Nicolau’s syndrome following diclofenac administration: A report of two cases

Sir,

Nicolau’s syndrome (NS) is a rare injection site reaction, following intramuscular administration of drugs, with varying degrees of tissue damage. It is also synonymously described as embolia cutis medicamentosã [1] and livedoid dermatitis. [2] NS is characterized by development of an acute, severe pain and a localized erythematous rash following intramuscular injection. Subsequently cutaneous, subcutaneous and even muscular necrosis with a pale marble-like livedoid pattern results. [3] We report here two cases of NS following diclofenac administration.

Our first case was a 29 year-old man admitted with a history of snakebite, referred from the medicine department for evaluation of a painful ulcer over the right gluteal region. He had received two intravenous infusions of antivenin and an intramuscular (intragluteal) injection of diclofenac. After receiving diclofenac, he immediately noticed severe pain followed by blistering and ulceration.

Cutaneous examination showed a large tender, nonindurated ulcer with necrotic eschar covering almost the entire right gluteal region with minimal extension to the left [Figure 1]. There was no regional lymphadenopathy. Other cutaneous and systemic examinations were normal. Complete hemogram including bleeding time, clotting time and urine examinations were normal. Chest X-ray, blood urea, serum creatinine, liver function tests, creatine kinase were normal. Venereal disease research laboratory (VDRL), human immunodeficiency (HIV)-1 and 2 tests were negative. Culture from the ulcer showed growth of Staphylococcus aureus sensitive to ciprofloxacin.

The patient was treated with surgical debridement, sterile dressings, analgesics and oral ciprofloxacin 500 mg twice daily for 14 days. The ulcer healed completely with scarring in 14 weeks.

Our second case was a 70 year-old man who presented with a painful red lesion over the left gluteal region of three days’ duration. He had received an intramuscular (intragluteal) injection of diclofenac sodium for arthralgia prior to the onset. He noticed pain at the injection site immediately, followed by the development of the painful red lesion. There was no other drug intake or systemic illness.

Cutaneous examination showed a solitary, tender, nonindurated, nonblanchable, livedoid patch with dendritic extensions over the left gluteal region [Figure 2]. There was no regional or generalized lymphadenopathy. Other cutaneous and systemic examinations were normal. Routine hematological and urine examinations were normal. Blood VDRL, HIV-1 and 2, liver function tests, blood urea, serum creatinine, creatine kinase were normal. Skin biopsy showed thrombosis of blood vessels consistent with NS. The patient was started on analgesics and topical betamethasone ointment twice daily. Ulcer and crust were noted in a week and he was further managed conservatively with analgesics and sterile dressings. The lesion completely healed in ten weeks with atrophic scarring.

First described by Freudenthal in 1924 and Nicolau in 1925, [4] NS was recognized as an adverse effect of bismuth salts.
used in syphilis.[2] Subsequently NS has been associated with phenylbutazone, diclofenac, ibuprofen, vitamins K and B complex, sulfapyridine, tetracycline, streptomycin, sulfonamide, lidocaine, phenobarbital, chlorpromazine, dexamethasone, trimcinolone, diphenhydramine, interferon alfa, gentamicin, ketoprofen, influenza and diphtheria pertussis toxin (DPT) vaccination.[3]

The pathogenesis of NS is obscure. Intraarterial or periarterial injection of the drug may be the cause. The mechanism may involve direct trauma or arterial embolism caused by the drug or ischemia due to compression following paravascular injection.[1-3] Vascular pathogenesis involving arterial vasospasm with resultant ischemia-mediated livedoid necrosis, may be another possible mechanism.[2] Diclofenac was the drug responsible for NS in both of our cases. There are only a few reports of NS associated with diclofenac.[1,2,4-6] It possibly causes NS by vascular pathogenesis as it acts via the cyclooxygenase pathway, inhibiting prostaglandin synthesis with resultant vasoconstriction.[2]

Severe pain at the injection site may possibly be due to involvement of peripheral sensory nerves. Immediate pallor and edema occur, followed by a circumscribed red-violet, hemorrhagic plaque, with dendritic extensions. Necrotic plaques, ulcers, bullae, erosions and crusts may occur over the livedoid plaque.[1,3] Secondary bacterial infection may occur.[2] It heals in a few months with atrophic scarring. The gluteal region is most commonly affected although other sites of intramuscular injections such as the thighs may be involved.[1,3] NS needs to be differentiated from hematoma at the injection site.[2]

In the acute phase of NS, histopathology shows epidermal necrosis and thrombosis of small and medium blood vessels. Conservative treatment with dressings, debridement and pain control are the mainstay of therapy.[3] Therapy ranges from topical corticosteroids to excision.[1] Vasoactive medication may also be beneficial.[7]

This report highlights an uncommon adverse effect of a commonly used drug-diclofenac. Caution should be exercised during administration of parenteral NSAIDs especially diclofenac to prevent this rare reaction. Awareness and early recognition of NS will help in proper management.

**S. Chidambara Murthy, Karjigi Siddalingappa, T. Suresh**

Department of Dermatology and Venereology, Vijayanagara Institute of Medical Sciences, Bellary, Karnataka, India

Address for correspondence: Dr. S. Chidambara Murthy, Department of Dermatology and Venereology, Vijayanagara Institute of Medical Sciences, Bellary - 583 104, Karnataka, India. E-mail: chidumurthy@yahoo.com

### REFERENCES

4. Pillans PI, O’Connor N. Tissue necrosis and necrotizing fascitis after intramuscular administration of diclofenac.
Supernumerary digits associated with pachyonychia congenita type I

Sir,

An 18 year-old girl presented with thickening and discoloration of all nails, painful callosities on soles, whitening of tongue and papules over extensors of legs and back. On enquiry, it was discovered that discoloration of nails started within a few months of birth. Gradually nails became so thick and hard that by the age of four years, it was difficult to cut the nails. Whitening of tongue, callosities on the soles and papules over the extremities and back developed later. There was no history of palmo-plantar hyperhydrosis, bullae, cysts, hoarseness of voice, teeth malformation, hair loss or any systemic complaints. No family history of similar nail changes or other complaints was present. The patient was born of nonconsanguineous parents.

Examination revealed marked thickening and hardening with yellow-brown discoloration of all nail plates; fingernails were more thickened than toenails. The nail plates were uplifted by subungual hyperkeratosis with upward growth of the distal portion [Figure 1]. Hypercurvature on the transverse axis of all nail plates was present giving a pinched shape to the free edges. Focal palmoplantar keratoderma was present at pressure areas. She had oral leukokeratosis in the form of white diffuse opaque thickening of the tongue and buccal mucosa. Apart from this, there were hyperkeratotic follicular papules over extensors of the extremities and trunk. In addition, she had supernumerary digits on the ulnar aspect of her hands [Figure 2]. X-ray of hands did not reveal any bony element in the supernumerary digits. There was no history of supernumerary digits in the family. Hair, teeth, ophthalmological and otorhinolaryngological examinations were normal. Potassium hydroxide examination and fungal culture of nail plates and tongue scraping did not show any fungal element. Nail bed biopsy was refused. She was put

Letters to the Editor