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Introduction 1.

Paraguat (PO) is a bipyridylium herbicide widely-used to prepare the land for planting and for controlling weeds in many crops around the world.¹ It has been demonstrated to be a highly toxic compound for humans and animals.² Moreover, due to its inherent toxicity and lack of effective treatment, the rate of death caused by PO is very high. It can cause multiple-organ failure, involving the lung, liver, and kidney.1 For this reason, PQ has been banned in the European Union.3 As a potent free radical generator, PQ toxicity has been found mediated by the production of free radicals, which resulted in oxidative damage to cells.^{4,5} Therefore, free radical scavengers and antioxidants have been expected to be candidates for preventing PQ toxicity.^{1,5} Some substances possessing free radical scavenging and antioxidant properties have been tested for protective capability in PQ-induced damage, such as vitamin C, vitamin E, melatonin, glutathione, docosahexaenoic acid, etc.2,5-8

Cyanobacteria are considered biotechnologically interesting organisms since they produce many valuable products, such as phycobiliprotein.9 Some cyanobacteria with nitrogen-fixing ability are able to survive under nitrogen-free environments, which makes these microalgae a good option to cultivate on

Selenium-phycocyanin from selenium-enriched cultures of Nostoc sp. isolated from rice field prevents human kidney cells from paraquatinduced damage

Hongzhi He,^a Saijun Liu,^b Huashou Li^a and Tianfeng Chen^b*^c

Selenium-containing phycocyanin (Se-PC) was purified from Se-enriched cultures of a nitrogen-fixing cyanobacterium (Nostoc sp.) using fast protein liquid chromatography. Ammonium sulfate precipitation, DEAE-Sepharose ion-exchange chromatography, and Sephacry S-300 size exclusion chromatography were employed in the process of protein purification. The purity ratio and the separation factor of the purified Se-PC were found to be 5.97 and 14.73, respectively. The Se content in the Se-PC was 550 μ g g⁻¹ protein as determined by ICP-AES analysis. Interestingly, we also found that, Se-PC could effectively suppress the paraguat (PQ)-induced cell death and morphological changes, reverse PQ-induced DNA fragmentation and nuclear condensation, and block PQ-induced cell apoptosis through inhibition of superoxide overproduction in human kidney HK-2 cells. The results indicate that Se-PC is a promising selenium species for the protection of renal cells from paraguat-induced damage.

> a large scale for biotechnological purposes, may reduce contamination risk and production costs of cultivation.¹⁰ Indeed, as edible blue-green algae, Nostoc species including N. flagelliforme and N. commune have been used as food for thousands of years in China.11 There have been some reports about purification of PC from nitrogen-fixing cyanobacteria.12-14 Moreover, Sun et al.¹⁵ recorded that phycocyanin (PC) treatment significantly attenuated PQ-induced acute lung injury in rats. At the same time, oral supplementation inorganic Se showed beneficial effects on the overall antioxidant potential of the animals because it can protect cells against the effects of free radicals.16 There are some reports that show antioxidant activity of selenium-containing phycobiliproteins purified from Se-enriched S. platensis.17-19 In previous determinations, selenium-containing PC (Se-PC) purified from Se-enriched S. platensis showed enhanced antioxidant abilities against oxidative damage in human erythrocytes through inhibition of reactive oxygen species (ROS) generation.20 However, no information about Se-PC for PQ-induced injury treatment is available. In the current study, Se-PC was purified from a seleniumenriched diazotrophic cyanobacterium Nostoc sp., and the protective effects and mechanism of Se-PC against PQ-induced renal injury was investigated.

2. Materials and methods

2.1 Materials

The nitrogen-fixing cyanobacterium Nostoc sp. (FACHB-85) was obtained from the Freshwater Algae Culture Collection of the

[&]quot;Key Laboratory of Agro-Environment in the Tropics, Ministry of Agriculture, South China Agricultural University, Guangzhou 510642, China

^bDepartment of Dermatology and Venerology, The First Affiliated Hospital of Jinan University, Guangzhou 510630, China

^cDepartment of Chemistry, Jinan University, Room 303, Guangzhou 510632, China. E-mail: tchentf@jnu.edu.cn; Fax: +86 20 85221263; Tel: +86 20 85225962

Paper

Institute of Hydrobiology, the Chinese Academy of Sciences and which was originally isolated from rice field in south China. DEAE-Sepharose was purchased from Sigma-Aldrich. The ultrapure water used in all experiments was supplied by a Milli-Q water purification system from Millipore. All the reagents used for cell culture experiments were bought from GIBCO.

2.2 Cultivation of high Se-enriched Nostoc sp. cells

Nostoc sp. was cultured with sterilized N-free mineral medium BG11 at 27 °C with a light illumination of 3000 lx and a light : dark cycle of 16 h : 8 h. For the cultivation of Se-enriched *Nostoc* sp., a stepwise Se addition procedure was carried out based on the results of toxicological tests, where 2 mg L^{-1} Se in the form of sodium selenite was added to the medium on day 7, 8, and 9, respectively. The accumulative concentration of Se was 6 mg L^{-1} . On day 14, algal cells were collected by centrifuging at 10 000*g* for 10 min at -4 °C and then lyophilized.

2.3 Separation and purification of PC and Se-PC by FPLC

The separation and purification procedure of Se–PC and PC was conducted by using the method of stepwise precipitation as we previously described.^{17,21} The sample was eluted with increasing NaCl concentration gradient. The pooled fractions were centrifuged, purified and desalted on a Sephacryl S-300 column.

2.4 Spectroscopic determination

A spectrophotometer method was employed to measure the UV-vis absorption spectra of testing samples. The purity ratio (A_{620}/A_{280}) and the separation factor (A_{620}/A_{655}) were calculated based on the UV-vis spectra.

2.5 Measurement of total Se

ICP-AES method was used to determine Se concentration. The sample was digested with 1 mL of H_2O_2 and 3 mL of concentrated nitric acid in a digestive stove (Qian Jian Measuring Instrument Co., Ltd., China) at 180 °C for 3 h. The digested product was reconstituted to 10 mL with Milli-Q water and used for total Se determination.

2.6 Cell culture and PQ treatment

Human kidney (HK-2) proximal tubular cell line was purchased from American Type Culture Collection (ATCC, Manassas, VA). The cells were incubated in Dulbecco's modified Eagle's medium (DMEM), supplemented with 10% fetal bovine serum (FBS), 100 U mL⁻¹ penicillin, and 50 U mL⁻¹ streptomycin in a humidified incubator with 5% CO₂ atmosphere at 37 °C. HK-2 cells were seeded at a density of 6×10^4 cells mL⁻¹ and pretreated with PQ for 3 h before the application of Se–PC or PC. Both living and dead cells were counted under microscopy with staining by trypan blue solution.

2.7 ABTS⁺ free radical scavenging activities

 $ABTS^+$ free radical scavenging activities of antioxidants were measured according to the method described by Zhang and coworkers.¹⁷ Briefly, 50 μ L of the tested samples was mixed with



Fig. 1 (A) DEAE-Sepharose chromatography of Nostoc sp. selenium-phycocyanin (Se-PC): the sample was eluted with increasing NaCl concentration gradient, first at 0.12 M for 40 min, followed by 0.12-0.21 M for 40 to 180 min and then maintained at 0.21 M until 220 min. The flow rate was 1.5 mL min $^{-1}.$ (B) Sephacryl S-300 chromatography of Nostoc sp. Se-PC: the sample was eluted with 2 mM Na-phosphate buffer (pH 7.0) at 1 mL min⁻¹. (C) UV-vis spectra of Se–PC obtained in different steps of purification: (I) crude extract; (II) fraction obtained from 50% ammonium sulfate precipitation; (III) pooled fractions obtained from DEAE-Sepharose chromatography; (|V|)pooled fractions obtained from Sephacryl S-300 chromatography.

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1 mL of ABTS reagent with absorbance of 0.70 \pm 0.02 at 734 nm, and then the absorbance at 734 nm was measured after the initiation of mixing for 6 min.

2.8 Determination of caspase-3 activity

The caspase activity of the samples was assessed by measuring the fluorescence intensity of caspase substrates with the excitation and emission wavelength at 380 nm and 440 nm, respectively.²²

2.9 TUNEL assay and DAPI staining

DNA fragmentation of HK-2 cells was detected by TUNEL assay followed the same procedure that described previously.²²

2.10 Determination of ROS generation

The effects of PQ, Se–PC and PC on the ROS generation were assessed by using a DCFH-DA assay in HK-2 cells as described previously.²³ Intracellular ROS level was quantified by measuring the fluorescence intensity of treated cells under a microplate reader (Bio-tek), with the excitation and emission wavelength at 488 nm and 525 nm, respectively.

2.11 Statistical analysis

All results are presented as mean \pm standard deviation (SD) (n = 3). Statistical analysis was carried out by using SPSS statistical package (SPSS 13.0 for Windows; SPSS, Inc., Chicago, IL). The difference between two groups was analyzed by two-tailed Student's t test. Differences with P < 0.05 (*) were considered statistically significant. The difference between three or more groups was conducted using the one-way analysis of ANOVA followed by the Student-Newman-Keuls *post hoc* test. Bars with different characters are statistically different at P < 0.05.

Results and discussion

3.1 Purification of Se–PC and PC by FPLC and measurement of Se concentration

The results for purification of Se–PC from Se-enriched *Nostoc* sp. are shown in Fig. 1 and Table 1. First, a stepwise precipitation procedure using $(NH_4)_2SO_4$ at 30% and 50% saturation degree was conducted.²⁰ Contaminating proteins were

Table 1	Purity ratio,	separation	factor and	Se conc	entration	of Se-Po	2
obtained	d in different	purification	1 steps				

Purification step	Purity ratio (A_{620}/A_{280})	Separation factor (A_{620}/A_{655})	Se concentration $(\mu g g^{-1} \text{ protein})$
Crude extract	0.93	1.69	243
Precipitation with	2.94	5.17	317
50% saturation of			
$(NH_4)_2SO_4$			
DEAE-Sepharose fast	4.26	10.86	_
flow chromatography			
Sephacryl S-300 chromatography	5.97	14.73	550

precipitated in $(NH_4)_2SO_4$ and Se–PC fraction was precipitated in $(NH_4)_2SO_4$ at 50% saturation. As shown in Table 1, the stepwise precipitation enhanced the purity ratio (A_{620}/A_{280}) and the separation factor (A_{620}/A_{655}) of Se–PC to 2.94 and 5.17, respectively. The peak of Se–PC appeared at about 176 min and a rational separation of Se–PC from other algal proteins was visualized (Fig. 1A). The purity ratio and separation factor of Se– PC reached 4.26 and 10.86, respectively. An effective removal of



Fig. 2 (A) Antioxidant activity of selenium–phycocyanin (Se–PC) by scavenging ABTS free radicals. (B) Antioxidant activity of PC by scavenging ABTS free radicals. (C) The IC_{50} indicating the half maximal inhibitory concentration at 30 min.



Fig. 3 (A) Effect of phycocyanin (PC) and selenium–phycocyanin (Se–PC) on cell viability of HK2 cells after 24 h exposure to paraquat (PQ). (B) Effect of PC and Se–PC on cell viability of HK2 cells after 48 h exposure to PQ. (C) Effect of different concentrations of Se–PC on cell viability in PQ-induced HK2 cells. (D) Effect of PC and Se–PC on cell morphology of PQ-induced HK2 cells. Different letters indicate statistically significant differences (P < 0.05).

other algal proteins before 120 min might be the key reason for achieving the high purity ratio of Se–PC. The purity was similar to previous works also employing DEAE ion exchange chromatography for PC purification from cyanobacteria.^{24–26} Sephacryl S-300 size exclusion chromatography was used for further purification of Se–PC. As a result, a sharp peak of Se–PC appeared in Fig. 1B. The purity ratio and separation factor of Se– PC after this step were increased to 5.97 and 14.73, respectively. As compared with methods addressed by other authors, the purification method in the present study gave a higher purity of Se–PC.^{24–26} Se content of protein obtained in different purification steps was shown in Table 1. After purification by ion exchange and size exclusion chromatography, Se–PC with the Se content at 550 μ g g⁻¹ was ultimately obtained. It is higher than that of Se–PC purified from *S. platensis* in our previous study.²⁰

3.2 Characterization of Se-PC by UV-vis

The characteristic absorption peak at 620 nm and strong absorbance below 300 nm in the UV-vis spectra of various Se–PC fractions (shown in Fig. 1C) corroborated the presence of the Se–PC protein. The reduced absorption intensity of peaks at 280 nm and 655 nm indicated the enhanced purity of Se–PC in terms of purity ratio (A_{620}/A_{280}) and separation factor (A_{620}/A_{655}) (Table 1).

3.3 Antioxidant activity of PC and Se-PC

ABTS' free radical scavenging assay was used to determine the antioxidant activity of PC and Se–PC.²¹ In the present study, the antioxidant activity of Se–PC is represented as the inhibition percentage. As shown in Fig. 2, the suppression effect of Se–PC on ABTS oxidation was stronger than that of PC, indicating the greater antioxidant activity of Se–PC in hydrophilic environment. For instance, Se–PC shows dose-dependent inhibition of ABTS oxidation, and the IC₅₀ (half maximal inhibitory concentration) of Se–PC (1.60 μ M) and PC (3.44 μ M) further confirmed this better antioxidant activity of Se–PC than that of PC.

3.4 Se–PC attenuates PQ-induced nephrotoxicity in HK-2 cells

Our previous studies have showed that selenium-containing allophycocyanin purified from selenium-enriched S. platensis showed enhanced hepatoprotective effects through reversing oxidative stress-induced cytotoxicity in HepG2 human hepatoma carcinoma cells.27 To confirm the possibility that Se incorporation into PC is able to enhance its properties in prevention of PQ-induced nephrotoxicity, MTT reduction assay was used to evaluate the possible protective effects of Se-PC on HK-2 cells. As shown in Fig. 3A and B, the treatment of HK-2 cells with PQ time-dependently decreased HK-2 cell viability. For instance, cells treated with 800 μ M PQ for 24 and 48 h decreased the cell viability by 25.7% and 46.8%, respectively. However, the pretreatment with Se-PC significantly relieved this cytotoxic effect, and Se-PC exhibited better protective effect than that of PC. Pretreatment with 4 µM Se-PC significantly increased the cell viability from 53.2% (PQ, 48 h) to 77.3% (Se-PC) while with 4 μ M PC the cell viability did not significantly change in contrasted with PQ treatment. For instance, Se-PC treatment (less than 8 µM) showed no significant cytotoxicity toward HK-2 cells (Fig. 3C). Only with 16 µM Se-PC, the viability of HK-2 cells was drastically decreased by 14.7%.

In order to confirm the protective effect of Se–PC, a phasecontrast microscopy was employed to observe the cell morphological changes. As shown in Fig. 3D, the exposure of HK-2 cells to 800 μ M PQ for 24 and 48 h caused a significant drop of cell density and elimination of cell-to-cell contact. However, these changes caused by PQ treatment were significantly reversed by the combined treatment with 4 μ M Se–PC. In contrast, PC treatment showed worse protective effect than that of Se–PC. Based on the above results, it suggested that Se incorporation into PC enhances its ability to suppress PQcaused cytotoxicity in HK-2 cells.

3.5 The effects of PC and Se–PC on PQ-induced apoptosis in HK-2 cells

Induction of cell apoptosis by exotic insults is an effective way to inhibit cell proliferation.^{21,28} Caspase is fundamentally important for regulating cell apoptosis.²⁹ Caspase-3 as the executor of apoptosis can inactivate PARP and initiate cell apoptosis.³⁰ Accumulated evidences have indicated that cisplatin-induced nephrotoxicity involves activation of caspase-3 in renal tubule epithelial.³¹ Therefore, our results suggest that PQ-induced cytotoxicity against HK-2 cells was achieved by induction of apoptosis, as convinced by the activation of caspase-3.

As shown in Fig. 4A, the treatment of HK-2 cells with 800 μ M PQ significantly activated the caspase-3 from 100% to 194.2%, indicating the existence of apoptosis in HK-2 cells. However, PQ-induced activation of caspase-3 was markedly suppressed by Se-PC co-treatment. For instance, treatment with 4 μ M Se-PC dramatically decreased the caspase-3 activity from 194.2% (PQ) to 131.1% (Se-PC). Moreover, 4 μ M Se-PC alone caused no change in caspase-3 activity. The results above clearly suggest that Se-PC inhibits PQ-induced apoptosis in HK-2 cells through the inactivation of caspase-3, and Se incorporation into PC enhances its ability in inhibition of PQ-induced HK-2 cell apoptosis.

Apoptotic cells may undergo nuclear condensation and DNA fragmentation.²² DNA fragmentation is a critical biochemical hallmark of apoptotic cells. TUNEL enzymatic labeling can be employed to especially detect the DNA fragmentation in early stage of apoptotic cells.³² Fig. 4B showed that 800 μ M PQ treatments caused evident DNA fragmentation of HK-2 cells, as



Fig. 4 (A) Selenium–phycocyanin (Se–PC) inhibits paraquat (PQ) induced activation of caspase-3. (B) Se–PC reverses PQ-induced DNA fragmentation and nuclear condensation in HK-2 cells.

Paper

convinced by the bright green fluorescence. However, PQinduced DNA fragmentation was completely reversed by Se-PC co-treatment, as convinced by the decreased number of TUNEL-positive cells. This protective effect of Se-PC on HK-2 cells was further confirmed by DAPI staining. For instance, cells exposed to 800 μ M PQ exhibited typical nuclear condensation. However, Se-PC co-treatment effectively blocked PQinduced morphological alterations. Moreover, Se-PC showed better protective effect than that of PC. These results further confirmed that Se incorporation in PC reduces DNA fragmentation and nuclear condensation, which are indicators of apoptosis. These effects contribute to the general trend of preventing PQ-induced apoptosis.

3.6 Se-PC prevents PQ-induced superoxide overproduction in HK-2 cells

Reactive oxygen species (ROS), including hydroxyl radical and superoxide anion, bring out severe harm to intracellular proteins, membrane lipids and DNA which ultimately results in cell apoptosis.^{22,23} Emerging studies have reported that ROS production is related to several types of human pathological conditions, including cisplatin-induced nephrotoxicity.^{33,34}





Fig. 5 (A) Selenium–phycocyanin (Se–PC) blocks paraquat (PQ) induced superoxide overproduction in HK-2 cells. (B) Se–PC reduces PQ-induced intracellular superoxide anion level examined using DHE fluorescent probe.

Therefore, intracellular superoxide anion level was examined by DHE fluorescent probe.

As shown in Fig. 5, PQ caused significant increase of superoxide anion level in HK-2 cells. For instance, treatment of cells with 800 μ M PQ notably promoted the superoxide anion level to 260.4%. However, co-treatment of the HK-2 cells with 4 μ M Se–PC significantly prevented the superoxide anion rate to 142.6%. The protective effect of PC at 4 μ M is similar to that of Se–PC at 2 μ M. However, the effect of Se–PC at 4 μ M was the strongest achieved. Moreover, no significant change in super-oxide anion level was detected after treatment with 4 μ M Se–PC alone. Fluorescent images detected by microscope further confirmed the protective effects of Se–PC, as convinced by weaken red fluorescence (Fig. 5B). The results indicate that Se–PC prevents PQ-induced superoxide overproduction in HK-2 cells, and Se incorporation into PC enhances its antioxidant activity.

In summary, a fast liquid chromatographic method for separation and purification of Se–PC from Se-enriched *Nostoc* sp. has been established. The purified Se–PC drastically prevented PQ-caused oxidative damage and nephrotoxicity in HK-2 proximal tubular cells. Mechanism investigation revealed that Se–PC blocked PQ-induced cell apoptosis through restraining the overproduction of superoxide. In conclusion, we suggest that Se–PC is a promising Se species for the advanced treatment of oxidative stress-mediated renal injury.

Conflicts of interest

There are no conflicts to declare.

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