

Methylene Blue for the Treatment of Intractable Pain From Oral Mucositis Related to Cancer Treatment: An Uncontrolled Cohort

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ABSTRACT

Background: Oral mucositis is a complication of cancer therapy, causing severe pain that affects oral functioning, nutrition, and quality of life, as well as therapy nonadherence or dose-limiting toxicity. Anecdotal experience has suggested that methylene blue (MB) oral rinse may be an effective and safe treatment of this oral pain. **Methods:** To evaluate the efficacy and safety of MB oral rinse for the treatment of oral pain due to mucositis in patients with cancer, we retrospectively evaluated patients who experienced refractory pain despite conventional therapy. **Results:** We identified 281 patients who received MB oral rinse. Most were receiving treatment for leukemia (n=85; 30.3%) and head and neck squamous cell carcinoma (n=84; 29.9%). The most common treatments were radiation therapy alone (n=108; 38.4%) and chemoradiation (n=86; 30.6%). Median duration of symptoms was 14 days. Mean (SD) numeric rating scale pain scores were 7.7 (1.83; median, 8) before MB oral rinse and 2.51 (2.76; median, 2) after MB oral rinse ($P<.0001$). Most patients achieved pain control within the first 3 doses. The effectiveness of MB oral rinse was independent of patient age, sex, cancer type, cancer stage, MB dilution, and pain duration or baseline pain scores. The lowest response to treatment was reported in individuals with esophageal mucositis. Few patients experienced adverse effects of MB oral rinse (n=13; 4.6%); 10 had a transient burning sensation, 2 had transient blue discoloration of the teeth and mouth, and 1 had increased pain. **Conclusions:** MB oral rinse is an effective and safe treatment for refractory pain from oral mucositis related to cancer treatment.

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Background

Oral mucositis (OM) is a painful inflammation and ulceration of the oral mucous membranes that affects millions of patients with cancer. It is a treatment-related toxicity associated with chemotherapy, radiation therapy, and hematopoietic stem cell transplantation (SCT).¹ Oral lesions cause pain, resulting in dehydration, malnutrition, medication noncompliance, interruption in therapy, or increased risk of infection. Pain caused by OM may lead to impairment of oral function, which may require more aggressive supportive measures, including placement of a transcutaneous gastric feeding tube, parenteral nutrition, or frequent hospitalizations, exponentially increasing morbidity and cost.²

Lesions that develop after chemotherapy or during preparation for SCT occur approximately 3 to 5 days after treatment initiation. In contrast, postradiation OM exhibits a gradual progression and potentially lasts for weeks after completion of radiation therapy. Ulcerations caused by mucositis are typically nonuniform and can merge to form one large lesion.³ The ulcerations are not the result of an inflammatory process and the borders are not erythematous unless there is a concomitant infection. In addition, lesions tend to develop on less keratinized areas and therefore are rarely found on the dorsum of the tongue, gingiva, and hard palate.³ The pathophysiology of OM includes mucosal and epithelial damage secondary to endothelial injury in the lamina propria.⁴ Symptoms correlate with the severity of mucositis, fluctuating between local irritation and a burning sensation to severe debilitating pain.

Efforts have increased to prevent OM and histologically reverse oral lesions; however, pain is the major problem causing interruption of treatment and less favorable prognosis.⁵ Current strategies to manage OM include oral hygiene, prophylactic antimicrobials, selective diets to decrease bacterial growth, and avoidance of foods that exacerbate pain, in addition to topical therapies, including coatings, artificial saliva, and neutralizing

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solutions. Preventive agents such as palifermin, a keratinocyte growth factor, and amifostine, a free-radical scavenger, as well as low-level laser therapy have shown promising results but have varying levels of evidence for reducing the severity of mucositis; therefore, expert and empirical judgment have been the recommended approach in guidelines for managing mucositis.^{6–9}

Managing oral pain has been the most substantial challenge in addressing OM. Conventional alternatives include use of topical anesthetics found in different mixes, compounded products, and oral rinses. Emerging options for pain control include oral doxepin rinse, ketamine mouthwash, bupivacaine lozenge, polyvinylpyrrolidone-sodium hyaluronate gel, and CAM2028. These therapies have varied yet encouraging results but are not exempt from adverse effects and toxicity.^{10–15} To date, there is no standard efficacious therapy to prevent or treat OM-related pain in patients with cancer.¹⁶ Escalating doses of opiate analgesics remains a suboptimal but commonly used alternative. Unfortunately, many patients lack access to or cannot afford opioids, and concerns exist regarding opioid dependency and addiction-related drug diversion.

In the search for effective options to treat OM, novel topical and systemic agents have failed. More recently, studies have been conducted of methylene blue (MB), a partially liposoluble vital dye used for a variety of diagnostic and therapeutic indications. The structural composition of MB seems to affect different levels of the nociceptive pathways, and has been useful in reducing pain syndromes. It causes inhibition of nitric oxide synthetase, guanylyl cyclase, and histamine¹⁷; pain and analgesia have been shown to be in part modulated by guanylyl cyclase. A study by Salman et al¹⁸ demonstrated a decrease in pain intensity during induction of anesthesia when MB was first injected compared with placebo. A systematic review by Fransiska et al¹⁹ described a local analgesic effect of MB after anorectal surgery pain by temporarily disrupting the sensory nerve conduction.

Prior to this, MB had never been used to treat oral pain caused by mucositis secondary to cancer treatment. Thus, in 2017 we pilot-tested a diluted MB oral rinse to treat uncontrolled oral pain from cancer therapy. Our preliminary clinical data suggested that MB oral rinse was safe and effective for controlling pain in patients with this condition.²⁰ We soon learned that providers within other specialties at The University of Texas MD Anderson Cancer Center had rapidly adopted our mix as a second-line treatment of OM in patients for whom conventional therapy had failed. Therefore, we initiated the current study to investigate the efficacy and safety of MB oral rinse for OM-related pain in patients receiving cancer treatment.

Methods

Study Setting and Population

This retrospective study was performed at our academic tertiary care cancer center and was approved by the Institutional Review Board of The University of Texas MD Anderson Cancer Center. No written informed consent from participants was required.

Data Collection

Study candidates were electronically identified by searching an institutional pharmacy database using the 2Know Request System for billing codes associated with a diagnosis of OM (ICD-9 and ICD-10 codes 99232, 99213, and 99214) between December 1, 2017, and December 1, 2019. From this search, we identified patients who were prescribed MB oral rinse to treat pain from OM. We then selected and reviewed the medical records of patients whose medication was dispensed to use.

Selection and Description of Participants

We considered patients of any age, sex, cancer diagnosis, and disease stage, and included all treatment modalities (chemotherapy, radiation therapy, SCT, surgery, and combinations). Pain intensity among patients with documented oral pain from cancer treatment-related OM was measured using a numeric rating scale (NRS; scores ranged from 1–10).

Patient information was stored in a secure database, and access to the database for secondary projects such as the current report required a separate approval by the Institutional Review Board, which was obtained.

Procedures

Patients with intractable oral pain from OM were instructed to use MB oral rinse. Various dilutions of MB were prescribed (0.025%, 0.05%, and 0.1%) and patients were instructed to swish, gargle, and spit every 6 hours until achieving pain control. The dilutions given seemed to be based on provider preference or pharmacy availability. Other modalities of pain management were not discontinued, including systemic analgesics and oral rinses; however, patients had not experienced sufficient pain relief despite use of these treatments.

Outcome Measures

We aimed to compare self-rated NRS scores of oral pain before treatment with MB oral rinse with those reported after treatment. If documentation was available, we also reviewed the morphine equivalent daily dose (MEDD) used, WHO grade of OM, and the location, nature, and duration of pain before treatment with MB oral rinse. In addition, we collected information about the duration of the analgesic effect experienced with the medication and any adverse effects reported.

Statistical Analysis

Our plan was to include 300 patients in the retrospective study. With a sample size of 300, a 2-sided 95% confidence interval for the mean pain reduction would extend to 0.113 times the standard deviation from the observed mean based on the large-sample z statistic (nQuery Advisor 7.0; Statsols). Continuous variables were summarized using descriptive statistics, including mean, standard deviation, median, and range. Continuous variables included patient age, pain medication reduction, NRS before and after treatment, duration of pain before treatment, duration of pain relief, duration of pain reduction after treatment, dose of MB, and concentration of MB used for treatment. Categorical variables were tabulated by frequencies and the corresponding percentages, and included WHO grade before treatment, adverse effects of MB oral rinse, and patient sex. Chi-square or Fisher exact test was used to evaluate the association between categorical variables. Wilcoxon signed rank test was used to evaluate whether the change in pain score from before to after treatment was significantly different from zero ($P < .05$ indicated statistical significance). Wilcoxon rank sum test was used to evaluate the difference in a continuous variable between patient groups. A boxplot was generated as a visual aid to show the difference in the distribution of a continuous variable between or among patient groups. Statistical software SAS 9.4 (SAS Institute Inc.) and Splus 8.2 (TIBCO Software Inc) were used for all analyses.

Results

Patient Characteristics

We identified 350 adult patients who were prescribed MB oral rinse for oral pain from mucositis related to cancer therapy. Per pharmacy files, 31 individuals (all outpatient) could not afford the compounding charges from the pharmacy. In addition, 6 patients declined to use the medication and 32 medical records had inadequate documentation, precluding imputation of the data. The final study population consisted of 281 patients. All patients had documentation of actively receiving oral rinses, and most received concomitant opiate analgesics in oral or parenteral form.

The providers prescribing MB oral rinse were from several departments, including supportive care, emergency medicine, critical care, leukemia, lymphoma, sarcoma, breast cancer, radiation oncology, head/neck oncology, SCT, urology, and internal medicine.

Demographic Characteristics

Demographic information for the 281 patients included in the analysis is summarized in Table 1. Mean age was 55 years (range, 18–83 years), and there were 114 women

Table 1. Patient Demographic Characteristics

Characteristic	n (%)
Patients, N	281
Mean age (range), y	55 (18–83)
Sex	
Male	167 (59.4)
Female	114 (40.6)
Cancer diagnosis	
Leukemia	85 (30.3)
Head and neck SCC	84 (29.9)
Lymphoma	43 (15.3)
Germ cell	13 (5.0)
MDS	11 (3.9)
Multiple myeloma	10 (3.6)
Gastrointestinal	9 (3.2)
Sarcoma	6 (2.1)
Other	20 (7.1)
Cancer therapy, N	261
Radiation therapy	114 (43.7)
Chemoradiation	91 (34.9)
Stem cell transplantation	3 (1.1)
Chemotherapy alone	39 (15.0)
Surgery and radiation	14 (5.4)
Mean duration of pain (\pm SD)/median, days	23.31 (\pm 38.34)/14

Abbreviations: MDS, myelodysplastic syndromes; SCC, squamous cell carcinoma.

and 167 men. Ethnicity was not considered relevant. The most common diagnoses included leukemia ($n=85$; 30.3%) and head and neck squamous cell carcinoma ($n=84$; 29.9%). Most patients were actively undergoing cancer therapy ($n=261$; 92.9%). The most common therapeutics used were radiation alone ($n=108$; 38.4%) and chemoradiation ($n=86$; 30.6%). Median duration of mucositis-related pain was 14 days.

Clinical Characteristics

Clinical patient characteristics are summarized in Table 2. Most patients ($n=207$; 73.7%) had painful mucosal lesions of the oral and esophageal structures; the oral mucosa was the most commonly reported ($n=188$; 66.9%). Most commonly, pain description was burning ($n=155$; 70.1%). Only 126 patients had WHO grade of mucositis documented, with most grade III ($n=82$; 65.1%). The MB dilution used most commonly was 0.025% ($n=179$; 63.7%).

Efficacy of MB

Among the 281 patients, 243 had pain scores reported both before and after MB oral rinse therapy. Mean (SD)

Table 2. Patient Clinical Characteristics

Characteristic	n (%)
Patients, N	281
Location of pain	
Oral mucosa	188 (66.9)
Sublingual	86 (30.6)
Oropharynx	26 (9.3)
Esophageal	23 (8.2)
Soft palate	21 (7.5)
Inner lips	12 (4.3)
Multiple locations	207 (73.7)
Pain description, N	221
Oral burning	155 (70.1)
Aching	56 (25.3)
Sharp	10 (4.5)
Cutting blades	27 (12.2)
WHO grade, N	126
IV	14 (11.1)
III	82 (65.1)
II	30 (23.8)
Adverse effects	
Transient burning sensation	10 (3.6)
Transient discoloration	2 (0.7)
Increased pain	1 (0.4)
MB dilution used	
0.025%	179 (63.7)
0.05%	90 (32.0)
0.1%	12 (4.2)

Abbreviation: MB, methylene blue.

NRS pain score before MB oral rinse therapy was 7.7 (1.83; median, 8) and after therapy was 2.51 (2.76; median, 2). Mean (SD) NRS pain score reduction was 5.12 (2.97; median, 5), and the Wilcoxon signed rank test indicated that the change in pain score from before to after treatment was significantly different ($P < .0001$; Figure 1). Among the 243 patients, 205 (84.4%) had a baseline NRS pain score ≥ 7 and 38 had a baseline NRS pain score < 7 (15.6%). The difference in pain score change among the different MB dilution groups was not statistically significant ($P = .66$; Figure 2).

The maximum pain relief was reported within minutes of the first dose in most patients ($n = 124$; 58.5%); the remaining patients required several doses—up to 9 doses (48–72 hours)—to achieve sufficient pain relief (Table 3). Although more than half of the patients in whom pain scores were reported achieved sustained pain relief after the first dose ($n = 131$; 55.5%), others reported pain recurrence between 4 and 8 hours ($n = 58$; 24.6%). However,

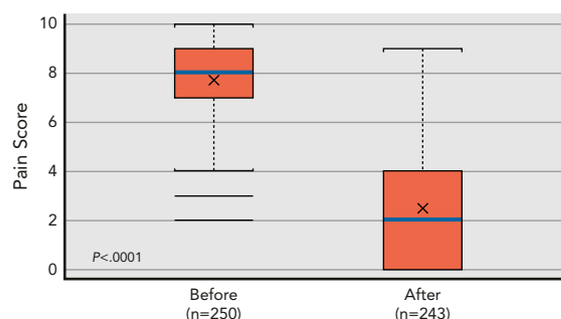


Figure 1. NRS pain scores before and after treatment with MB oral rinse. Bar graph shows the mean (X inside the bar), median (white bar inside the box), and outliers (horizontal lines beyond the end of the vertical dotted lines) of the pain scores before (left) and after (right) treatment with MB oral rinse. Pain reduction was significantly different from zero according to the Wilcoxon signed rank test ($P < .0001$). Abbreviations: MB, methylene blue; NRS, numeric rating scale.

the subsequent intensity was reported at lower scores than at baseline, which suggested an accumulative analgesic effect up to complete control within 3 days. In the remaining cases, recurrence was not clearly documented.

Many patients had documented multisite pain beyond OM; therefore, conclusions regarding the effect of MB oral rinse on MEDD could not be drawn.

Oral functioning (ability to talk, chew, and swallow) before and after MB was inconsistently documented, and 5 patients had already had a gastric tube placed before using the MB oral rinse.

Association Between Demographic Characteristics and MB Efficacy

Reduction in NRS pain scores during a specified time frame was documented in 212 patients, who were divided into interval groups according to reduction in pain scores: > 5 , 2 to 5, and < 2 (Table 3). Most patients whose pain scores were reported achieved pain reduction of ≥ 5 on the NRS ($n = 138$; 65.0%), and only 11 patients (5.2%) reported pain reduction of < 2 . The difference in pain score change among age groups was not statistically significant ($P = .48$). Similarly, the difference in pain score change between sexes was insignificant ($P = .1853$). Patients with esophageal lesions reported lower pain score changes compared with others ($P < .05$).

Adverse Effects

Few mild and transient adverse effects were reported ($n = 13$; 4.6%; Table 2). A total of 10 patients experienced an oral burning sensation during first-time use, with 1 subsequently discontinuing therapy. Two patients reported transient blue discoloration of the teeth and mouth, which resolved with oral hygiene. One patient experienced increased pain during MB oral rinse use, resulting in discontinuation of therapy.

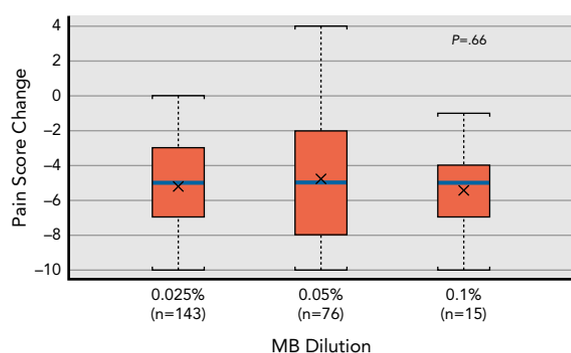


Figure 2. NRS pain scores after treatment with MB oral rinse at 3 different dilutions. Bar graph shows the mean (X inside the bar) and median (white bar inside the box) change in pain score after treatment with MB oral rinse at 0.025%, 0.05%, and 0.1% dilution. The graph shows no statistical difference among the groups according to the Wilcoxon rank sum test ($P=.66$).

Abbreviations: MB, methylene blue; NRS, numeric rating scale.

Discussion

In 2017, we conducted a pilot clinical test of MB oral rinse (0.05%) in 5 patients with severe oral pain from mucositis related to cancer therapy.²⁰ Based on promising results, a phase II prospective trial of MB oral rinse was designed and implemented. It included 4 treatment arms: 3 using MB oral rinse at 0.025%, 0.05%, and 0.1%, respectively, and 1 using conventional therapy alone (ClinicalTrials.gov identifier: NCT03469284). Although data were being collected, preliminary results of safety and efficiency became visible to providers from different departments within our institution who adopted our protocol for clinical use (including MB doses and frequency of use). Because of the increasing number of documented prescriptions of MB oral rinse, we were able to collect sufficient data for the current study. Our results support the use of MB oral rinse as an easy-to-use, safe, and efficient therapy for intractable pain associated with OM in patients with cancer.

When administered orally, MB is absorbed through the gastrointestinal tract, with peak plasma concentration

occurring in 1 to 2 hours.²¹ However, this concentration is 100-fold less than concentrations achieved after intravenous administration of an equivalent dose.²² This significant difference is attributable to first-pass metabolism in the liver after oral administration.²³ Therefore, the pharmacokinetics of MB suggest that if diluted to 0.025% to 0.1%, and if strictly used as an oral rinse, the plasma concentration of MB absorbed is potentially negligible and therefore exempt from pharmacologic interactions or toxicity. In this study, no toxicities or medication interactions were reported, suggesting that MB is safe when used as an oral rinse. If accidentally ingested, MB is metabolized to leukomethylene blue and excreted primarily in urine, turning the urine a blue-green color.²⁴ The terminal half-life is estimated to be 5.25 hours.²⁵ This is our rationale for prescribing its use only every 6 hours, to avoid any remote chance of systemic accumulation.

The only on-label use of MB approved by the FDA is for the treatment of methemoglobinemia, for which MB is administered intravenously.²⁶ Through this route, MB has also been used for the management of vasoplegia (a high-output shock state with poor systemic vascular resistance, a common occurrence in patients undergoing cardiothoracic surgery) refractory to catecholamines, and as an adjuvant to treat cyanide toxicity.^{27,28} Oral MB has been used in the treatment of malaria in sub-Saharan Africa²¹ and for the treatment of ifosfamide-induced encephalopathy.²⁹ For diagnostic purposes, MB is used for tissue staining to help detect various gastrointestinal pathologies.^{30–32} Similarly, intra-articular MB is used to establish the integrity of the capsule of the knee joint in trauma patients.³³ When coupled with visible light therapy, antimicrobial properties of MB are used to inactivate lipid-enveloped viruses.³¹ Furthermore, as a 2% topical solution, MB has been effective in treating onychomycosis.³⁴ Lastly, a 0.1% topical solution of MB in combination with fluorescent light has been used in the treatment of genital herpes simplex infections.³⁵

Table 3. Peak Pain Reduction Per Time and Dose Received of MB Oral Rinse (N=212)

Time of Peak Pain Relief	Any Pain Reduction n (%)	Pain Reduction >5 on NRS ^a n (%)	Pain Reduction 2–5 on NRS ^a n (%)	Pain Reduction <2 on NRS ^a n (%)
0 h (dose 1)	124 (58.5)	76 (61.3)	41 (33.1)	7 (5.6)
0–2 h (dose 1)	6 (2.8)	4 (66.7)	2 (33.3)	0 (0.0)
2–12 h (dose 1–2)	11 (5.2)	9 (81.8)	2 (18.2)	0 (0.0)
12–24 h (dose 2–4)	52 (24.5)	36 (69.2)	14 (26.9)	2 (3.8)
24–36 h (dose 4–6)	1 (0.5)	1 (100.0)	0 (0.0)	0 (0.0)
36–48 h (dose 6–8)	10 (4.7)	5 (50.0)	4 (40.0)	1 (10.0)
48–72 h (dose 8–10)	8 (3.8)	7 (87.5)	0 (0.0)	1 (12.5)

Abbreviations: MB, methylene blue; NRS, numeric rating scale.

^aPercentages in these columns refer to the number of patients who reported any level of peak pain reduction at that time point.

Systemic use of MB is contraindicated in pregnant and nursing women, patients with glucose-6-phosphate dehydrogenase deficiency, and in individuals allergic to it.³⁶ MB can cause hemolytic anemia in patients with glucose-6-phosphate dehydrogenase deficiency because of its decreased ability to metabolize MB.¹⁸ In addition, because MB is a monoamine oxidase inhibitor, it must be used with caution in patients receiving other proserotonin medications.³⁷ In addition, MB can stain tissue, skin, clothing, or whatever else it may come in contact with.

MB seems to be a long-term inhibitor of peripheral axons and an antioxidant and anti-inflammatory agent, which is why it is a useful analgesic.³⁸ When used as an oral rinse, MB is thought to denature free nociceptive nerve endings, inhibit the nitric oxide inflammatory pathway, and block *N*-methyl-D-aspartate receptors.^{39–41} In this study, the lower pain response in patients with esophageal lesions supports the need for direct contact between MB and the compromised mucosa, given that it was not swallowed.

Unlike other oral rinses containing topical anesthetics, MB does not cause local anesthesia, which is manifested as intraoral numbness. Thus, MB does not alter the taste of ingested meals. In addition, MB seems to have an accumulative analgesic effect, as opposed to the short-lasting pain relief encountered with other rinses compounded with local anesthetics.

Given that the MB available in our setting is only in the form of vials for intravenous use, high cost is a limitation to its use. In addition, compounding charges are a tremendous barrier for many patients, even those who have health insurance. Surprisingly, while visiting a

developing country we found MB easily accessible and inexpensive at local pharmacies in a nonintravenous form. It was tested by local providers with results similar to those we have reported.

This study had a few limitations. Inadequacies in the data acquisition could not be addressed due to the retrospective nature of the study. In addition, a discrepancy in the size of the comparison groups could not be reconciled. Lastly, we could not objectively address MEDD before and after MB oral rinse therapy.

Conclusions

MB oral rinse is a promising therapy for pain from OM related to cancer therapy, although long-term benefits and risks have yet to be explored. MB is a low-risk, efficient, easy-to-use treatment for refractory pain in OM related to cancer therapy, and potentially accessible to patients of all socioeconomic backgrounds in the absence of compounding charges.

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