Chlormethine Gel is Efficient and Safe in Mycosis Fungoides Skin Lesions

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Chlormethine is a bifunctional cytotoxic alkylating agent that binds to DNA, resulting in cell death (apoptosis). Chlormethine (also known as mechlorethamine) gel (CL gel) was approved in the European Union in 2017 and was first used in 2019. The aim of the study is to examine evidence regarding the efficacy and safety of chlormethine gel in everyday clinical experience from a cutaneous lymphoma centre. Twenty-three patients with stage IA–IIB mycosis fungoides received chlormethine gel between September 2020 and May 2021. All patients started by applying the gel daily and were monitored every month. At 1, 3, 6 and 9 months, 0%, 43.47%, 56.52% and 65.22% of patients, respectively, achieved an overall response. Five out of 23 patients (21.73%) achieved near complete response at a mean time of 6 months. Chlormethine gel was given as monotherapy in 12 patients (52.17%), and in addition to systemic treatments (methotrexate and peginterferon alpha-2a) in 11 patients (47.82%). Adverse events (AE) were recorded in 43.47% of patients, but only 3 discontinued treatment, due to dermatitis. Scale down of the treatment to application 3-times per week led to better patient compliance. This study shows that chlormethine gel is effective and safe in patients with mycosis fungoides with different types of skin lesions.

Key words: chlormethine; cutaneous lymphoma; mycosis fungoides; modified Severity Weighted Assessment score; mSWAT.

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The choice of treatment depends on the type of PCL and the stage of the disease. Recommendations are based on consensus by lymphoma societies (3, 4). The 3 most common skin treatments for early-stage MF are topical steroids, phototherapy, and topical chlormethine (also known as mechlorethamine), a bifunctional cytotoxic alkylating agent that results in cell death (apoptosis). In 1950, aqueous chlormethine (Caryolysine) received initial approval in the USA for topical treatment of MF (and was also used in Europe). In the early 1980s, chlormethine compounded in Aquaphor (a petroleum-based ointment) was introduced. All compounded forms had the disadvantage of instability and lack of reproducibility of results.

A novel topical chlormethine (CL) gel was developed in 2004, and was approved for use in the USA and EU in 2013 and 2017, respectively, based on a randomized controlled trial 201 (5), in which the efficacy and safety of 0.02% CL gel were found to be non-inferior to 0.02% compounded ointment (5). The overall response to treatment with 0.02% CL gel (59%) was greater than with the compounded 0.02% CL ointment (48%). In study 201 (5), 20% of enrolled patients treated with CL gel and 17% treated with CL ointment withdrew due to drug-related skin irritation. Among the adverse events (AEs) reported, dermatitis was the main reason for discontinuing treatment. The use of steroids was not allowed within the registration trial, but concomitant use of corticosteroids with CL gel is currently being studied, with the aim of better understanding the potential use of corticosteroids for management of such skin reactions. A number of case studies and clinical practice experiences have been published, highlighting that, in the real-world setting, CL gel is well-tolerated, with treatment-emergent...
AEs generally mild in nature and effectively managed with appropriate topical interventions and dosing modifications (6, 7).

The aim of this study was to evaluate the real-life efficacy and side-effect profile of CL gel in patients with MF.

PATIENTS AND METHODS

Patients with different types of classic MF skin lesions (patches, plaques, tumours) were treated with CL gel at the National Center of excellence for the treatment of CL, Attikon University Hospital, Athens, Greece, between September 2020 and May 2021. All patients had a histologically confirmed diagnosis of MF, except for 1 patient with gamma-delta epidermotropic PCL, which was diagnosed as MF on clinical grounds.

All patients were evaluated at baseline, and at months 1, 3, 6 and 9, for efficacy and AEs. Overall response was recorded using the modified Severity Weighted Assessment Tool (mSWAT) score, time to response (TR), and time to next treatment (TNT).

Patients were instructed to use the CL gel at night. In case of dermatitis, a dose reduction to 3 times per week was advised, in addition to emollient creams. In cases of severe dermatitis, use of clobetasol and temporary discontinuation of CL gel was advised. Patients were instructed to continue with treatment with close-monitoring and frequent visits.

RESULTS

The baseline demographic and clinical characteristics of patients treated with CL gel are shown in Table I. Twenty-three patients, 16 men and 7 women (male:female ratio 2.3:1), median age 69.5 years (range 52–87) were treated. Median duration of disease was 8 years (range 2–15 years). Before gel application, patients had a median mSWAT of 54.73 (range 4–105.45) and 18/23 (78.26%) were early-stage (7/23 stage IA, 11/23 stage IB), and 5/23 (21.73%) advanced-stage IIB MF. CL gel was given as monotherapy in 12/23 patients, and in combination with methotrexate (MTX) or peginterferon alpha-2a in 10/23 patients.

The results for efficacy and AEs are shown in Table II. At month 1, all patients had stable disease (SD). At month 3, 10/23 patients showed partial response (PR) with mean reduction in mSWAT 60% from baseline, while 8/23 patients remained at SD. At month 6, 13/23 patients showed overall response, with 5/13 near complete response (NRCR, 90% mSWAT improvement) and 8/13 with PR. At month 9, 15/23 patients achieved overall response; 5/15 showed CR with mSWAT 0, 2/15 NRCR, and 8/15 showed PR with reduction in mSWAT of 75%.

Overall response in stages IB/IIB was delayed compared with IA (this result was not statistically significant due to the small number of patients, and remains to be clarified in future studies).

At month 1 of gel treatment, AEs were observed in 10/23 patients with MF and at month 3 these had resolved (Table II). Mild dermatitis was observed in 7/10 patients, and 3/23 patients developed severe dermatitis at month 1, leading to ulceration in 2/3 patients at month 3 with high overall response at months 6 or 9 (2 of these achieved PR and 1 CR). All patients continued treatment with CL gel with resolution of AEs at months 6 and 9. No skin malignancy was observed during treatment with CL gel.

One stage IIB patient died due to respiratory infection, which was not related to the treatment, 3 discontinued treatment in the first month due to dermatitis that could not be tolerated, and 1 patient with gamma-delta epidermotropic MF discontinued treatment due to disease progression.

| Characteristics | Age, years, median (range) | 69 (52–87) | Male sex, n (%) | 16 (69.5) | Disease duration, years, median (range) | 8 (2–15) | mSWAT prior to gel treatment, median (range) | 54. (4–105.45) | Stage, n (%) | IA | 7 (30.43) | IB | 11 (47.82) | IIB | 5 (21.73) |
|-----------------|-----------------------------|---------------|-----------------|-------------|--------------------------------------|-----------|------------------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|

Table I. Baseline demographics and clinical characteristics of 23 patients treated with chlormethine (CL) gel

Table II. Efficacy and adverse events in 23 patients treated with chlormethine (CL) gel

| Efficacy per month: n (%) or median (range) | Month 1 | Month 3 | Month 6 | Month 9 |
|---------------------------------------------|---------|---------|---------|---------|
| IA: 7/23 | SD: 3-> stop treatment | PR: 4/4 | NCR: 4/4 | CR: 4/4 |
| IB: 11/23 | SD: 1-> stop treatment | PR: 5/10 | NCR: 3/10 | PR: 6/10 |
| IIB: 5/23 | SD: 1-> death IIB stage death cause: respiratory infection | PR: 1/4 | NCR: 1/4 | CR: 1/4 |
| Total: 23/23 (100%) | SD: 3/23 (13.04%) | PR: 2/4 | NCR: 2/4 | PR: 2/4 |

Adverse events:
- Mild dermatitis 7/23 (30.43%) | Medium-to-severe dermatitis 3/23 (13.04%) |
- Itching (dermatitis) 8/23 (34.78%) | Ulceration (dermatitis) 2/23 (8.69%) |
- Skin hyperpigmentation: 8/23 (34.78%) | SD: stable disease; OR: overall response; PR: partial response; NRCR: near complete response; CR: complete response.
DISCUSSION

This study of a small group of patients found that CL gel is a well-tolerated and safe treatment for MF lesions, such as patches, plaques and tumours. A better and more rapid response was observed in patients with early and limited disease (IA). All patients with MF treated with CL gel achieved stable disease at month 1, and at month 3 half of the patients achieved PR. Despite close monitoring 3 out of 7 stage IA patients discontinued treatment in the first month due to dermatitis. In order achieve better compliance the frequency of application of CL gel was reduced to 3 times per week. At month 3, all stage IA patients who continued the gel achieved PR, while at month 9 all achieved CR. Of the 11 stage IB patients, 6 patients achieved PR, while at month 9 4 patients achieved NRCR and 2 achieved CR. One out of 4 stage IIB patients achieved NRCR and 2 achieved PR from month 3. All stage IIB patients were on systemic treatment, and achieved clinical response when CL gel was added to treat a few localized tumour lesions refractory to systemic treatment. Notably, 1 stage IIB patient on peginterferon alpha-2a with partial response achieved CR at month 9 of combination treatment with CL gel. To the best of our knowledge, this is the first study to use CL gel in all clinical MF lesions including tumours, which were not included in study 201.(5). The response rates at all months were similar to those in study 201 (5). CL gel was well tolerated, with good clinical outcomes reported from month 1. Since CL gel was available in June 2019 in Greece, some of the current study patients had not yet reached 1-year treatment, and peak clinical response among the stage IA–IIB patients was higher at month 9 (65.22%). In the PROVE study, which has a longer follow-up, peak response was seen later (at 18 months, 66.7%) among stage IA–IB patients (6).

CL gel was generally well tolerated, except for in 10 patients who developed dermatitis. Throughout the current study 43.47% of patients developed some degree of dermatitis, 13.04% developed medium-to-severe dermatitis during the first month of treatment, but only 3 patients using CL gel discontinued treatment due to dermatitis. Based on results from clinical practice, the management of gel application was adjusted from daily to 3 times per week, which led to better patient compliance. Management of patients with severe dermatitis was through close monitoring and down-tapering of gel use to 3 times per week and use of potent topical steroids (Table III). It was observed that the presence of dermatitis did not have an impact on the efficacy of CL gel. More specifically, 3 patients with severe acute dermatitis finally achieved overall response. In the PROVE study, among 298 adult patients with MF-CLTf, dermatitis/skin irritation rates (12.8%/7.4%, respectively) were lower than observed in study 201, possibly due to concomitant steroid treatment and/or dosing modifications (8). In the current study no malignancy was observed at gel application sites at 9-month follow-up. In the MIDAS study, which is investigating the incidence and severity of contact dermatitis following treatment with CL gel in patients MF-CTCL stages IA–IB, mild-to-moderate dermatitis may not require suspension of treatment, but may require emollients or topical steroids or decreased dosing frequency (7, 9) (Table IV).

Table III. Treatment proposals of adverse events

| Treatment | Response rates (month 9) |
|-----------|-------------------------|
| Mild dermatitis | Reduce frequency of chlormethine gel application every other day | 3 PR |
| 7/23 patients | Combined with topical emollients | 3 CR |
| | When dermatitis is persistent add very potent topical steroids | 1 SD |
| Severe dermatitis | Very potent steroids (clobetasol propionate) initially once/daily | 2 PR |
| 3/23 patients | Reduce frequency chlormethine gel application 2 or 3 times per week | 1 CR |

SD: stable disease; PR: partial response; CR: complete response.

Table IV. List of studies on chlormethine (CL) gel

| Study | Year | Type of study | Patients N | Efficacy | Dermatitis, % |
|-------|------|---------------|------------|----------|---------------|
| 201   | 2013 | RCT           | 260        | RR 59% (CAILS) 46.9% (mSWAT) | 15–20 |
| PROVE | Ongoing | Prospective observational study of real world | 301/298 during MG | Time to response demonstrated superiority of CL gel to ointment (p<0.012) 1/3 (33.3%) of participants with Stage IA–IB MF-CTCL CR at month 12 2/3 (66.6%) CR at month 18 CL gel was well tolerated with responders reporting significantly improved HR-QoL compared with non-responders | 12.8 |
| MIDAS | Ongoing | Non-randomized open-label split-face 2-arm study | 78 | CAILS assessment showed similar clinical responses to CL gel with or without concomitant corticosteroid treatment over 6 months | 33.3 |
| Current study | 2019 | Non-randomized open-label | 23 | OR: 15/23 (65.22%) NRCR: 2/15 (13.33%) CR: 5/15 (33.33%) PR: 8/15 (53.33%) at month 9 | 43.47 |

mSWAT: modified Severity Weighted Assessment score; CR: complete response; NRCR: near complete response; OR: overall response; PR: partial response; SD: stable disease; HR-QoL: health-related quality of life; MF-CTCL: mycosis fungoides cutaneous T cell lymphoma; MIDAS: mechlorethamine induced dermatitis assessment study (9); PROVE: the prove study (8); RCT: randomised control trial; CAILS: composite assessment of index lesion severity; MG: monitoring.
In our experience, close monitoring is very important in order to maintain patients on treatment. We monitor patients closely throughout treatment, especially during the first months when AEs usually occur. During the national lockdown due to the COVID-19 pandemic, we continued to monitor most of our patients via teleconferencing, urging them not to stop treatment. In 5 patients with difficulty getting the treatment on time, a relapse in skin disease, but not progression, was observed. In the meantime, patients were instructed to continue treatment with clobetasol propionate. After the end of national lockdown, approximately 3 months, patients restarted treatment with CL gel with no recurrence of AEs. In a large study of 4,922 patients, it was shown that, in settings with higher patient volume, patients sustained longer treatment duration and, importantly, avoided early discontinuation due to better management of the disease and dermatitis (10). In the current study, only 4 patients discontinued treatment, which is attributed to our increasing experience and good physician–patient inter-relationship.

The current study has some limitations. First, the chronic period of CL gel use was short due to the availability of CL gel in Greece. However, our last follow-up at 12 months has not shown significant differences. From the 15 patients with overall response at month 9, 2/15 (13.3%) patients achieved CR at month 12, 8/15 (53.3%) patients achieved PR at month 12, and 5/15 (33.3%) patients completed treatment with CL gel, maintaining the response for at least 1 month after treatment. Finally, in patients who had to discontinue treatment, the study did not perform a patch test in order to differentiate allergic reactions from irritant reactions.

Both physicians and patients should be educated in the management of MF with CL gel treatment and close monitoring, in order to avoid premature discontinuation and loss of compliance. Continuous use of CL gel treatment is required in order to achieve maximum response. Larger studies with more data are important in order to assess the use of CL gel in patients with different MF lesions.

The authors have no conflicts of interest to declare.

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