures and a lack of reproducibility [7,8,22,23].

We therefore designed a larger study of imatinib in SR-cGVHD to prospectively evaluate the response rate using the Couriel criteria [22-23] and the NIH criteria [21]. Outcome according to response and NIH global score improvement [2-5] at 6 months was also evaluated.
PATIENTS AND METHODS

Study design

This prospective, multicenter, phase 2 study aimed to evaluate the safety and efficacy of imatinib in patients with SR-cGVHD. The protocol was approved by the co-ordinating center of Potenza Hospital’s ethical committee (EUDRACT number: 2009-012927-27) and at all participating sites. This study was conducted in accordance with the Declaration of Helsinki and was supported by AIFA (Italian Drugs Agency, CODE: FARM7ZWZ7Y).

The primary end points were as follows: 1) overall response rate (ORR) after 6 months of treatment with imatinib, evaluated according to the Couriel and NIH criteria [21-23], and 2) acute and long-term toxicity of grade >2, evaluated according to the World Health Organization (WHO) scale.

The secondary end points were as follows: a) Overall survival (OS) and event-free survival (EFS). OS was measured from the initiation of imatinib until death, while EFS was measured from the initiation of imatinib until death, secondary neoplasia, or treatment failure. Treatment failure was defined as: cGVHD progression or death because of cGVHD; relapse of the underlying disease; addition (or increase) of immunosuppressive drug/procedure, excluding a brief, transient steroid dose increase in the event of a GVHD flare [21]; or severe toxicity [grade 3–4 according to the common toxicity criteria (CTC)], requiring permanent discontinuation of imatinib. b) Response duration (RD), which was defined as time to loss of response (i.e. no response according to Couriel criteria) in patients achieving ≥ minor response/stable disease (MR/SD) at 6 months, according to Couriel criteria. Among patients with SD, only those with stable pulmonary function and a >50% decrease in steroid dose were included [22]. c) Changes in the NIH cGVHD global and organ-specific severity score during imatinib treatment [2-3-24]. d) Evaluation of stimulating anti-PDGF-R antibodies at baseline and during treatment to assess the correlation with the response to imatinib.

Statistical analysis

Using the Simon 2-stage, mini-max design, on the basis of the results of previous experimental treatment of SR-cGVHD [1], we set a minimal overall response rate (ORR, according to Couriel [22-23]) of 30% (p0) and an expected ORR of 50% (p1). Nineteen patients were recruited in the first stage, with an early stopping rule for futility if less than 7 responses occurred and a toxicity stopping rule if more than 1 death occurred potentially correlated to the treatment. In the second step, 20 more patients were to be recruited. This design yields a type I error probability of 0.05 (alpha), and a
power of 0.80 (1-beta); the true response rate is likely to be higher than 30% (p0) if at least 17 of 39 patients exhibited a response [25].

Data were analyzed using the Stata 12.0 package (Statacorp, College Station, TX, USA) and Microsoft Excel (Microsoft Corporation, Redmond WA, USA). Actuarial curves were plotted using the Kaplan–Meier method [26]. Univariate analyses for the outcome of OS were conducted for all the baseline characteristics reported in Table 1. All reported p-values in the text are 2-tailed (alpha = 0.05).

Intention-to-treat (ITT) analysis for response and outcome evaluation included the 39 patients receiving imatinib (regardless of the duration of treatment).

**Patient characteristics**

Forty consecutive adults with median age of 48 (range, 28–73) years, most with multiorgan cGVHD, were enrolled between February 7, 2008 and May 3, 2011. Patient characteristics are detailed in Table 1.

Eligibility criteria included active, moderate–severe cGVHD as per the NIH criteria [24] (requiring >0.4 mg/day of prednisone), with or without sclerotic features and refractory to at least 2 immunosuppressive lines of therapy. The NIH scoring system[24] was used at baseline for scoring cGVHD involvement in 5 organs (skin, mouth, gut, liver, and lungs); the global NIH severity (mild, moderate, severe) score was then calculated for all evaluable patients [3] and re-assessed after 6 and 12 months of treatment. Four patients had moderate cGVHD and 36 had severe cGVHD (Fig. 1). Severe scoring was attributable to multiple severe organ involvement in 13 patients and single severe organ involvement in 23 (13 with severe skin involvement and 10 with moderate/severe lung involvement). Thirty-one patients had skin involvement (25 with sclerotic features) and 33 had clinical lung involvement, with compromised forced expiratory volume and/or diffusion capacity of the lung for carbon monoxide. Twenty-four patients exhibited oral mucosal involvement, 9 exhibited gut involvement, and 6 exhibited liver involvement. A patient could have involvement of 1 or more of the abovementioned organs.

**Treatment schedule and concomitant therapy**

Imatinib treatment was planned for at least 6 months, starting at 100 mg/day during the first 15 days. In the absence of severe (grade 3–4 WHO) toxicity, the dosage was gradually increased to 400
mg/day [9]. Patients who achieved measurable response at 3 months continued imatinib up to 6 months. After 6 months, imatinib was continued at the discretion of the physician.

Patients on steroids, cyclosporine, or other concomitant immunosuppressive treatment (including ECP) were allowed to continue treatment, while those on myelotoxic drugs such as cyclophosphamide, pentostatin, or methotrexate were not. Patients treated with rituximab (RTX) in the previous 6 months were excluded to avoid a possible interference with the evaluation of anti-PDGFR antibodies. Supportive care/other medications were allowed according to institutional guidelines and/or local practice.

**Organ-specific and global response evaluation**

A center-assigned global response (Center Response), i.e. a clinical judgement reported by the local investigator on the basis of the Couriel criteria [22-23], expressed on a 4-point scale [Complete response (CR), Partial Response (PR), MR/SD, or no response/progressive disease (NR/PD)], was evaluated every 3 months after imatinib therapy. The decision to continue imatinib at 3 and 6 months was based on Center Response. Data about steroid dose modifications during imatinib treatment were also collected and integrated into the Center Response.

At baseline and at 6 months, NIH forms regarding clinician-assessed measures were completed, from which we derived 5 organ-specific core clinical measures of GVHD activity. Genital assessments, for which there were no robust objective measures; fascial/articular evaluations, which could partially overlap with sclerotic involvement; and eyes, whose damage is often irreversible, were not included [8, 21]. The data were collected centrally and organ-specific together with global responses were calculated according to the NIH response criteria proposed by Pavletic and coworkers (Appendix D) as CR, PR, SD and PD [21]. For the skin, an overall district response was evaluated from the 3 scales resulting from the 3 different skin manifestations (erythematous rash, moveable, and hidebound sclerosis). In the case of discordance between the response in 2 manifestations (i.e., PD in the erythematous and PR in moveable), the response was calculated on the overall body surface area; however, PD in the hidebound skin always indicated PD. For patients with lung involvement, improvement of at least 1 point on the lung functional score (LFS) without any evident clinical deterioration, was defined as PR. With regard to the other organs, the original NIH criteria were used according to the algorithm reported in the *Supplementary file n.1 (Algorithm).*
Patients were also classified according to modifications of the organ-specific NIH severity score (NIH SS) [3-24]. Briefly, NIH SS was calculated in each district at baseline and after 6 months. Changes were recorded as CR (complete resolution), PR (improvement of at least 1 point), SD (stable), and PD (worsening of at least 1 point).

After calculating organ-specific responses with NIH criteria and NIH SS, global response was determined as follows: CR was defined as CR in all affected organs; PR as CR/PR in at least 1 organ, without evidence of PD in any organ; SD as SD in each affected organ; and PD as PD in at least 1 organ.

**Antibody (Ab) purification and bioassay for anti-PDGFR Ab**

Immunoglobulins (Igs) from patients with cGVHD were purified from serum as previously described [17, 18]. Briefly, the stimulating ability of these antibodies to induce PDGF-R phosphorylation was evaluated using an in vitro fibroblast bioassay. Murine fibroblasts transduced with human PDGF-R were incubated with the patients’ purified Igs, and the reactive oxygen species (ROS) output was measured at baseline and after treatment with imatinib [17]. These tests were conducted for patients who received imatinib for at least 6 months, according to patient and center availability. Only the samples from patients free of imatinib treatment for at least 1 week were evaluated in order to avoid a possible interference with the PDGF-R pathway.

**RESULTS**

Forty patients were entered in the trial. One patient never began imatinib because the informed consent was reviewed with the local investigator and retracted. This patient received an alternative treatment and eventually died. The 39 patients who received imatinib were evaluated for toxicity, response, and outcome, regardless of the duration of treatment; 34 received treatment for ≥3 months. Imatinib was continued at 200 mg/day in 18 patients, increased to 300 mg/day in 8, and increased to 400 mg/day in 8.

Imatinib was administered at a mean dose of 270 mg/day for a median duration of 16 (range, 1–45) months. After 6 months, most patients (22) were receiving imatinib at 200 mg/day.

Thirteen patients discontinued imatinib within 1 year, while 14 were still receiving imatinib at the last follow-up.

**Toxicity**

No toxic deaths were observed. Details about the patients who experienced grade 1-2, 3 and 4 toxicity are reported in Tab 2. During imatinib treatment, 153 grade 1–2 and 23 grade 3–4 events
according to the CTC were recorded (Table 1S, Supplementary file n.2). Most events occurred during the first 3 months of treatment (90% of grade 3–4 events and 69% grade 1–2 events). Common adverse events (AEs) included fatigue, muscle pain, and weakness. Fluid retention was uncommon, and no severe heart or liver toxicity was observed. Hematological toxicity was mainly represented by anemia and thrombocytopenia (3 grade 3–4 events).

Twelve patients developed 13 infectious episodes: 7 episodes of pneumonia (2 due to *Aspergillus* and 1 due to *Candida albicans*), 2 of cytomegalovirus reactivation, 1 of sepsis (*Pseudomonas aeruginosa*), 1 of zoster reactivation, 1 of enteritis (salmonella), and 1 of influenza pneumonia (H1N1 virus) associated with the Aspergillus pneumonia. The 2 patients who developed Aspergillus pneumonia eventually died.

**Response**

Of the 39 patients receiving imatinib, 7 were not evaluable for response at 6 months because of treatment failure. Three patients died (1 each at 8 days, 6 weeks, and 3 months after initiating imatinib); 2 due to rapid cGVHD progression and 1 due to a pulmonary infection. Two patients experienced severe AEs after 2 and 10 weeks of imatinib treatment and received an alternative treatment (both were alive at the last follow-up). Two patients discontinued imatinib within 3 months because of cGVHD progression and received an alternative treatment (both were alive at the last follow-up).

The response at 6 months was evaluated using Center response, NIH response criteria and NIH SS change (Table 3, see details in Table S2, Supplementary file n.2). One patient with a GVHD flare after 5 months was treated with transient steroid increase and experienced notable improvement; because of the rapid changing situation, the local investigator judged him not evaluable for Center Response.

According to Center Response, we observed 14 PR and 12 MR/SD; of these, 4 achieved significant and sustained steroid sparing (>50% of the daily dose). Therefore, the ORR (including the criterion of steroid sparing used in our first trial [9]) was 46.1% (18/39 patients). According to the NIH criteria, 20/39 patients were classified as PR (51.3%) and 7 as SD. The best response rates were observed in the gut and lungs, with 50% and 35% of affected patients achieving PR, respectively. A retrospective evaluation of response based on changes in NIH SS revealed 20 PR (51.3%) and 9 SD. The response evaluation was grossly different in 2 patients (i.e., changing the status from ≥ SD/MR to NR/PD or vice-versa) as per the 3 response criteria. One patient exhibited a strong improvement in LFS (from 8 to 3) as well as an evident worsening in the skin according to the NIH criteria; exhibiting PD in the skin resulted in overall PD. Regarding the NIH SS, this patient had already the highest category of skin score (3) at baseline; therefore, he was judged to have SD; in the Center
Response, the local investigator classified him as a MR. The second patient exhibited severe lung involvement with the highest LFS (12), and pulmonary function further worsened at 6 months. This patient was judged to have SD by NIH response and NIH SS change and NR by Center Response. A few other minor discordances were observed. For example, a patient was classified as PR according to the NIH criteria because of an improvement in LFS (9 to 8), but as SD according to NIH SS changes: in fact the lung improvement was not enough to improve the NIH severity category in the lung; other affected organs were stable. This patient was judged as MR by the local investigator according to the Couriel criteria.

On initiation of imatinib, 14 patients were on concomitant steroid therapy. In 13 patients evaluable at 6 months (1 patient died), the median dose of prednisone (PDN) was 0.16 mg/kg at baseline and 0.03 mg/kg after 6 months. Two patients introduced PDN following imatinib discontinuation because of treatment failure; 4 discontinued PDN. According to Center Response, PDN dose was decreased in all 5 patients classified as PR at 6 months. Of the 7 patients with MR/SD, the dose was decreased in 5 and was unchanged in 2. Of the 3 patients with NR, the PDN dose was increased in 2 and discontinued in 1 because of inefficacy. Twelve months after the initiation of imatinib, 18 of 26 patients with ≥ MR/SD maintained their improvement: 9 maintained unchanged or decreased their previous immunosuppressive treatments and 9 were able to permanently discontinue them.

No significant differences (p = 0.29) were found in the mean imatinib dose between responders (202 mg/day) and nonresponders (221 mg/day).

Outcome

The median follow-up period was 40 months (minimum 12). Twenty-eight of the 39 patients were alive at the last follow-up. We observed 2 early deaths (before completing the first month of imatinib therapy) due to lung cGVHD progression. Five patients died because of cGVHD progression after 3, 7, 9, 15, and 16 months, respectively; 3 died from infection (1 each from necrotizing enteritis, lung candidiasis, and septic shock) after 6, 26, and 51 months, respectively; and 1 from relapsed acute myelogenous leukemia 24 months after imatinib discontinuation.

The 3-year OS and EFS were 72% [95% confidence interval (CI), 54%–84%] and 46% (95% CI, 29%–61%), respectively. In the 26 patients who achieved ≥MR/SD after 6 months, RD was 69% at 36 months (95% CI, 46%–84%; Fig. 2). In the 14 patients with ≥PR according to Center Response, RD was 83.1% at 36 months (data not shown). A landmark analysis for OS at 6 months according to response status revealed that achieving ≥ PR at this time-point (regardless of the response criteria utilized) strongly predicted the outcome (Fig. 3A, B, C), suggesting that adopting a centralized NIH
response criteria is a reliable tool for predicting outcome. According to univariate analyses, the baseline characteristics reported in Table 1 were not significantly associated with a difference in OS.

**Anti-PDGF-R antibodies and response**

All the 11 patients evaluated with the functional bioassay [17-18] had relevant anti-PDGF-R antibodies activity before treatment; of these 7 responded to imatinib, whereas 4 did not. The relative decrease in PDGFR agonistic activity in terms of ROS stimulation (ROS index) was evaluated at different time points, and a significant difference was observed between the two groups (p = 0.006). The ROS index was decreased in all responders, while in the nonresponders, the anti-PDGF-R activity did not substantially change compared with that at baseline, suggesting a relationship between clinical response and the resolution of agonistic antibody-mediated PDGF-R stimulation (Fig. 4). There was no significant difference in the mean imatinib dose between responders and nonresponders (p = 0.38), and no correlation was observed between the imatinib dose and change in ROS index (r = −0.12).

**DISCUSSION**

The management of SR-cGVHD is problematic, and the evidences are limited to phase II trials or retrospective studies [7-9-28-29].

Survival at 5 years among patients with high-risk cGVHD is reported to be approximately 50% [1-5-31], with a median survival of only 30 months [2]. In a recent study, the NIH SS was an independent factor for predicting cGVHD-specific survival [31]. A recent retrospective study in 312 patients receiving second-line treatment for cGVHD showed a 56% failure-free survival (FFS) at 6 months, and NIH SS emerged again as an independent factor influencing outcome. Moreover, this study suggested that the 6-month FFS may represent an early surrogate end point for predicting the long-term efficacy of a second-line treatment for SR-cGVHD in clinical trials [27].

Our study demonstrates that response at 6 months strongly correlates with OS, regardless of the criteria adopted, suggesting that this early end point is highly predictive of the long-term outcome. There was a low discordance rate using the 3 different response criteria; however, the feasibility of the NIH response criteria has been extensively evaluated in both retrospective [32] and prospective [33] studies. Our study is the first one to assess the feasibility of NIH response criteria in the context of a therapeutic trial and validate their prognostic impact on survival. Although further confirmation in larger randomized trials and in different populations of cGVHD patients is required,
NIH criteria should be considered the new standard for measuring therapeutic response in future prospective trials of cGVHD. The use of an automatic algorithm defining the kind of response according to NIH recommendations (Supplementary File n.1) may facilitate application in clinical practice.

As per the Couriel criteria, the response rate in this trial was lower than that in our first trial [9], but it should be noted that the median age of the population treated in this trial was significantly higher than of the first cohort (48 vs 27 years), which also included pediatric patients. The incidence of severe toxicity was acceptable; 2 patients died of lung aspergillosis, a common complication of SR-cGVHD regardless the kind of immunosuppressive treatment.

In our first trial, only patients with skin fibrotic features were treated, while this study also included patients with visceral involvement, suggesting that imatinib is effective against all types of cGVHD. In a retrospective French study, 39 patients with SR-cGVHD on imatinib, were evaluated by the physician, according to his perception of the change in skin involvement, as "improvement," "stability," or "worsening." According to this empirical approach, the authors found an overall improvement rate of 30%. Systemic corticosteroids were tapered or discontinued in 41% patients [13]. Other experiences with imatinib for SR-cGVHD have yielded less encouraging results: Chen et al. [12] evaluated 15 patients according to Hopkins criteria and reported an ORR of 40% at the last follow-up. Stadler [11] reported only 2 PR among 9 patients with severe SR-cGVHD of the lungs. Several factors such as differences in baseline characteristics of patients, duration of imatinib treatment, higher variability because of a smaller sample size, and evaluation of only some organs with different response criteria can explain, in part, the lower ORR found in these studies.

Despite the low number of patients, our data suggest a relationship between response to imatinib and anti-PDGF-R antibody activity. In contrast to the findings of Chen et al. [12], we observed that while patients had a relevant anti-PDGF-R activity at baseline, the stimulating activity of these antibodies significantly decreased in the responding patients. Chen et al. used a different assay for detecting the anti-PDGF-R antibodies; therefore, we cannot exclude that these authors assessed antibodies directed against different PDGF-R epitopes. Moreover, the behavior of the anti-PDGF-R antibodies during treatment suggests that imatinib may also act through an immune-modulating effect, leading to a decrease in the level of these antibodies in the responders.

In conclusion, imatinib represents a valuable option for patients with SR-cGVHD who cannot access other treatments such as ECP; this treatment is simple because it requires neither hospitalization nor long term central venous access. In our study Imatinib resulted feasible in 85%
of the patients, achieving a clinically meaningful response in approximately half of patients. Our data support further investigation of this agent in SR-cGVHD; indeed the true efficacy of imatinib has to be confirmed and tested in larger randomized controlled trials.

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Explanation of Author Contributions:
A.O., A.B., M.C. conceived and designed the study; A.O., M.C., P.C., N.M., R.F., C.S., F.O., F.P., E.P. and P.B. provided study materials or patients; S.S., A.P. and A.G., performed, analyzed and interpreted the biological data. M.C., J.O., S.P., R.N. and A.O. collected, analyzed and interpreted the clinical data; A.O., and A.B. wrote the manuscript.

Conflict of Interest Disclosure:
The authors declare no competing financial interests.
REFERENCES


Blood: 2011 118: 4242-4249

[3] Patient-reported quality of life is associated with severity of chronic graft-versus-host disease as measured by NIH criteria: report on baseline data from the Chronic GVHD Consortium
Blood: 2011 117: 4651-4657

[4] Correlation between NIH composite skin score, patient-reported skin score, and outcome: results from the Chronic GVHD Consortium
Blood: 2012 120: 2545-2552

[5] Sclerotic-type chronic GVHD of the skin: clinical risk factors, laboratory markers, and burden of disease
Martires KJ, Baird K, Steinberg SM, Grkovic L, Joe GO, Kirsten M, Williams KM, Mitchell SA, Dattiles M, Hakim FT, Pavletic SZ and Cowen EW
Blood: 2011 118: 4250-4257


Blood: 2008 112: 2667-2674

[8] Phase II Study of Pentostatin in Patients With Corticosteroid-Refractory Chronic Graft-Versus-Host Disease
JCO September 20, 2007 vol. 25 no. 27 4255-4261
[9] Imatinib for refractory chronic graft-versus-host disease with fibrotic features.
Blood. 2009 Jul 16;114(3):709-18

[10] Imatinib mesylate as salvage therapy for refractory sclerotic chronic graft-versus-host disease.
Blood. 2009 Jul 16;114(3):719-22


Blood: 2011 118: 4070-4078

[13] Limited efficacy and tolerance of imatinib mesylate in steroid-refractory sclerodermatous chronic GVHD.
Blood. 2012 Dec 13;120(25):5089-90

[14] Dual inhibition of c-abl and PDGF receptor signaling by dasatinib and nilotinib for the treatment of dermal fibrosis

Oliveri J, Coluzzi S, Attolico I, Olivieri A.

McCormick LL, Zhang Y, Tootell E, Gilliam AC.

[17] Stimulatory autoantibodies to the PDGF receptor in systemic sclerosis.

[18] Stimulatory autoantibodies to PDGF receptor in patients with extensive chronic graft-versus-host disease.
Blood. 2007 Jul 1;110(1):237-41

[19] Imatinib inhibits T-cell receptor-mediated T-cell proliferation and activation in a dose-dependent manner.
Seggewiss R, Loré K, Greiner E, Magnusson MK, Price DA, Douek DC, Dunbar CE, Wiestner A.


[22] Extracorporeal photochemotherapy for the treatment of steroid-resistant chronic GVHD


Filipovich AH, Weisdorf D, Pavletic S et al.
Biology of Blood and Marrow Transplantation 11:945-955 (2005)

Simon R.

Gooley TA, Leisenring W, Crowley J, Storer BE.

[27] Failure-free survival after second-line systemic treatment of chronic graft-versus-host
disease.
Inamoto Y, Storer BE, Lee SJ, Carpenter PA, Sandmaier BM, Flowers ME, Martin PJ.


[29] Interleukin-2 and regulatory T cells in graft-versus-host disease.
VT, Treister NS, Bienfang DC, Prasad S, Tzachanis D, Joyce RM, Avigan DE, Antin JH, Ritz J,

Lee SJ, Vogelsang G, Flowers MED

[31] Incidence and outcome of chronic graft-versus-host disease using National Institutes of
Health consensus criteria.
S, Greer J, Kassim A, Morgan D, Ruffner K, Schuening F.

Leukemia. 2009 Jan;23(1):78-84.

[33] A multicenter pilot evaluation of the National Institutes of Health chronic graft-versus-
host disease (cGVHD) therapeutic response measures: feasibility, interrater reliability, and
minimum detectable change.
Mitchell SA, Jacobsohn D, Thormann Powers KE, Carpenter PA, Flowers ME, Cowen EW,
Schubert M, Turner ML, Lee SJ, Martin P, Bishop MR, Baird K, Bolaños-Meade J, Boyd K, Fall-
Dickson JM, Gerber LH, Guadagnini JP, Imanguli M, Krumlauf MC, Lawley L, Li L, Reeve BB,
Clayton JA, Vogelsang GB, Pavletic SZ.
### Patient characteristics (n = 40)

<table>
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<tr>
<th>SEX:</th>
<th>Female/Male n (%)</th>
<th>12/28 (32%/68%)</th>
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<td>AGE:</td>
<td>Median (range)</td>
<td>48 (28–73)</td>
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#### Disease, n (%)

| Myeloproliferative disorders: CML/MF/MDS | 3/1/1 (12%) |
| MM                                      | 8 (20%)     |
| Lymphoproliferative disorders: NHL/HD/CLL | 5/2/0 (18%) |
| Others: solid tumor                      | 1 (2%)      |

#### Transplant, n (%)

| MUD | 8 (20%) |
| HLA identical sibling | 32 (80%) |
| PBSC | 30 (75%) |
| BM | 10 (25%) |

#### Conditioning regimen, including TBI

| Conditioning regimen, including TBI | 17 (44%) |

#### cGVHD

| Duration: Median (range) in months | 26 (3–148) |

#### Risk score (according to Inamoto et al.): high/intermediate/low, n

| Risk score | 24/15/1 |

#### FIRST-LINE THERAPY, n

| Prednisone (alone or associated with other drugs) | 36 |
| Miscellaneous (including MF, tacrolimus, ECP, CSA) | 4 |

#### SECOND AND SUCCESSIVE LINES OF THERAPY, n

| Rituximab+/miscellaneous (No ECP) | 5 |
| Rituximab+ECP+/−miscellaneous | 8 |
| Miscellaneous, including ECP (no rituximab) | 19 |
| Miscellaneous, excluding ECP and rituximab | 19 |

#### Karnofsky performance status at enrollment, n (%)

| Karnofsky performance status at enrollment | <70 (28 (70%)) |
| 80 | 9 (22%) |
| 90–100 | 3 (8%) |

#### Platelet count before imatinib initiation, 10⁹/L, n (%)

| Platelet count before imatinib initiation | <100 (2 (5%)) |
| >100 | 38 (95%) |

#### Main cGVHD TARGETS (Different organ involvements are not mutually exclusive)

| Skin, n (%) | 32 (80%) |
| Mean affected BSA% for erythematous/moveable/nonmoveable sclerosis | 19/27/27 |
| Lung, n (%) | 33 (82.5%) |
| Median values of FEV1/DLCO | 61/55 |
| Mouth, n (%) | 23 (57.5%) |
| Median Schubert Scale value | 3 |
| Liver, n (%) | 6 (15%) |
| Median values of ALT (U/l)/total bilirubin (mg/dL) | 74/3.0 |
| Gut, n (%) | 9 (22.5%) |
| Mean value of the GI scale (0–3) | 1.5 |


---

**Table 1: Baseline characteristics of the 40 enrolled patients**
<table>
<thead>
<tr>
<th>TOXICITIES</th>
<th>Patients experiencing toxicities of grade:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1–2</td>
</tr>
<tr>
<td><strong>Extrahematological</strong></td>
<td></td>
</tr>
<tr>
<td>Nausea</td>
<td>7 (17%)</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>5 (12%)</td>
</tr>
<tr>
<td>Cramps</td>
<td>14 (35%)</td>
</tr>
<tr>
<td>Muscle pain</td>
<td>11 (28%)</td>
</tr>
<tr>
<td>Peripheral edema</td>
<td>6 (15%)</td>
</tr>
<tr>
<td>Periorbital edema</td>
<td>5 (12%)</td>
</tr>
<tr>
<td>Asthenia</td>
<td>13 (33%)</td>
</tr>
<tr>
<td>Emesis</td>
<td>2 (5%)</td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>5 (12%)</td>
</tr>
<tr>
<td>Rash</td>
<td>2 (5%)</td>
</tr>
<tr>
<td>Kidney</td>
<td>5 (12%)</td>
</tr>
<tr>
<td>Muscle weakness</td>
<td>9 (23%)</td>
</tr>
<tr>
<td>Liver</td>
<td>1 (2%)</td>
</tr>
<tr>
<td>Neuropathy</td>
<td>1 (2%)</td>
</tr>
<tr>
<td>Weight gain</td>
<td>2 (5%)</td>
</tr>
<tr>
<td>Heart failure</td>
<td>2 (5%)</td>
</tr>
<tr>
<td>Bullous dermatitis</td>
<td>0</td>
</tr>
<tr>
<td><strong>Hematological</strong></td>
<td></td>
</tr>
<tr>
<td>Anemia</td>
<td>25 (64%)</td>
</tr>
<tr>
<td>Neutropenia</td>
<td>6 (15%)</td>
</tr>
<tr>
<td>Thrombocytopenia</td>
<td>4 (10%)</td>
</tr>
</tbody>
</table>

Table 2: Main extrahematological and hematological toxicities in the 39 patients treated with imatinib
<table>
<thead>
<tr>
<th>CENTER (Couriel)</th>
<th>NIH CRITERIA</th>
<th>CHANGES IN NIH SEVERITY SCORE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>RESPONSE</strong></td>
<td><strong>OVERALL</strong></td>
<td><strong>RESPONSE</strong></td>
</tr>
<tr>
<td>ORR corr.* 46%</td>
<td>ORR 36%</td>
<td>ORR 32%</td>
</tr>
<tr>
<td>ORR 36%</td>
<td>CR 0</td>
<td>CR 3</td>
</tr>
<tr>
<td>MR/SD 12</td>
<td>MR/SD 12</td>
<td>PD 2</td>
</tr>
<tr>
<td>NR/PD 5</td>
<td>NE 8</td>
<td>NE 4</td>
</tr>
<tr>
<td>TOT. 39</td>
<td>TOT. 39</td>
<td>TOT. 31</td>
</tr>
<tr>
<td>EVAL. 31</td>
<td>EVAL. 31</td>
<td>EVAL. 27</td>
</tr>
</tbody>
</table>

Table 3: Global and organ-specific response according to center evaluation (response based on Couriel criteria), NIH response criteria, and changes in NIH severity score (NIH SS).

ORR: overall response rate, calculated as (CR + PR)/TOT. (total number of patients receiving imatinib), ORR corr.*: denotes corrected ORR, taking into account patients with MR/SD with concomitant steroid sparing (>50% of the initial dose), CR: complete response, PR: partial response, MR: minor response, SD: stable disease, NR: no response; PD: progressive disease, NE, not evaluable [NE category includes patients who did not undergo a complete response evaluation at 6 months because of treatment failure within 6 months (7) or other reasons (1 for Center Response)], TOT.: total patients receiving imatinib included in the intention-to-treat analysis, EVAL.: includes patients undergoing a complete response evaluation at 6 months according to NIH, Center, and changes in NIH SS response.

According to Center (Couriel) criteria, patients with SD and without significant steroid-sparing were considered NR: in the category MR/SD we included patients with SD and >50% steroid reduction, and all patients with MR [22].


**Figure legends**

**Fig. 1:** Individual organ severity scoring within global severity categories

**Fig. 2:** Overall survival (A) and event-free survival (B) in the 39 patients receiving imatinib and response duration (C) in the 26 patients who achieved ≥MR/SD (minor response or stable disease with steroid sparing) at 6 months according to Center Response. OS was measured from the initiation of imatinib until death, while EFS was measured from the initiation of imatinib until death, secondary neoplasia, or treatment failure. Response duration (RD) was defined as time from response evaluation at 6 months to loss of response (NR/PD) in patients achieving ≥ minor response/stable disease (MR/SD) at this timepoint, according to Center Response. Among patients with SD, only those with stable pulmonary function and a >50% decrease in steroid dose were included [22].

**Fig. 3:** Thirty-six of the 39 patients receiving imatinib were alive at 6 months and were included in a landmark analysis for overall survival (OS) according to response status at 6 months (in this analysis responders were those patients included in the ORR reported in Table 3, not corrected for steroid sparing). The 36-month OS was significantly higher for responders (dotted line) than for nonresponders (solid line): 94% vs. 55%, 100% vs. 60%, 94% vs. 58% according to changes in NIH SS (A), Center Response (B), and NIH criteria (C), respectively.

**Fig. 4:** Mean change in the ROS stimulatory index (ROI) of cGVHD immunoglobulins before and after imatinib treatment according to response status (Center Response).
Fig. 2

A. Overall Survival

B. Event-Free Survival

C. Duration of Response
Fig. 3

A. Overall Survival by ORR (changes in NIH SS at 6m) 

B. Overall Survival by ORR (Center Response at 6m) 

C. Overall Survival by ORR (NIH response criteria at 6m) 

$p = 0.004$ 

$p = 0.02$ 

$p = 0.007$ 

Number at risk 

Non-responders 

Responders 

Non-responders 

Responders 

Non-responders 

Responders 

0

0

0

0

0

0
Fig. 4

Change in ROI (ROS index) before and after Imatinib treatment

According to response status

% Change in ROI

Non-responder

Responder
Long-term outcome and prospective validation of NIH response criteria in 39 patients receiving imatinib for steroid-refractory chronic GVHD

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