Role of lasers in treatment of connective tissue diseases

Connective tissue diseases (CTD) is a group of systemic disorders, characterized by clinically unique skin manifestations often resistant to conventional therapies [1]. In this article, Brauer et al have reviewed the role of lasers in treatment of these manifestations. By searching MEDLINE, they found 39 related articles.

Lupus erythematosus (LE): Its skin manifestations include in acute and chronic cutaneous lupus, oral and nasal ulcers, and non-scarring alopecia. The efficacy of pulsed dye laser (PDL) in treating all kinds of cutaneous LE has been revealed in 8 articles. The relative efficacy of argon laser in improving discoid LE (DLE) has been reported in 2 articles. Successful improvement of DLE with prolonged remission has been reported with ablative carbon dioxide (CO₂) and ablative yttrium aluminum garnet (YAG) lasers.

Scleroderma and morphea: Skin manifestations of scleroderma are summarized in skin fibrosis, Raynaud’s phenomenon, and telangiectasias. The efficacy of PDL in treating telangiectasias has been revealed, but about en coup de sabre and plaque morphea, the results have been variable. Intense pulsed light (IPL) has successfully been used for treating microstomia. The efficacy of ablative and fractional ablative CO₂ lasers has been demonstrated in improving rhytides, calcinosis of the digits, and joint contractures. Additionally, the efficacies of excimer and 1064-nm neodymium-doped YAG (Nd-YAG) have been reported in treating morphea lesions and severe Raynaud’s disease, respectively.

Sarcoidosis: Regarding existence of non-caseating granulomas, its skin manifestations are classified into specific and non-specific lesions. PDL has successfully been used in treating cutaneous sarcoidosis. In remodeling of lupus pernio, CO₂ laser has provided positive responses. However, among the CTDs, sarcoidosis has had the most adverse events from laser therapy.

Dermatomyositis: Its skin lesions include calcinosis, poikiloderma, Gottron’s papules, periungual telangiectasias, heliotrope eruption, shawl and holster signs, and mechanic’s hands. The efficacy of PDL and argon laser have been reported in improving telangiectasias, poikiloderma, and Gottron’s papules.

Commentary: Lasers can find proper placement in the treatment of skin manifestations of CTDs.

Therapeutic options for atrophic acne scarring

Acne vulgaris is the most common skin disease among adolescents [2]. Atrophic scars, as its common sequel, result in significant psychological distress. They are subdivided into boxcar, icepick, and rolling scars. In this article, Hession and Graber reviewed options for treating atrophic acne scars.
its treatment especially in early stages is important. Classically, the curettage, cautery, and surgical excision are standard modalities for the treatment of KA. Other therapeutic options include topical imiquimod, topical 5-flurouracil, and photodynamic therapy with topical 8-aminolaevulinic acid.

Podophyllin is an antimitotic, cytotoxic, and caustic agent. It can cross cell membranes, and inhibit cell mitosis and DNA synthesis through binding to tubulin. It also prevents polymerization of tubulin into microtubules, resulting in arrest in cell division. Furthermore, podophyllin can block oxidation enzymes in tricarboxylic acid cycle, interfere with nutrition of cells, and inhibit axonal transport, protein, RNA, and DNA synthesis. It also inhibits mitochondrial activity and reduces cytochrome oxidase activity. This toxic agent can induce local necrosis and death of tumor cells.

In previous studies, the efficacy of podophyllin has been shown in treating genital wart, cutaneous leishmaniasis, basal cell carcinoma, and solar keratosis. In the current study, Sharqie and Noaimi introduced podophyllin as an effective and safe therapeutic option for KA. They recruited three cases with KA (2 female xerodermoid patients and 1 male non-xerodermoid patient). Topical 25% podophylline solution was administered once weekly for maximum of 6 weeks. The patients were followed up for 18 months.

Sharqie and Noaimi showed that podophylline was very effective for treating KA with good cosmetic results. Complications or side effects were not reported in this study. Eventually, they concluded that topical podophyllin can be an appropriate option for the treatment of KA in patients refusing surgical approaches.

Commentary: Podophylline can be an effective and safe agent for the treatment of keratoacanthoma.

Mechanisms underlying aggressive manner of melanoma

Melanoma has an aggressive manner, with metastasizing to multiple organs including lung, liver, brain, bone, and lymph nodes.[4]. In this article, Braeuer et al have comprehensively reviewed the probable mechanisms underlying this manner:

Antigenicity: Melanoma cells are highly antigenic and share most of their antigens with vascular cells. These antigens allow melanoma cells to adhere to the vessel wall in distant organs, and extravasate into the parenchyma. Furthermore, these molecules result in homotypic melanoma cell adhesion and emboli formation.

Regarding variability of antigens on endothelial cells in different organs, depending on those shared with melanoma cells, melanoma has site-specific metastatic characteristics.

Local invasion: Expression of melanoma cell adhesion molecule MUC18 in melanoma is associated with activation of matrix metalloproteinases, leading to degradation of the basement membrane. Furthermore, expression of cadherins in melanoma cells is responsible for adhesion of malignant cells to keratinocytes and fibroblasts.

Vasculogenic mimicry: Expression of VE-cadherin in melanoma causes formation of capillary-like structures in tumor.

Angiogenesis and lymphangiogenesis: This tumor is highly angiogenic and lymphangiogenic, attributed to inflammatory molecules secreted by both malignant cells and the microenvironment.

Seed and soil hypothesis: Genetically unstable melanoma cells (seed) find the appropriate organ microenvironment (soil).

Premetastatic niche: Tumor produced-soluble molecules and bone marrow-derived cells result in melanoma cell proliferation, invasion, and metastasis via promoting formation of the metastatic microenvironment.

Stemness: Melanoma cells can act as stem cells resulting in repopulation of new tumors.

Parallel vs. linear progression: There is two models of metastasis in melanoma; one parallel to primary tumor progression and another one, metastasis from metastasis.

Immunogenicity: Although melanoma is very immunogenic, it evade cytotoxicity via some methods.

Transcriptional regulation: The progression of melanoma from the radial growth phase to the vertical one is attributed to changes in the gene transcription factors.

Mutation: Melanoma has the highest mutation rate in comparison with other malignancies.

Complementary: With understanding the mechanisms underlying aggressive manner of melanoma, we can successfully treat this tumor with target-targeting weapons.

The efficacy of nalfurafine hydrochloride in the treatment of pruritus

Studies have shown that the μ- and κ-opioid receptors play a great role in the regulation of pruritus in the central nervous system; on the other hand, the epidermal keratinocytes also express these receptors.[5]. Nalfurafine hydrochloride is a potent and selective nonpeptide agonist of κ-opioid receptors with tyrosine-glycine moiety for endogenous opioid peptides, which was constructed in 1998 by a Japanese group. It appears that this agent exerts its antipruritic activity through the κ-receptors in the epidermis. In this study, Inui reviewed the efficacy of this drug in the management of pruritus induced by hemodialysis and other factors:

Hemodialysis-induced pruritus: Its exact pathophysiology is not well-understood. The probable roles of secondary hyperparathyroidism, histamine, mast cell-released tryptase, skin barrier dysfunction-induced increased epidermal neuron fibers, and the upper neuron system have been suggested in the pathogenesis of this kind of pruritus. The management of pruritus in patients under hemodialysis is a big challenge. Oral agents such as gabapentin, and thalidomide, topical agents such as endocannabinoid, capsaicin and tacrolimus, and ultraviolet B therapy have been administered with some success in managing of this kind of resistant pruritus. Different studies have shown the effectiveness and safety of nalfurafine in treating hemodialysis-induced pruritus.

Atopic dermatitis: As a causative factor in atopic dermatitis-induced pruritus, down-regulated expression of epidermal κ-opioid receptors has been suggested. It appears that nalfurafine can be effective in treating pruritus in patients with atopic dermatitis.

Psoriasis: Pruritus has been reported in some cases of psoriasis. It appears that the epidermal expression of κ-opioid receptor and dynorphin A is significantly decreased in these cases. Hence, nalfurafine can be useful in managing this kind of pruritus.

Cholestasis: Study on rats has revealed the potential effectiveness
of nalfurafine in treating cholestasis-induced pruritus.

**Drug-induced pruritus:** The effectiveness of nalfurafine has been demonstrated in chloroquine-induced pruritus.

**Commentary:** Nalfurafine hydrochloride can be a good option for treating recalcitrant pruritus.

**References**


