Identification of Allergens in the Box Jellyfish *Chironex yamaguchii* That Cause Sting Dermatitis

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**Key Words**
Specific allergen · Box jellyfish dermatitis · *Chironex yamaguchii* · Nematocyst · Venom immune reaction

**Abstract**

**Background:** Jellyfish stings cause painful, papular-urticarial eruptions due to the immediate allergic, acute toxic and persistent inflammatory responses. In spite of many marine accidents and their economic impact, modes of first-aid treatment remain conventional and specific allergen and medical treatment are not yet available. The purpose of this study was to define the specific allergen of the box jellyfish *Chironex yamaguchii* and to study the precise mechanism of the resulting dermatitis.

**Methods:** We comprehensively studied the immunoglobulin-binding molecules from the box jellyfish *C. yamaguchii* with a purification procedure and Western blotting, using sera from 1 patient and from several controls.

**Results:** From the nematocyst wall and spine, we detected IgG-binding acidic glycoprotein (of 66 and 30 kDa) as determined by Western blot and ion-exchange chromatography. In addition, the 66-kDa protein was found to be an asparagine residue-coupled N-linked glycoprotein and the epitope resided in the protein fraction. We found that CqTX-A, the major toxic protein of the nematocyst, is also a heat-stable IgE-binding allergen. This was confirmed as a 45-kDa protein by Western blot from both nematocyst extracts and purified CqTX-A.

**Conclusions:** The detection of these proteins may, in part, explain the combined immediate allergic-toxic and persistent allergic responses. Hopefully, our findings will lead to the development of specific venom immunotherapy for marine professional workers and tourists for jellyfish-sting dermatitis and anaphylaxis.

**Introduction**

The box jellyfish *Chironex yamaguchii* (also known as *Chiropsalmus quadrigatus*) inhabits the tropical Indo-Pacific region [1]. Envenomation of *C. yamaguchii* occurs in coastal areas from June to September, especially in Okinawa in the tropical region of Japan. The species is distinguished by the cube-shaped medusa, and is classified into Cubozoa (a so-called box jellyfish or sea wasp) of cnidarian. Box jellyfish stings, including those from *C. yamaguchii*, *Chironex fleckeri* and *Carukia barnesi* as well as the hydrozoan jellyfish *Physalia physalis* (Portuguese man-of-war), can be fatal [2, 3]. Contact with the jellyfish triggers the nematocysts (harpoon-shaped microscopic...
mechanisms) that exist on the tentacles to inject venom into the victim’s skin. A worldwide epidemiological survey recorded that there are 150 million cases of jellyfish stings a year [4]. In Okinawa, 103/132 jellyfish stings officially reported in 2013 were caused by C. yamaguchii [5].

Jellyfish stings cause painful, papular-urticarial eruptions due to immediate allergic, acute toxic and persistent inflammatory responses [6]. The most characteristic signs and symptoms are painful skin eruptions at the sting site. These acute lesions usually last for minutes to hours, and the rash is combined with urticaria. Further progression can lead to vesicles, hemorrhaging or necrotizing lesions. Delayed reactions such as pruritic urticarial lesions may relapse after 2–4 weeks [7]. Systemic reactions may occur and these often result in cardiovascular and respiratory failure, probably due to a combination of toxicity and anaphylaxis. It was assumed that these are allergic reactions, but the nature of the allergen(s) has not yet been elucidated [8]. In spite of the many accidents and the risk to fishery employees and marine tourists, no definite consensus exists for effective first-aid, clinical management or specific medical treatment.

Cnidarians have countless numbers of nematocysts on the surface of their tentacles. The nematocyst is specialized to inject venom into the victim. It is filled with other components such as protein toxins, inflammatory molecules, polypeptides and enzymes [9, 10]. It also has a mechanical apparatus that encloses a folding spine, like a harpoon. The nematocyst capsule wall and spine (NCS) are made of hard tissues, formed by the strong binding of minicollagen and glycoprotein [11, 12]. The nematocyst injects venom in response to physical and/or chemical stimulation [13]. A major protein toxin, CqTX-A, with a molecular weight of 45 kDa has already been isolated from the venom of C. yamaguchii and the deduced amino acid sequence clarified [14]. CqTX-A exhibits potent hemolytic activity and crayfish-lethal toxicity [14]. A small amount of cytotoxic protein (43 kDa) was found in the venom [15].

Recently, many allergens from marine organisms have been reported, such as in sea urchins and red soft coral [16, 17]. The purpose of this study was to define the specific allergen of the box jellyfish C. yamaguchii and the precise mechanism of the resulting dermatitis. From our comprehensive investigation, starting with this species, we detected the IgG-binding acidic glycoprotein in the NCS, and identified CqTX-A, in the nematocyst, as the critical allergen and main toxic protein.

Materials and Methods

Collection of C. yamaguchii
The box jellyfish C. yamaguchii was captured and collected at Ginowan, Okinawa, Japan in the month of August. The tentacles were excised from the live specimen immediately after capture and frozen at −30°C. The remaining body was discarded. The purification process is charted in figure 1a.

Preparation of Crude Extracts from Tentacles
The frozen tentacles of C. yamaguchii were excised in 10 mM PBS (pH 7.0) using scissors, and were then sonicated with an ultrasonicator MUS-400 (EYELA, Tokyo Rikakikai Co., Tokyo, Japan). The obtained suspension was centrifuged at 10,000 g for 5 min. The supernatant was filtered with a 0.45-μm-disk filter (ADVANTEC, Toyo Roshi Kaisha, Tokyo, Japan). The filtrate was defined as the tentacle extracts (fig. 1a).
**Preparation of Isolated Nematocysts**

Nematocysts (wet weight: 15 g) were isolated from the frozen tentacles of *C. yamaguchii* (fig. 1a) as described previously [18]. The frozen tentacles (wet weight: 100 g) were autolyzed in 1.0 M sodium citrate solution (200 ml) for 5 days at 4 °C. The autolyzed suspension was filtered through fine-mesh nylon netting. The filtrate was allowed to settle in a beaker for 3 h at 4°C. The supernatant was then removed by decanting and the nematocysts were washed with 1.0 M of sodium citrate solution for isolation.

**Preparation of Nematocyst Extracts**

Nematocyst extracts were prepared by bead mill homogenization as described by Carrette and Seymour [19]. The nematocysts were washed with 10 mM phosphate buffer (pH 6.0), and then suspended in the same buffer and ruptured using a Mini-Beadbeater (BioSpec Products, Bartlesville, Okla., USA) and 0.5-mm glass beads. Disruption of the nematocyst was monitored microscopicly. The suspension was then centrifuged at 10,000 g for 5 min and the supernatant was filtered through a 0.45-μm-disk filter. The filtrates were considered as nematocyst extracts (fig. 1a).

**Isolation of CqTX-A from *C. yamaguchii***

The isolation of CqTX-A, a main protein toxin of *C. yamaguchii*, was conducted as described [14]. Nematocyst extracts (7.0 g) were put on a cation-exchange HPLC, TSK-GEL CM-5PW column (7.5 × 75 mm, 0.5 ml/min; Tosoh, Tokyo, Japan) which had been equilibrated with 10 mM phosphate buffer containing 0.3 M NaCl (pH 6.0), and the column was then washed with 10 mM phosphate buffer (pH 6.0). The adsorbed components, which included CqTX-A, were eluted with a gradient solvent system (0–0.7 M NaCl for 0–90 min). The toxicity was checked by means of a hemolytic activity test as described below. The column-adsorbed toxic fraction was concentrated using a cation-exchange HPLC, TSK-GEL CM-5PW column [14]. The concentrated active fraction was applied to a gel permeation chromatography, which had been equilibrated with 10 mM phosphate buffer containing 0.3 M NaCl (pH 6.0). The column was then washed with 10 mM phosphate buffer (pH 7.5). The column was then washed with 10 mM phosphate buffer (pH 7.5). The adsorbed fraction was eluted with 10 mM phosphate buffer containing 1.0 M NaCl (pH 7.5) without a concentration gradient, in order to get a higher yield. For cation-exchange chromatography, the same methods were applied as for anion-exchange except that a HiTrap-CM (GE Healthcare) column and 10 mM of phosphate buffer (pH 6.0) were used instead. Adsorbed and nonadsorbed fractions were obtained from both anion- and cation-exchange chromatography, followed by SDS-PAGE and Western blotting.

**Western Blot**

Sample proteins were analyzed by 10% SDS-PAGE under a condition of reducing by 2-mercaptoethanol or a nonreducing condition. Proteins in the gels were transferred onto a PVDF membrane (Bio-Rad) in a Mini Trans-Blot (Bio-Rad) for 4 h at 60 V. After transfer, the PVDF membrane was blocked with 2.5% skim milk in Tris-buffered saline (pH 7.6) containing 0.1% Tween 20 (TBS-T) for 1 h at room temperature. The membrane was incubated for 1 h at room temperature with primary antibody (1:2,500 dilutions) in TBS-T. After washing 4 times with TBS-T, the membrane was reacted with HRP-conjugated secondary antibody in TBS-T containing 1% skim milk for 1 h at room temperature. After washing 2 times, immunoreactive proteins were detected with ECL Plus (GE Healthcare). Serum from the patient (with written informed consent) was taken by the medical doctor (S.K.), and stored at −30 °C to be used as a primary antibody. The serum from a patient who was assumed to not have had any contact with *C. yamaguchii* was used as a negative control. HRP-conjugated goat anti-human IgG (1:50,000 dilutions, polyclonal; Zymed®/Invitrogen™, Thermo Fisher Scientific), HRP-conjugated mouse anti-human IgG1 (1:10,000 dilutions, monoclonal; Zymed/Invitrogen), HRP-conjugated mouse anti-human IgG2 (1:10,000 dilutions, monoclonal; Zymed/Invitrogen) and HRP-conjugated goat anti-human IgE (1:20,000 dilutions, polyclonal; Kirkegaard & Perry Lab., Gaithersburg, Wash., USA) were used as secondary antibodies.

**Glycosylation of IgG-Binding Allergen Protein**

The NCS extracts were deglycosylated to examine whether they contained amino acid-linked sugar residues of N-linked or O-linked oligosaccharides. The extent of deglycosylation was assessed by mobility on SDS-PAGE using the enzymatic protein deglycosylation kit (Sigma-Aldrich, St. Louis, Mo., USA) according to the manufacturer’s instructions. N-linked sugars were removed by digestion with the enzyme PNGase F [peptide-N4-(acetyl-β-glucosaminyl)-asparagine amidase] and O-linked sugars were removed by using

Identification of Specific Allergens in *C. yamaguchii* Lab., Tokyo, Japan) in 10 mM PBS at 37°C for 2 h. Cell suspensions were centrifuged at 2,000 g for 10 min. The concentration of hemoglobin released into supernatant was measured at 570 nm with a microtiter plate reader (Model 550; Bio-Rad Lab., Hercules, Calif., USA). In this assay, 1 unit was defined as the dose which caused 50% hemolysis.

**Ion-Exchange Chromatography on NCS Extracts**

To characterize NCS extracts, ion-exchange chromatography using an anion or cation column was performed. For anion-exchange chromatography, NCS extracts (20 mg) were loaded to HiTrap-DEAE (GE Healthcare) chromatography, which had been equilibrated with 10 mM of phosphate buffer containing 0.3 M NaCl (pH 7.5). The column was then washed with 10 mM phosphate buffer (pH 7.5). The adsorbed fraction was eluted with 10 mM phosphate buffer containing 1.0 M NaCl (pH 7.5) without a concentration gradient, in order to get a higher yield. For cation-exchange chromatography, the same methods were applied as for anion-exchange except that a HiTrap-CM (GE Healthcare) column and 10 mM of phosphate buffer (pH 6.0) were used instead. Adsorbed and nonadsorbed fractions were obtained from both anion- and cation-exchange chromatography, followed by SDS-PAGE and Western blotting.
the enzymes, glycosidase (endo-α-N-acetylgalactosaminidase) and α-2 (3, 6, 8, 9)-neuraminidase, respectively.

**Skin-Prick Test**

The doctor put a drop each of solutions containing intact and thermally deactivated CqTX-A on the patient’s forearm using a sterilized needle, and then pricked the skin to allow the toxin to enter beneath the epidermis for 20 min in order for a reaction to appear. PBS was used as a negative control.

**Skin Eruption from the Jellyfish Stings**

A 43-year-old male patient, who had a history of repeated sting injury by jellyfish over several years, visited our health clinic for skin eruptions after being stung by jellyfish during maritime fieldwork in the Okinawa islands. His immunoglobulin profiles were IgE: 460 IU/ml (high), IgG: 927 mg/dl, IgA: 269 mg/dl and IgM: 40 mg/dl. He was sensitized to the house-dust mite, *Derma-tophagoides pteronyssinus*. His C-reactive protein was negative, white blood cell count 4,600/μl and hemoglobin 14.3 g/dl. The pain and papular eruption appeared immediately after the sting injury and the prick application, likely indicating a toxic reaction (fig. 2a, b). Erythema, wheal formation, induration and itching followed

**Lymphocyte Transformation Test**

A lymphocyte transformation test was done using heparinized blood followed by the isolation culture. The stimulation by the tentacle extracts was monitored by means of 3H-thymidine incorporation.

**Results**

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15 min later as allergic reactions (fig. 2c). One hour later, the erythema and wheal persisted and fused with each other, in spite of a decrease in pain and the hot sensation (fig. 2d). After 24 h, the papules and pigmentation were still conspicuous (fig. 2e). The skin eruption subsided gradually after this, but the erythema relapsed after 7 days.

**Comprehensive Examination of the Reactive Protein from C. yamaguchii**

For detection of IgG-binding protein molecules in C. yamaguchii (fig. 1a), each sample, i.e. the extracts of the tentacle, the nematocyst and the NCS, was subjected to Western blotting with the patient’s serum after the sting and negative control serum, and secondary antibodies: HRP-conjugated goat anti-human IgE (lanes 1 and 5), HRP-conjugated goat anti-human IgG (lanes 2 and 6), HRP-conjugated mouse anti-human IgG1 (lanes 3 and 7) and HRP-conjugated mouse anti-human IgG4 (lanes 4 and 8). When using the patient’s serum, a band of 66 kDa was detected with IgE, IgG and IgG4 (lanes 1, 2 and 4) and a band of 30 kDa was detected with IgE, IgG1 and IgG4 (lanes 1, 3 and 4). When using the negative control serum, faint bands of 66- and 30-kDa proteins were detected in IgE and IgG4 (lanes 5 and 8) and no band was detected in IgG and IgG1 (lanes 6 and 7).

**Isotype Specificity of the Reactive Protein in the NCS Extracts**

To determine isotype specificity to the protein, the NCS extracts were subjected to SDS-PAGE followed by Western blotting with human IgE, IgG, IgG1 and IgG4. The 66- and 30-kDa proteins reacted with HRP-conjugated goat anti-human IgE (fig. 4: lane 1). The 66-kDa protein reacted with HRP-conjugated goat anti-human IgG (fig. 4: lane 2) similarly to in figure 3. The 30-kDa protein reacted with HRP-conjugated mouse anti-human IgG1 (fig. 4: lane 3). The 66- and 30-kDa proteins also reacted with HRP-conjugated mouse anti-human IgG4 (fig. 4: lane 4). The 30-kDa protein was not detected under the nonreducing condition. In the serum from the negative control, faint bands of the 66- and 30-kDa proteins were detected in IgE and IgG4, but no band was detected in IgG1 (fig. 4: lanes 5–8).

**Characterization of the Protein in the NCS Extracts by Ion-Exchange Chromatography**

To characterize IgG-binding protein, NCS extracts were applied to cation-exchange HPLC (Hitrap-CM) and
anion-exchange HPLC (Hitrap-DEAE). On Hitrap-CM chromatography, we found that the 30-kDa protein in the nonabsorbed fraction was detected by Western blot with the patient’s serum and HRP-conjugated goat anti-human IgG (fig. 5a). Similarly, on Hitrap-DEAE chromatography, the 30-kDa protein was detected in the absorbed fraction (fig. 5b: lanes 9, 10). The 66-kDa protein was not detected in any fractions. These results revealed that the IgG-binding protein was an acidic protein.

Glycosylation of Specific IgG-Binding Protein
To determine the glycosylation of IgG-binding protein in the NCS extracts, we analyzed the mobility shifts of IgG-binding protein by digestion with PNGase F, O-glycosidase and α-2 (3,6,8,9)-neuraminidase. The validity of these enzymes was confirmed using bovine fetuin, which contains sialylated N-linked and O-linked oligosaccharides (data not shown). Samples without enzyme digestion were used as a control. Samples and control were incubated for 3 h at 37°C, and their mobility shifts were then analyzed by SDS-PAGE. PNGase F decreased the protein into a smaller amount of molecular weight (fig. 6: lanes 1,2,4), whereas O-glycosidase and α-2 (3,6,8,9)-neuraminidase did not affect the mobility of the allergen when compared with the control (fig. 6: lanes C, 3). These results indicated that the protein was asparagine residue-coupled N-linked glycoprotein and that the epitope resided not in the glycoside but in the protein fraction.

Identification of the Allergen in the Nematocyst Extracts
The nematocyst extracts were subjected to Western blotting with HRP-conjugated goat anti-human IgE, HRP-conjugated goat anti-human IgG, HRP-conjugated goat anti-human IgG. An IgG-binding allergen was detected at 30 kDa in the Hitrap-CM nonabsorbed fraction and in the Hitrap-DEAE fractions 9 and 10. Major proteins of NCS extracts were detected in the Hitrap-DEAE fractions 8–16 and in the Hitrap-CM nonabsorbed fraction by silver-staining SDS-PAGE. N = Nonabsorbed fraction.
mouse anti-human IgG1 and HRP-conjugated mouse anti-human IgG4. The result shows that these extracts had a main protein around 45 kDa and a minor protein in 43 kDa (fig. 7: left). The main protein reacted with IgE and IgG4 (fig. 7: lanes 1, 3). Similarly, the minor protein (43 kDa) reacted with IgE (fig. 7: lane 1). We did not find any bands that reacted with IgG1 (fig. 7: lane 2). In the negative control serum, no bands were detected in IgG1 and a faint band of 45-kDa protein was detected in IgE and IgG4 (fig. 7: lanes 4–6).

Antigen-Antibody Reaction of CqTX-A with Human IgE

From the above findings (fig. 7), we assumed that the IgE-binding protein of the nematocyst extracts was the main protein toxin CqTX-A, identified in a previous study [14]. We isolated CqTX-A (19.7 mg) based on the hemolytic activity (122,880 units) and detected it in SDS-PAGE (45 kDa; fig. 8: left). The results of Western blotting showed that IgE-binding CqTX-A was detected at the position identical to a major band of nematocyst extracts (fig. 8: lane 1 vs. lane 2). In addition, CqTX-A also reacted with IgG4 (fig. 8: lane 3). No band was detected in anti-IgE (fig. 8: lane 4) or anti-IgG4 (fig. 8: lane 5) in the serum from the negative control. In the skin-prick test, both intact and thermally deactivated CqTX-A caused erythematous skin eruption in the patient; the mean diameters of induration and erythema were 8 × 5 and 11 × 8 mm for intact CqTX-A, 6 × 6 and 12 × 18 mm for thermally deactivated CqTX-A and 5 × 4 and 6 × 6 mm for PBS, respectively. Thus, the IgE epitope of CqTX-A is a heat-stable protein and it does not change its conformation, allowing IgE to recognize the molecule after heating.

### Discussion

Common signs of jellyfish stings include pain, skin eruptions, itching, swelling or numbness. The stings of many species of box jellyfish, i.e. *C. yamaguchii*, *C. fleckeri* and *C. barnesi* and also of the hydrozoan jellyfish *P. physalis* can be fatal [2, 3].

In this study, we comprehensively investigated the specific allergens in the box jellyfish *C. yamaguchii*, which was separated into 3 parts, the tentacles, the nematocyst and the NCS, in order to study the allergen (fig. 1a). We detected an IgG-binding protein and an IgE-binding allergen in *C. yamaguchii*.

The major protein toxin with a molecular weight of 45 kDa, CqTX-A, in the nematocyst, was found to be an IgE- and an IgG4-binding allergen. Nagai et al. [14] reported that they isolated this main protein toxin of *C. yamaguchii* from the venom obtained from the nematocyst and also deduced its amino acid sequence. They also isolated *Carybdea arborifera* (formerly *Carybdea alata*) protein toxins (CaTX-A, CaTX-B) and *Carybdea brevipedalia* (formerly *Carybdea rastoni*) protein toxins (CrTX-A, CrTX-B) and designated them the ‘jellyfish toxin family’ [21, 22]. *C. fleckeri* protein toxins (CfTX-1, CfTX-2) show 89 and 72% sequence homology to CqTX-A by amino acid sequence alignment [23]. These box jellyfishes are well known to cause injury to fishermen, fisherwomen, scuba divers, surfers and sea-bathers in Japan, Hawaii and Australia.

Dermatitis caused by jellyfish stings can be characterized as 3 types of immune reaction, i.e. immediate allergic, late-phase reaction and relapsing allergic response, which correspond to clinical reactions [6]. The pathogenesis caused by the jellyfish stings includes at least both jellyfish toxic effects and a significant immediate allergic response. Intact CqTX-As and thermally deactivated
CqTX-As were used as allergens for the prick test. The test showed only an immediate allergic reaction, i.e. itching and erythema. The lymphocyte transformation test on the tentacle extracts showed a stimulation index of 0.87 (subnormal range) compared with >139 with phytohemagglutinin. This result indicated the absence of a T cell-mediated reaction. Taken together, it was shown that CqTX-A caused a mixed venom immune reaction, although the precise mechanisms remain unknown. In particular, it is not known whether IgG (IgG4) is involved in the late-phase reaction or if it reflects the repeated sting injury [24, 25]. IgG1 has been found to be involved in systemic anaphylaxis in a mouse model [26] and in the late asthmatic response in an epidemiological survey [27]. It has also been recognized in canine mast cell activations [25]. The pathogenesis caused by the venom from wasp and bee stings and that of aspergillosis or anisakis disease cannot be explained by allergic or toxic reactions alone. In fact, both these types of reactions occur simultaneously or sequentially and modify the symptoms and signs in the subacute and chronic phase. Bee venom contains melittin, histamine and phospholipase A, causing toxic effects along with an IgE reaction in the second sensitization [28]. The clinical progression of allergic bronchopulmonary aspergillosis and aspergilloma, however, is interwoven, and could worsen to chronic pulmonary aspergillosis in the case of infection by fungi of the genus *Aspergillus*.
The IgG- and IgG_1-binding proteins that we detected were a 66- and a 30-kDa protein from the NCS; the 66-kDa protein possibly resulted from the dimerization of the 30-kDa protein. It has been suggested that a minicollagen is the main component of the NCS [29], so this glycoprotein reacting with IgG might be a minicollagen in the nematocyst. Nematocysts are toxin-containing capsules with apparatus for catching prey, and are located in the jellyfish tentacles (fig. 1b). They have a ‘barb’ in the spine and discharged toxins can reside in a victim’s skin tissue for a long time [13]. Thus, a minicollagen in the wall of the nematocyst could cause IgG_1 and IgG reactions that trigger anaphylaxis, late-phase allergic reactions or a delayed hypersensitive reaction.

First-aid for jellyfish stings includes: (1) the inhibition of discharge of unfired adherent nematocysts by applying vinegar, (2) the removal of the tentacle and (3) applying ice to the skin to control the pain. Victims must be treated depending on their symptoms and the size of the affected area. Current clinical treatment is palliative and according to symptoms, i.e. with topical steroids and anti-inflammatory drugs [30]. To date, no effective treatment based on the evidence exists for stings. Recently, the Cochrane Collaboration reviewed interventions for stings of both adults and children, and did not produce any significant evidence for the efficacy of applying vinegar or ice [31].

In 1902, Portier and Richet [32], who had started studies on cnidarians, injected dogs with sea anemone venom twice, and expected a protective immune response (or prophylaxis), but found the opposite: ‘the hypersensitive reaction anaphylaxis’. From an occupational and hygienic point of view, fishery employees suffer anaphylaxis and allergic symptoms from cnidian injuries and beekeepers can become allergic to bee stings. In the case of beekeepers, hyposensitization therapy using allergens has already been developed as a preventive method [33]. Antivenom antibody was developed against box jellyfish venom and has been used over the last 30 years with limited efficacy [34].

It is plausible that CqTX-A has cross-reactivity with CaTX-A and CaTX-B, CrTX-A and CrTX-B and CfTX-1 and CfTX-2 because of their amino acid sequence similarities. Nagai et al. [35] and Iguchi et al. [36] managed to isolate other cnidian protein toxins, i.e. in the sea anemone Phyllophora semoni and the fire coral Millepora sp.; these cause a similar immune reaction in the skin. If we can reveal the IgG epitope in cnidian nematocyst minicollagen or the IgE epitope in the family of versatile toxins, more effective treatments will be made possible, such as hyposensitization therapy for relapsing allergic reactions and specific anaphylaxis (venom immunotherapy) from cnidian injury for marine professionals and tourists.

In our study, we found a 45-kDa IgE-binding allergen to CqTX-A and 66- and 30-kDa IgG-binding proteins in the NCS extracts. The detection of these proteins does, in part, explain the combined immediate allergic-toxic and persistent inflammatory response; this will hopefully lead to the development of specific and effective clinical applications for marine-sting dermatitis.

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Disclosure Statement

The authors declare no conflicts of interest.

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