Jellyfish envenomation: A chilling toxidrome of seizures and cyanosis – a case report

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Jellyfish envenomation is a common incident in coastal areas all over the world. While the majority of reported cases are self-limiting with few lasting complications, a few deadly species can cause life-threatening and debilitating illnesses with a prolonged recovery time. Chrysaora spp. have been known to cause a diverse spectrum of manifestations. We report the case of a 7-year-old boy with jellyfish envenomation presenting with cyanosis, seizures and hypertension not previously described in association with the Chrysaora spp.


Jellyfish are intriguing and are found ubiquitously in all oceans. However, species predominance is seen across different marine ecosystems. The most common species seen across coastal India are Aurelia and Physalia.1,2 Chrysaora spp. are widely dispersed across the Atlantic, Pacific and Indian oceans.2,3 Most Chrysaora spp. envenomations are mild, with irritation of the skin and pain, requiring outpatient medical attention.1,4 These species have rarely been reported to cause neurotoxicity or digital ischaemia, although they are an expected complication of the more toxic species.1,4 In this case report, we describe a rare case of a Chrysaora spp. sting associated with dermatitis, neurotoxicity and vasospasm leading to hypertension.

Case report
A 7-year-old boy of European origin who was on vacation in coastal Karnataka, India, was brought to triage with complaints of a jellyfish sting belonging to species Chrysaora while bathing in the sea, resulting in a rash over both his hands and legs which had been progressively increasing over 12 hours. The parents described the rash as spread out over the sites of attachment of the tentacles of the jellyfish which had to be forcibly separated from the boy. Swelling progressively increasing over 12 hours. The parents described the rash as spread out over the sites of attachment of the tentacles of the jellyfish which had to be forcibly separated from the boy. Swelling and diffuse subcutaneous oedema. The renal artery Doppler and 2D flow velocities and spectral waveforms with low biphasic resistance, all other pulses were well felt. SpO2-right upper limb was not recordable. Left upper limb: 98%; right lower limb: 96%; left lower limb: 98%. The third, fourth and fifth digits of the right hand were cyanosed and swollen, with painful movements up to the proximal interphalangeal joints including the nail beds (Fig. 1). An erythematous maculopapular rash was present on both thighs, legs and forearms, and flagellate pigmentation was present over both thighs. Purpuric lesions over the right thigh and popliteal fossa, with a local rise in temperature and extreme pain on movement were also seen (Fig. 2). The respiratory, abdominal and cardiovascular examinations were normal. Neurological examination at admission showed no focal neurological deficits.

The patient’s laboratory parameters were as follows: alanine transaminase 13.0 IU/L; aspartate transaminase, 45.0 IU/L; C-reactive protein, 1.4 mg/L; serum creatinine, 0.4 mg/dL; serum potassium, 4.6 mmol/L; sodium, 134.0 mmol/L; urea, 25 mg/dL; creatinine, 2.0 mg/dL; haemoglobin, 13.6 g/dL; haematocrit, 41.8%; mean corpuscular volume, 82.8 fl; platelet count, 299.0 × 10³/µL; red blood cell count, 5.06 × 10⁶/µL; RDW, 14.0%; total white blood cells, 15.8 × 10³/µL; serum calcium, 9.3 mg/dL; serum chloride, 93.8 mmol/L; magnesium, 2.0 mg/dL; phosphorus, 5.7 mg/dL; complement C3 level, 131.0 mg/dL; complement C4 level, 25 mg/dL; ESR, 13.6 g/dL; C-reactive protein, 1.4 mg/L; serum creatinine, 0.4 mg/dL; serum potassium, 4.6 mmol/L; sodium, 134.0 mmol/L; urea, 25 mg/dL; creatinine, 2.0 mg/dL; haemoglobin, 13.6 g/dL; haematocrit, 41.8%; mean corpuscular volume, 82.8 fl; platelet count, 299.0 × 10³/µL; red blood cell count, 5.06 × 10⁶/µL; RDW, 14.0%; total white blood cells, 15.8 × 10³/µL; serum calcium, 9.3 mg/dL; serum chloride, 93.8 mmol/L; magnesium, 2.0 mg/dL; phosphorus, 5.7 mg/dL; complement C3 level, 131.0 mg/dL; complement C4 level, 19.0 mg/dL; anti-nuclear antibody profile, negative; urine examination: normal.

The right upper limb arterial Doppler showed attenuation of the axillary, brachial, radial and ulnar arteries with a regular colour flow, flow velocities and spectral waveforms with low biphasic resistance and diffuse subcutaneous oedema. The renal artery Doppler and 2D
The echocardiograph were normal. An ultrasound demonstrated diffuse subcutaneous oedema with moderate knee-joint effusion with septation. Computed tomography angiography showed attenuation in the calibre of the distal two-thirds of the right brachial artery, radial and ulnar arteries and faint contrast opacification (Fig. 3).

Treatment was initiated with subcutaneous enoxaparin, intravenous dexamethasone, oral aspirin and oral cilostazol. On the third day following the sting, the patient developed five episodes of generalised tonic-clonic seizures which required levetiracetam (40 mg/kg/day) and valproate (10 mg/kg/day). Magnetic resonance imaging of the brain revealed patchy bilateral asymmetrical T2/FLAIR hyperintensities involving bilateral frontal-parietooccipital sulci in the cortex predominantly, with mild extension into the adjacent subcortical white matter and focal extension into the deep white matter (Fig. 4). Electroencephalography indicated a generalised disturbance of electrical function. The boy was found to be hypertensive on the fourth day, with a blood pressure reading of 140/100. The hypertension was controlled with oral nifedipine. The rash over the arms worsened over the course of five days and required therapy with topical antibiotics, antihistamines and steroids. The right knee effusion was treated conservatively.

The reappearance of the right radial pulse and reperfusion of the digits was observed after 7 days. He remained normotensive and seizure-free after the 8th day. However, a right-sided foot drop was noted on the 10th day, which responded minimally to therapy and rehabilitation. A residual reddish hue persisted as a remnant of the rash. He was normotensive and seizure-free on maintenance antiepileptics at discharge with a right foot drop.

**Discussion**

Jellyfish belong to the phylum **Cnidaria**, which has 10 000 species and only ~100 of those are toxic to humans.\(^7\) Although only a few species contain tentacles, nematocysts are found in all. The length of the nematocysts in the vast majority are insufficient to penetrate the dermis and are therefore harmless.\(^8\) Upon skin contact, the toxin produced by a number of species causes a local reaction involving the phospholipase A2 pathway, which leads to pain, swelling and skin necrosis in severe conditions.\(^9\) Once in the general circulation, the toxin has gastrointestinal, cardiac, neurological, muscular and immunogenic effects. The venom also contains haemolytic and lethal fractions affecting the heart.\(^10\)

Jellyfish stings most frequently result in pain, itching, an intense burning sensation and redness in the majority of cases. However, a few studies show that allergic reactions and anaphylaxis are remote complications as the jellyfish toxin is known to act by inducing the release of inflammatory mediators.\(^11\) The classical jellyfish dermatitis, also called seabather's eruption, is believed to occur due to the extended action of the jellyfish toxin leading to intense burning, pain, itching and erythematous papules for a prolonged time after the exposure.\(^11,12\) Synonymous with this, the patient in our case presented as described and responded to topical corticosteroids and antihistamines.

The venom of the *Chrysaora* spp. has been demonstrated to cause calcium ion influx through voltage-gated channels and cause severe arterial spasm in experimental studies.\(^13\) Jellyfish envenomation leads to acute regional vasospasm leading to cyanosis of the affected areas and absence of sweating and piloerrection.\(^14,15\) *Chrysaora* spp. causing these effects have been reported in rare cases across the Indian coast.\(^16\) Analogously, our patient developed a delayed response to the sting with a severe spasm of the right brachial artery, swelling, pain and cyanosis of the right third and fourth digits. Sympathomimetic action of the venom is known to cause severe hypertension in a syndrome commonly called the Irukandji, which is accompanied by severe cardio-toxic manifestations mimicking a catecholamine crisis.\(^14\) Hypertension, as described in these studies, was noted in our patient who was controlled with a direct arteriolar dilator such as cilostazol, a phosphodiesterase inhibitor, and nifedipine, a calcium channel blocker.

The toxin isolated from nematocysts of *Chrysaora* is known to depolarise nerve and muscle membranes and to increase the frequency...
of miniature end-plate potentials. Motor and sensory neurotoxic effects of cnidarian venoms have been reported in several cases of mononeuritis multiplex produced by different jellyfish. There have also been reports of Guillain–Barré syndrome and dysphonia following jellyfish stings. However, Chrysaora stings leading to neurological complications are seldom seen. Our patient developed seizures three days following exposure. Patchy bilateral hyper intensities were seen over the cortical areas on MRI. A point to consider here is that these changes may be due to a postictal state. However, after ten days he had a residual foot drop on the right side.

Treatment of jellyfish stings should begin by administration of basic life support. Appropriate methods for tentacle removal, such as flushing with sea water, using tweezers or scraping off with a card, should be done carefully as they carry the risk of further discharge of nematocysts. The application of vinegar and an ice pack to the local area to reduce pain is recommended. The affected part should then be immersed in water as hot as is bearable if no proprietary meds are available. Skin inhibitor creams are commercially available which provide protection for swimmers against jellyfish stings.

Conclusion

Jellyfish envenomation is usually harmless or causes mild illness. However, in rare instances, it can manifest as a toxidrome of seizures, cyanosis and hypertension due to prolonged and delayed multisystem effects of the toxin, requiring continuous intensive monitoring during the acute phase. Residual neurological deficits can be anticipated following envenomation. Adequate awareness by seabathers, personal protection, and prompt medical intervention is necessary to prevent morbidity.

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