Heavy Metals Affect Nematocysts Discharge Response and Biological Activity of Crude Venom in the Jellyfish *Pelagia noctiluca* (Cnidaria, Scyphozoa)

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**Key Words**

Heavy metals • Nematocysts • Discharge • Crude venom • Jellyfish • *Pelagia noctiluca*

**Abstract**

*Background:* Pollution of marine ecosystems and, specifically, heavy metals contamination may compromise the physiology of marine animals with events occurring on a cellular and molecular level. The present study focuses on the effect of short-term exposure to heavy metals like Zinc, Cadmium, Cobalt and Lanthanum (2–10 mM) on the homeostasis of *Pelagia noctiluca* (Cnidaria, Scyphozoa), a jellyfish abundant in the Mediterranean sea. This species possesses stinging organoids, termed nematocysts, whose discharge and concomitant delivery of venom underlie the survival of all Cnidaria. *Methods:* Nematocysts discharge response, elicited by combined chemico-physical stimulation, was verified on excised oral arms exposed to heavy metals for 20 min. In addition, the hemolytic activity of toxins, contained in the crude venom extracted from nematocysts isolated from oral arms, was tested on human erythrocytes, in the presence of heavy metals or their mixture. *Results:* Treatment with heavy metals significantly inhibited both nematocysts discharge response and hemolytic activity of crude venom, in a dose-dependent manner, not involving oxidative events, that was irreversible in the case of Lanthanum. *Conclusion:* Our findings show that the homeostasis of *Pelagia noctiluca*, in terms of nematocysts discharge capability and effectiveness of venom toxins, is dramatically and rapidly compromised by heavy metals and confirm that this jellyfish is eligible as a model for ecotoxicological investigations.
Introduction

Cnidaria, representing the first stage of Metazoan evolution, comprise different classes, such as Anthozoa, Scyphozoa, Hydrozoa and Cubozoa, all worldwide distributed and possessing highly specialized stinging cells, referred to as nematocytes. These cells produce an organoid, the nematocyst, which occupies most of the cytoplasm and consists of a large capsular structure with a spiny tubule inside, immersed in the crude venom - a fluid matrix containing toxins [1, 2]. When an appropriate chemico-physical stimulus is applied, both tubule and toxins are promptly delivered via an exocytic process termed discharge. Discharge is one of the most rapid biological responses known so far, whose activation mechanism has been focused on by many authors [2-5]. The first stage for discharge activation is chemosensitization, occurring via binding of exogenous compounds to chemoreceptors located on cells, other than nematocytes, modulating in turn the mechanosensitive apparatus (cilia). Contact with the prey and release of substances from its tissues further activates the process [6-9]. Therefore, discharge is reasonably acknowledged as the physiologic reaction responsible for Cnidaria survival through the ages, since it is used for either prey capture or defensive and locomotory functions [10, 11].

As already mentioned, discharge response is strictly associated with toxins release, being the crude venom delivered as a consequence of the ejection of the inverted tubule contained in the nematocyst. Crude venom from different Cnidaria specimens is reported to contain diverse compounds showing hemolytic, cytolytic, clastogenic, enzymatic, cardiotoxic, neurotoxic and even insecticidal effects, as revealed by different biological assays [12-18]. However, the current understanding on Cnidaria venom composition and biological activity is still incomplete for many species.

With regard to discharge response and biological activity of crude venom in the jellyfish Pelagia noctiluca, several investigations added novel contribution to the understanding of the physiology of this Cnidarian [4, 5, 19-27]. The current knowledge of Pelagia noctiluca biology [17, 28] and physiology [5, 29, 30] may allow to propose this specimen and its homeostatic features as new models to monitor the possible presence and ecological impact of marine pollutants, such as hydrocarbons and heavy metals, and ocean acidification. Concerning this latter phenomenon, it has been recently proven that acidification of the external medium significantly impairs discharge response and cell volume regulation capability following hypotonic shock (regulatory volume decrease, RVD) [20] of Pelagia noctiluca nematocytes. Moreover, in the same species, RVD is also inhibited by heavy metals such as cadmium, zinc and cobalt [19].

Pelagia noctiluca specimens are particularly abundant in the Mediterranean sea [31], especially in the Strait of Messina, an exemplary habitat where anthropic factors may critically contribute to pollution, including heavy metals contamination. Therefore, the aim of the present investigation was to verify the effect of heavy metals on Pelagia noctiluca homeostasis, with specific regard to discharge response and hemolytic power of crude venom. With this purpose, holotrichous isorhiza [32] nematocysts discharge has been studied on isolated oral arms, according to the Thorington and Hessinger [33] and Morabito et al. [5] technique. The biologic activity of toxins contained in the crude venom was assessed via an hemolytic assay on human erythrocytes, according to Marino et al. [21, 22].

Materials and Methods

Specimens collection

Specimens of Pelagia noctiluca (Cnidaria, Scyphozoa) were collected during Summer 2013 in the Strait of Messina (Italy) and immediately used for experiments. Oral arms were utilized for experiments evaluating nematocysts discharge and crude venom extraction.

In situ discharge

Segments of oral arms, approximately of the same size, were excised with ophthalmic scissors and collected by fire-polished silicon-coated Pasteur pipettes. Segments were then rinsed with low-Ca²⁺ artificial
sea water (ASW, in mM: NaCl 520, KCl 9.7, CaCl₂ 0.01, MgCl₂ 24, MgSO₄ 28, imidazole 5, pH=7.65, 1100 mOsm/Kg) to remove mucus. Tissues were then transferred into 5 ml Petri dishes previously treated with Sylgard (Dow Corning), fixed with Opuntia spines and repeatedly rinsed with standard ASW (in mM: NaCl 520, KCl 9.7, CaCl₂ 10, MgCl₂ 24, MgSO₄ 28, imidazole 5, pH 7.65; 1100 mOsm/Kg). Oral arms were carefully checked under a light microscope (100x magnification) to assess their structural integrity and then submitted to a combined chemical-mechanical stimulation, as described in the following.

Control experiments. Nematocysts discharge was elicited in situ by combined chemical-mechanical stimulation of oral arms, according to Thorington and Hessinger [33] and Morabito et al. [5]. Glutamate was chosen as the chemosensitizer compound [5]. The chemical-mechanical stimulation consisted in a 20 min incubation of the oral arms in 5 ml standard ASW plus 10⁻³ M glutamate (chemosensitization), followed by mechanical stimulation with a gelatine-coated non-vibrating test probe. Test probes consisted of 2 cm segments of 0.8±0.01 mm diameter nylon fishing line coated at one end with 30% (w/v) gelatine with a thickness of approximately 0.06 mm. After storage for 24 h at 4 °C and 100% humidity, the probes were inserted, at the uncoated end, into glass capillary tubes and fixed to a micromanipulator (Leitz). Test probes were then moved into the tissue by careful checking with an inverted microscope (Zeiss, 200x magnification). This manoeuvre induced nematocysts discharge and allowed for the adhesion of fired capsules to the gelatine. The individual gelatine-coated probes, possibly bearing fired nematocysts, were placed in separate microtiter wells (Microtest 11, Falcon Plastics) each containing 50 µL of 1% enzyme/detergent mixture (Trizyme; Amway Products, Ada, MI, USA). After a 4 h incubation at room temperature the gelatine was completely dissolved and probes were discarded. Discharged nematocysts were counted by inspecting with an inverted microscope (Zeiss, 400x magnification) and their number (nematocysts/probe) was considered as an index for discharge response.

Evaluation of the effect of heavy metals on nematocysts discharge. To test for a possible modulation of in situ discharge following exposure to heavy metals, oral arms were incubated for 20 min in 5 ml standard ASW containing 2 mM ZnCl₂, CdSO₄, CoCl₂ or La(NO₃)₃. After incubation, the ASW containing heavy metals was removed and chemical-mechanical stimulation immediately followed as previously described. To exclude a possible direct discharge-inducing effect of heavy metals on oral arms, the latter were exposed to heavy metals and then mechanically stimulated by the non-vibrating test probe, without previous chemosensitization with glutamate. Discharge response was then evaluated as reported previously.

Nematocysts isolation and crude venom extraction

In order to perform the hemolytic assay, it was firstly necessary to isolate nematocysts from oral arms of the collected jellyfish [29] and then extract the crude venom [21].

The oral arms were excised and submerged in distilled water for 2 h at 4 °C. In these conditions, undischarged nematocysts are released following osmotic lysis of nematocytes. After a complete detachment of the epidermis, the underlying tissue was removed from the suspension containing both the epidermis and nematocysts. Further release of nematocysts still attached to the epidermal tissue was induced by stirring. The nematocysts were repeatedly washed in distilled water and filtered through plankton nets (100, 60 and 40 µm mesh nets were used in this order) to remove most of the tissue debris, and then centrifuged at 4 °C (ALC PK 120R centrifuge, 4000 g for 5 min). The suspension was immediately used or, alternatively, frozen at -20 °C until use. The nematocysts obtained with this method were classified as holotrichous isorhizas, according to Morabito et al. [32].

The nematocysts were defrosted, filtered and washed again in distilled water before each experiment. Nematocysts were then re-suspended in 0.01 M phosphate buffer containing 0.9% NaCl (pH 7.4, 300 mOsm/Kg). Crude venom was extracted by sonicating (Sonopuls, 70 mHz, 30 times, 20 s) on ice a suspension of 90 nematocysts/µL. After sonication, the crude venom extract was separated from crushed capsules by centrifugation (4 °C, 4000 g, 10 min) and used for the hemolytic assay.

Hemolytic assay

Control experiments. After obtaining informed consent from human healthy volunteers, erythrocytes were prepared by washing fresh human whole blood thrice with 0.01 M phosphate buffer containing 0.9% NaCl, pH=7.4, 300 mOsm/kg. The erythrocyte suspension (0.05% in 0.01 M phosphate buffer containing 0.9% NaCl, pH=7.4, 300 mOsm/kg) was incubated with 5%, 10% or 20% v/v crude venom at 37 °C for 1 h and then centrifuged at 2000 g for 5 min to precipitate both intact erythrocytes and ghosts. Aliquots of the
supernatants were then taken and the optical density at $\lambda=414$ nm was spectrophotometrically determined to assess the amount of hemoglobin released from the lysed erythrocytes. The lytic power of the venom was expressed as % absorbance compared to that observed after maximal lysis of erythrocytes suspended in distilled water. The supernatant of an untreated 0.05% erythrocytes suspension in isotonic buffer was taken as the negative control.

**Evaluation of the effect of heavy metals and their mixture on the hemolytic power of crude venom.** The effect of heavy metals on the hemolytic power of crude venom was determined after simultaneously adding either 2, 5 or 10 mM $\text{ZnCl}_2$, $\text{CdSO}_4$, $\text{CoCl}_2$, or $\text{La(NO}_3)_3$ and 10% or 20% v/v crude venom to 0.05% erythrocytes suspensions for 1 h. The 10% or 20% v/v crude venom concentrations were chosen as they were particularly effective in inducing hemolysis in control experiments. A similar protocol was applied to test the effect of the heavy metals mixtures on the hemolytic power of crude venom (total concentration: 1-10 mM; the individual heavy metals were present in equimolar amounts). The hemolytic activity of the crude venom was then assayed as described above. To exclude a possible direct hemolytic effect of heavy metals or their mixture, they were prior tested on 0.05% erythrocytes suspensions, without exposure to crude venom. Hemolysis was then spectrophotometrically determined, as described above.

**Salts and chemicals**

Stock solutions (1 M) were prepared dissolving heavy metals salts in distilled water. All salts and chemicals were pro analysis grade and purchased from Sigma (Milan, Italy).

**Statistics**

Data are shown as mean values ± standard error of the mean (s.e.m.). n values represent the number of independent experiments. With regard to discharge experiments, each data set is derived from 20 independent experimental sessions each including 2 oral arms (overall, 40 oral arms isolated from 10 animals collected in the same week were used for each condition tested). With regard to the hemolytic assay, each data set is derived from at least 20 independent experiments each performed on an individual blood sample (overall, 20 blood samples from 20 unrelated donors were used for each condition tested). Significance of the differences between mean values was tested using the one-way analysis of variance (ANOVA), followed by Bonferroni’s or Dunnet’s post hoc test. p<0.05 was considered as statistically significant.

**Results**

**Heavy metals significantly reduce in situ nematocysts discharge response in Pelagia noctiluca oral arms**

Simple mechanical stimulation of untreated oral arms with a non-vibrating test probe, without prior chemosensitization with glutamate, induced a very low nematocysts discharge (control, $6.87 \pm 0.75$ nematocysts/probe, Fig. 1). Combined chemical-mechanical stimulation of oral arms by incubation with the chemosensitizing compound glutamate ($10^{-3}$ M), followed by chemical stimulation with a non-vibrating test probe, elicited a discharge response significantly higher if compared to that elicited by simple mechanical stimulation (Glut, $39.72 \pm 0.82$ nematocysts/probe, p<0.001, Fig. 1). Treatment of oral arms for 20 min with 2 mM heavy metals salts followed by combined chemical-mechanical stimulation (incubation with $10^{-3}$ M glutamate, followed by mechanical stimulation) induced a significant reduction in discharge response compared to untreated oral arms (nematocysts/probe: $\text{ZnCl}_2: 31.43 \pm 0.70$, $\text{CdSO}_4: 15.37 \pm 2.12$, $\text{CoCl}_2: 21.80 \pm 0.94$, $\text{La(NO}_3)_3: 15.46 \pm 0.04$; p<0.001, Fig. 1). In all the conditions tested, discharge determined after heavy metals incubation was significantly higher than in control conditions, i.e. after simple mechanical stimulation (p<0.001, Fig. 1).

To test for the possible reversibility of the inhibitory effect of heavy metals on nematocysts discharge, $\text{ZnCl}_2$, $\text{CdSO}_4$, $\text{CoCl}_2$ and $\text{La(NO}_3)_3$-treated oral arms were rinsed with standard ASW for 20 min before glutamate exposure and mechanical stimulation by the non-vibrating test probe. In this case, discharge response was recovered, reaching values comparable to those observed in the same experimental conditions without prior exposure to heavy metals (nematocysts/probe: $\text{ZnCl}_2: 44.24 \pm 1.70$, $\text{CoCl}_2: 45.92 \pm 1.85$, $\text{CdSO}_4: 44.35 \pm 1.92$, p>0.05, Fig. 1).
2.25, n.s. vs glutamate, Fig. 2), with the notable exception of La(NO$_3$)$_3$ (nematocysts/probe: 16.46 ± 2.62, p<0.001 vs glutamate, Fig. 2).

In order to assess whether the inhibitory effect of heavy metals on discharge was due to oxidative stress events, oral arms were pre-incubated for 20 min in standard ASW containing heavy metal salts and 0.1 mM glutathione (GSH) or 1 mM ascorbic acid as antioxidants. Then, the solution containing heavy metals and the antioxidant was removed and combined chemical-mechanical stimulation was applied as described above. Simultaneous treatment with an antioxidant failed to reduce the inhibitory activity of heavy metals on discharge response (data not shown).

**Hemolytic assay**

Fig. 3 reports the hemolytic activity of crude venom extracted from holotrichous isorhizas nematocysts on 0.05% human erythrocytes suspensions. Treatment of erythrocytes for 1 h at 37 °C with crude venom induced a significant, dose-dependent hemolysis with respect to
Morabito et al.: Heavy Metals Affect the Homeostasis of the Jellyfish Pelagia noctiluca

Untreated erythrocytes (control). As shown, 5% and 10% v/v crude venom induced a 21.7 ± 2% and 45.63 ± 1.15% hemolysis respectively, while exposure to 20% v/v crude venom led to an almost complete hemolysis. Hemolysis obtained with 10% and 20% v/v venom concentrations was significantly higher with respect to that observed with 5% v/v crude venom (p<0.001). On this basis, to test for the possible effect of heavy metals on the hemolytic power of crude venom, 10% and 20% v/v venom concentrations were used and the 5% v/v venom concentration was excluded from the experimental design.

Incubation of erythrocyte suspensions with 2, 5 and 10 mM ZnCl₂, CdSO₄, CoCl₂ or La(NO₃)₃ did not elicit significant hemolysis compared to untreated erythrocytes (control, Table 1).

Interestingly, addition of 2, 5 and 10 mM ZnCl₂ or CdSO₄ to the erythrocyte suspensions significantly impaired the hemolytic activity of both 10 and 20% v/v crude venom (p<0.001, Table 1). In contrast, 2, 5 and 10 mM CoCl₂ treatment did not affect the hemolytic activity of both 10 and 20% v/v crude venom (Table 1). With regard to La(NO₃)₃ effect, 5 and 10 mM La(NO₃)₃ significantly impaired the hemolytic activity of both 10 and 20% v/v crude venom (p<0.001, Table 1), while 2 mM La(NO₃)₃ was ineffective (Table 1).

Incubation of erythrocyte suspensions with quaternary mixtures of ZnCl₂, CdSO₄, CoCl₂ and La(NO₃)₃ in equimolar concentrations (total concentration: 1 and 10 mM) did not elicit significant hemolysis compared to untreated erythrocytes (control, Table 1). The 10 mM heavy metals mixture completely blunted the hemolytic activity of 20% v/v crude venom, while the 1 mM heavy metals mixture was ineffective (Table 1).

**Discussion**

Metal pollution is a threat for ecological environment, affecting animals homeostasis at tissue and cell level [34]. Due to the detrimental effects of the heavy metals, research has focused on monitoring their concentrations and toxic effects in marine environments and organisms. In this regard, amongst marine vertebrates, fishes have been often used as bioindicators [35], while, among invertebrates, Cnidaria have not been exhaustively explored yet.

As described by Tarrant [36], pollutants may be internalized by Cnidaria by either uptake, ingestion or contact with chemicals contained in marine waters, suggesting a possible use of these specimens as a model for ecotoxicological investigations and biomonitoring of...
marine water quality. An interesting example is given by hydrazes, sensitive to metals and other environmental toxicants, including cadmium, zinc and butyltin [36-38].

The present investigation would propose the jellyfish *Pelagia noctiluca* as a novel model to verify and quantify heavy metals environmental pollution and bioaccumulation. This can be achieved by evaluating the effectiveness of nematocysts discharge and toxins delivered upon discharge, two features that, along with the cell volume regulation response under anisosmotic conditions, have allowed Cnidaria survival through the ages [1, 15, 30, 39]. What reported here completes our previous study on cell volume regulation of *Pelagia noctiluca* nematocytes [19], in an attempt to describe the homeostasis of this jellyfish in altered environmental conditions, i.e. in the presence of heavy metals. In this light, we tried to add novel information on the effect of metals on both nematocysts discharge and hemolytic power of crude venom. Hence, the same metals chosen for cell volume regulation investigations (i.e. Zinc, Cadmium, Cobalt and Lanthanum [19]) have been used. These pollutants are not only harmful for marine ecosystems, but are also of major concern for the human health [40, 41].

With regard to cell volume regulation capability, we previously demonstrated [19] that exposure of isolated nematocytes to heavy metals inhibits cell membrane transport systems underlying the regulatory volume decrease (RVD), the ubiquitous homeostatic response enabling cells [42, 43], including nematocytes [44], to counteract the osmolarity changes in the external medium. With regard to the hemolytic activity of crude venom extracted from *Pelagia noctiluca* nematocytes, Marino et al. [22] provided evidences of the inhibitory effect of divalent cations (1-10 mM), while nematocysts discharge and effectiveness of delivered toxins have not been considered yet to test heavy metal toxicity.

Here we show that a 20 minutes exposure of oral arms to heavy metal salts like ZnCl₂, CdSO₄, CoCl₂ or La(NO₃)₃ in concentrations similar to what reported elsewhere [22] and in line with concentrations measured in different basins [45, 46], dramatically impaired nematocysts discharge. This inhibitory effect seems to be in most cases - but not always - reversible, since extensive washing of oral arms with artificial sea water after exposure to heavy metals completely restored discharge response, with the exception of La(NO₃)₃. This observation is in line with what described by Morabito et al. [19], observing a reduced RVD response in isolated nematocytes of *Pelagia noctiluca* treated for 20 min with heavy metals. Inhibition of discharge response by metals has been demonstrated for the first time by Santoro and Salleo [47] and Salleo et al. [48] by using gadolinium, a metal of the lanthanide series of the periodic table and a specific Ca²⁺ channel blocker, thus defining discharge as a Ca²⁺-dependent phenomenon. Therefore, it is possible to hypothesize that heavy metals may similarly act by blocking ion channels involved in the transduction pathway triggering discharge. Moreover, Souza et al. [49] reported that Ca²⁺ channel blockers, such as nifedipine and verapamil, reduced cadmium uptake in several cell types, supporting the conclusion that cadmium may compete with Ca²⁺ for the uptake through Ca²⁺ channels. The presence of heavy metals in the extracellular environment may thus interfere with Ca²⁺ signaling and block discharge triggering.

Although discharge control has not been totally elucidated, some authors proposed that an osmotic influx of water, following ion uptake through channels, may represent the event underlying one of the first stages through which discharge normally occurs, i.e. an osmotic swelling of the nematocyte leading to delivery of tubule and toxins. On this basis, we suggest that heavy metals may affect discharge by impairing ion fluxes via membrane transport systems. As stated above, that ion transport in *Pelagia noctiluca* nematocytes is affected by heavy metals has been already shown [20], pointing out that heavy metals compromise ion fluxes through channels and/or co-transporters needed to perform RVD in isolated nematocytes. On a cellular level, the interaction of heavy metals with cell membrane transport mechanisms is the basis for their effect on biological systems. Examples of this interaction are given by Foulkes [50], describing the internalization of metals inside the cell, and Yu et al. [51], describing an alteration of the activity of epithelial sodium channels (ENaC) following divalent heavy metals exposure.
Our findings also show that the treatment of oral arms with heavy metals in the presence of GSH as antioxidant compound, did not protect the tissue from discharge inhibition. This result would indicate that heavy metals do not inhibit discharge response via an oxidative damage at tissue level. In contrast with this evidence, other authors described that metals may cause reactive oxygen species (ROS) production and, in turn, cell and DNA damage, impaired mitochondrial respiration, cell cycle arrest and apoptosis in various biological models [52-55]. Hence, the overall consideration is that the effect of metals such as Zinc, Cadmium, Cobalt and Lanthanum on biological substrates may be heterogeneous, also owing on the variety of the in vitro models used for toxicity assays [38, 56-58].

To further verify whether heavy metals totally compromise the defensive response of Cnidaria, that is based not only on nematocysts discharge, but also on effectiveness of delivered toxins, the hemolytic activity of crude venom from Pelagia noctiluca has been evaluated. Here we demonstrate that heavy metals, with the exception of Cobalt, substantially reduce the hemolytic power of crude venom extracted from nematocysts. The concentration and time of exposure of erythrocytes to metals allowed to exclude any direct hemolytic effect of Cadmium, Cobalt, Zinc and Lanthanum (see Table 1, control). The concentrations used (2-10 mM) were similar to those already employed by Marino et al. [22] for testing the hemolytic activity of Pelagia noctiluca crude venom in the presence of divalent cations or osmotic protectants.

In metal-exposed erythrocytes, oxidative damage of erythrocyte membrane leads to a reduction in membrane fluidity and increase in membrane fragility [59]. Accordingly, Hernández-García et al. [60] have recently described that heavy metals may induce hemolysis by lipoperoxidation. These observations differ to the results of our experiments, where no direct metal-dependent hemolysis was observed. We envision indeed that heavy metals may alter crude venom components, substantially preventing their hemolytic power. Accordingly, evidence exists that cations may affect the protein structure of toxins. In particular, previous studies have shown that, in venom of the jellyfish Carybdea marsupialis, Ca$^{2+}$ induced a significant decrease in hemolytic activity [12]. Moreover, our previous results [22] showed that metal ions like Ba$^{2+}$ and Cu$^{2+}$ block the hemolytic activity of Pelagia noctiluca crude venom.

Testing the effect of heavy metals mixtures instead of the single components may be a more realistic representation of sea water pollution [35, 45]. Therefore, a further set of experiments has been carried out to explore the effect of quaternary heavy metals mixtures on the hemolytic power of Pelagia noctiluca crude venom. A 10 mM (total concentration) Cadmium, Cobalt, Zinc and Lanthanum mixture completely impaired the hemolytic activity of crude venom, probably due to the presence of individual concentrations of Cadmium and Cobalt with maximal activity (Table 1). This observation leads to exclude possible antagonistic effects of the single components within the mixture. Conversely, a 1 mM (total concentration) mixture was completely ineffective, leading to exclude possible synergistic effects in our system, in agreement with previous findings on ternary and quaternary mixtures [35].

A mechanism of action suggested for many Cnidarian toxins [61], including Pelagia noctiluca [22], is the insertion of toxin molecules into the plasma membrane, the first phase of pore formation finally leading to colloid osmotic lysis of target cells. In this regard, the latter authors reported that the mechanism of toxin action mainly depends on pore-forming into the membrane of cell target rather than oxidative damage, since GSH, cysteine and ascorbic acid did not impair the hemolytic power. Based on these observations, an intriguing hypothesis is that the interaction of metals with cell membrane [50] or crude venom components may reduce the hemolytic activity of venom by impairing the binding and insertion of pore-forming toxins in the membrane of target cells. Accordingly, Miyake et al. [62] explained the inhibitory effect of divalent cations upon Vibrio metschnikovii cytolysin by changes in the erythrocytes membrane fluidity, so that pore formation due to the toxin was impaired, while, on the other hand, Iwase et al. [63] suggested that extracellular Ca$^{2+}$ may accelerate the rate of removal of the toxin by endocytosis.
Conclusion

Taken together, our findings show that nematocysts discharge response and effectiveness of toxins contained in the crude venom of the jellyfish Pelagia noctiluca are dramatically affected by short-term exposure to an external medium containing heavy metals, leading thus to reasonably hypothesize that the homeostasis of the entire animal may be compromised by these pollutants. Specifically, the strongest inhibition of nematocysts discharge was observed with Cadmium and Lanthanum and was irreversible with the latter, while Zinc and Cadmium were most effective in reducing the hemolytic activity of the crude venom.

Based on these evidences, we suggest that the jellyfish Pelagia noctiluca constitutes a good model for monitoring heavy metals environmental pollution and bioaccumulation. Evaluating Pelagia noctiluca physiological functions following pollutants exposure may provide novel contribution to eco-toxicological investigations. In this regard, further studies are warranted to elucidate the precise molecular mechanism of heavy metals in reducing nematocysts discharge and effectiveness of toxins and to monitor the effect of a long-term exposure to heavy metals on the homeostatic functions of this species.

Disclosure Statement

The authors have no conflict of interest to declare.

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