Lessons from practice

An unexpected case of pellagra

Clinical record

An anxious 50-year-old woman presented with recurrence of a painful, well demarcated violaceous eruption with erythematous margins and large areas of blistering on her ankles and dorsa of her feet (Box 1, A). She had an erythematous desquamating rash on her hands, forearms and periorbital skin, with surrounding hyperpigmentation (Box 1, B). There were no signs of chronic liver disease, arterial insufficiency or sensory deficit. Oral antibiotics and high-dose oral prednisolone had been ineffective. She had no history of trauma or previous skin conditions. Sun exposure was an aggravating factor. She had recently started taking a herbal diuretic tablet and occasionally used ibuprofen and paracetamol for headaches. Her sister had an undifferentiated connective tissue disease and a positive antinuclear antibody level. The patient was a smoker and reported drinking half a bottle of gin, amounting to at least 10 standard alcoholic drinks, per day.

The differential diagnoses we considered were infection, pseudoporphyria, phototoxic reaction caused by the herbal diuretic tablet or ibuprofen, bullous lupus and porphyria cutanea tarda. Directed investigations (Box 2) did not provide a diagnosis. Mild elevation of some urinary and faecal porphyrins were consistent with liver dysfunction, rather than confirming porphyria. Histopathological examination showed paucicellular interface dermatitis with subepidermal blistering and epidermal pallor. Direct immunofluorescence for immunoglobulin and complement proteins was negative. All parameters that were outside reference intervals, except mean cell volume and γ-glutamyl transferase level, returned to normal within 2 weeks of initial management.

Initial supportive management, oral cephalexin 500 mg four times daily, and tapering prednisolone were ineffective. Further history-taking revealed poor nutritional intake, purging using docusate sodium tablets for laxative effect, and intermittent diarrhoea. The patient reported restricting her diet for weight control. She was underweight, with a body mass index of 17 kg/m². A scaly hyperpigmented collarette on her upper chest (Box 1, C) resembled Casal necklace, suggesting the clinical diagnosis of vitamin B₃ deficiency.

Administering oral nicotinamide 50 mg twice daily resulted in significant improvement in tenderness, burning discomfort and erythema within 2 days, and desquamation within 1 week. After 3 months of supplementation, only mild postinflammatory hyperpigmentation remained, subjective skin sensation had returned, mood had improved and diarrhoea had abated. The patient declined further investigations, including colonoscopy for other potential causes of malabsorption. The eventual resolution of her gastrointestinal symptoms suggested they were due to the disease process rather than a secondary cause such as colitis. She also declined referral to psychology, psychiatry or drug and alcohol counselling but accepted advice from a dietitian.

Pellagra is caused by a cellular deficiency of niacin or its precursor amino acid, tryptophan. Niacin, also known as nicotinic acid or vitamin B₃, is essential for carbohydrate, fat, protein and alcohol metabolism, detoxification of drugs and reactive oxygen species, cell signalling and DNA repair.¹ ²

Early gastrointestinal and neurological symptoms are usually subtle and non-specific but if untreated can progress to death from multiorgan failure. Therefore, mucocutaneous signs provide important diagnostic clues. An acute, symmetrical, well demarcated, tender erythematous eruption occurs at sites of sun exposure, heat, friction and pressure. Exacerbations involving the face, neck, chest and dorsa of extremities occur in spring and summer.¹ ³ The eruption can mimic sunburn with vesicles and bullae, termed wet pellagra.⁴ As in this patient, pemphigus pellagrosus with exudative bullae, pruritus and burning may occur on solar re-exposure.² In chronic

1 The patient’s skin lesions

A: Well demarcated violaceous eruption with erythematous margins and large areas of blistering. B: Erythematous desquamating rash, with surrounding hyperpigmentation. C: Scaly hyperpigmented collarette on her upper chest, resembling Casal necklace.
cases the skin is hyperpigmented and a cinnamon-brown colour. Hyperkeratosis and desquamation produce a dry, rough, scaly or cracked appearance.

Primary pellagra occurs in dietary deficiency of tryptophan or niacin. Secondary pellagra is caused by conditions characterised by interference with absorption or metabolism of tryptophan and niacin, including anorexia nervosa, chronic alcoholism, prolonged diarrhoea, ileitis, colitis, cirrhosis, carcinoid syndrome, Hartnup disease and HIV. Medicines including azathioprine, 5-flourouracil, 6-mercaptopurine and phenobarbitone inhibit the conversion of tryptophan to niacin. Isoniazid is an analogue of niacin and suppresses endogenous formation.1,2

Clinical diagnosis is confirmed by rapid response to oral nicotinamide when up to 500 mg daily in divided doses is given. Nicotinamide is the preferred supplement, as niacin causes flushing and headaches.2,3,5

Isolated cases of pellagra still occur in developed countries despite dietary fortification and the accessibility of niacin-rich foods. There are no known published reports from Australia. A retrospective study in a hospital population in Spain found the annual incidence was 0.5 cases per 100 000 population,3 and the most common risk factors for pellagra were alcoholism and poor diet. As in our patient, the diagnosis is often delayed, possibly owing to the rarity of pellagra.3,6

Factors contributing to alcohol-induced pellagra include poor dietary intake, niacin depletion due to alcohol metabolism, malabsorption due to the toxic effect of alcohol on the gut, and cirrhosis.5 Of note, patients with severe psychiatric and neurological forms of the disease such as pellagrous encephalopathy and pellagra psychosis may present without cutaneous signs. The incidence reported in the Spanish study may have underestimated true incidence, as their diagnostic criteria included dermatological manifestations.3 One study of 20 cases has suggested that lack of cutaneous involvement in chronic alcoholics with neuropsychiatric symptoms may be due to negligible sun exposure, both in social life and in an acute care setting.6 Severe neurological symptoms may progress so quickly that cutaneous signs do not have time to manifest.5 The important lesson from these reports is that niacin deficiency is common in chronic alcoholism, and pellagra should be considered a differential diagnosis of delirium tremens and psychosis, especially if signs of gait disturbance, limb rigidity and hyperreflexia are present.5-8

Associations with anorexia nervosa and medication interactions are also important. In patients with anorexia nervosa, signs of pellagra may be atypical or overlap with other nutritional deficiencies.9 These features may be an ominous sign, as demonstrated by a recent fatality.10 Further, isoniazid, azathioprine and antiepileptic drugs have been implicated recently.3,11,12 Our literature search revealed no reports of pellagra being caused by excessive use of docusate sodium, but this may have contributed to the disease in our patient, owing to malabsorption.

The purpose of this review is to draw attention to the presence of nutritional deficiencies such as pellagra in the developed world. Our patient did not present initially with obvious signs of poor self-care, alcoholism, anorexia nervosa or body-image issues. The diagnosis was only made after clinical suspicion led to a careful social and dietary history-taking and examination. Thus, awareness of the condition and of high-risk populations can prevent delayed diagnosis and treatment, with its associated morbidity and mortality.

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