Russell’s viper (Daboia Russellii) Venom Toxicity Neutralizing Efficacy of Curcumin- Gold Nanoparticle (C-GNP) in Experimental Animal Model

Keywords: Snake venom; Russell’s viper venom; Snake venom neutralization; Curcumin; Gold nanoparticle

Abstract

Herbs/herbal compounds possess anti-snake venom activity. Anti-snake venom activity of Curcuma longa in animal model has been reported earlier. In the present study, Curcumin (C), an active compound present in Curcuma longa was Conjugated with Gold Nanoparticle (C-GNP) and its neutralizing activity against viper (Daboia russellii) venom induced toxicity was evaluated in animal models. C-GNP was synthesized by adsorption method using chemically synthesized gold nanoparticles. Physicochemical characterization of C-GNP was done by DLS (size + zeta potential), UV-visible spectra, XRD and FESEM. Snake venom neutralizing activity of curcumin/C-GNP was done using in vivo and in vitro models. Animals were divided into gr. 1: Sham control, gr. 2: RVV control, gr. 3: Curcumin treated and gr. 4: C-GNP treated. Animal ethical clearance was availed before experiments. [IAEC/IV/Proposal/AG-001/2015; Dt: 10.04.2015]. One way ANOVA was done (n=4 and P<0.05 was considered as statistically significant). DLS size showed the hydrodynamic diameter of C-GNP to be 230-260 nm with polydispersity index of 0.103 and zeta potential was -18.32 mV. Maximum absorption of C-GNP was found to be 420 and 538 nm. XRD data confirmed the presence of crystalline gold in C-GNP, and FESEM indicated the presence of nearly spherical particle with size 18-24 nm. C-GNP could not antagonize RVV induced lethality, but it significantly antagonized RVV induced toxicity (edema, deturgescence, hemorrhage) and serum biochemical parameters (AST, ACP, LDH, urea, creatinine) in animal model. Significant neutralization of RVV phospholipase activity and plasma recalcification (in vitro) was observed after C-GNP treatment. The present study confirmed the formation of C-GNP and its efficacy against RVV induced toxicity. It may act as supportive treatment of snake bite victims. Further detailed studies on the antivenom activity and molecular mechanism of C-GNP are warranted.

Introduction

Russell’s viper envenomation is associated with local swelling, incoagulable blood, thrombocytopathy, spontaneous systemic bleeding, hypotension, increased capillary permeability, oliguria, proteinuria and urinary blood loss [1]. The only available treatment for RVV is anti snake RVV (ASVS), which was first developed in 1894. In viper envenomation, rapid initial absorption followed by slow absorption of venom. The delayed absorption has been linked with absorption of venom. The delayed absorption has been linked with the recurrence of envenomation when ASVS levels in blood decreases [2]. ASVS cannot neutralize viper venom induced local effects such as swelling, spontaneous systemic bleeding, increased capillary permeability, incoagulable blood, oliguria, thrombocytopathy, hypotension, proteinuria and urinary blood loss [3]. ASVS treatment carries a high risk of serum sickness and a lesser risk of anaphylaxis [4]. There is a need for alternative therapeutics that can effectively neutralize RVV induced toxicities. From the present laboratory, many herbal antioxidants have been identified against viper and cobra venom tested in animal models [5-13]. Recently, Gomes et al. reported that efficacy of herbal compound against RVV was increased by gold nanoparticle conjugation [14]. Use of nanotechnology increases the efficacy and bioavailability of drugs [15]. Karain et al. has shown that C60 fullerene nanoparticles decreased the lethality of Pacific Rattle snake venom in Acheta domestica model [16]. In the present study, curcumin (an active compound present in the rhizomes of Curcuma longa) was conjugated with gold nanoparticle and its Russell’s viper venom (RVV)-induced toxicity neutralizing efficacy was studied in animal model.

Materials and Methods

Venom: Lyophilized RVV was purchased from Calcutta Snake Park, Kolkata, India and preserved in desiccators at 4 °C in an amber colored glass vial until further use. It was dissolved in Milli-Q water, kept at 2-8 °C for overnight and centrifuged at 3000 rpm for 10 min. The supernatant was used as venom and kept at 2-8 °C until further use. The venom concentration was expressed in terms of dry weight (mg/ml, w/v).

Herbal compound: Curcumin was obtained commercially from Sigma-Aldrich (USA, Catalogue No: C7727). It (10 mg) was dissolved in 1 ml absolute alcohol and diluted 10 times by slowly adding distilled water. The final concentration of curcumin solution was 1 mg/ml and expressed in terms of dry weight (mg/ml, w/v).

Animals: Swiss male albino mice (20±1 gm) aged 8 weeks were obtained from authorized animal suppliers of Calcutta University. The animals were kept in polycarbonate cages, acclimatized and maintained in a controlled environment (temperature: 25±2 °C, humidity: 60±5% and 12 hr light/dark cycle). Food (pellet diet, Bengal gram and fresh green vegetables) and water was provided in ad libitum. All experimental protocols described in this study were approved by the animal ethics committee, Dept of Physiology, University of Calcutta and were in accordance with the guideline of the committee.

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for the purpose of control and supervision of experiments on animal (CPCSEA), Government of India, (Animal ethical clearance no: IAEC/IV/Proposal/AG-01/2015; Dated 10.04.2015). Swiss male albino mice (20±1 gm) were divided into: Gr.1- sham control, Gr.2- RVV control, Gr.3-curcumin treated and Gr.4- C-GNP treated.

Gold Nanoparticle Conjugation with Curcumin (C-GNP): Gold salt was purchased from Sigma-Aldrich (USA, Catalogue No: 484385). Gold Nanoparticles (GNP) were synthesized by citrate reduction method [17]. Briefly, an aqueous solution of HAuCl4 (250 µM, 25 mL) was brought to almost boiling condition up to 100 ºC stirring continuously on a magnetic stirrer. Freshly prepared trisodium isocitrate solution (600 µM final concentration) was added quickly to it which brought change in the solution color from pale yellow to bluish and finally to deep red. After obtaining a stable deep red color, the solution was allowed to be cooled in room temperature with continuous stirring. Gold nanoparticle prepared was then conjugated with curcumin to synthesize curcumin-conjugated GNP (C-GNP) by adsorption method. 1 mL of Curcumin (1 mg/mL) was added slowly to 2 mL of freshly prepared GNP on a vortex mixture, kept at 4 ºC for further use, and termed as C-GNP. To check the stability of the nanoparticles, it was kept at different temperatures for 3 months: at room temperature (28±3 ºC), at incubator (37±1 ºC) and at refrigerator (8±1 ºC).

Characterization of C-GNP: UV-visible spectra of curcumin and C-GNP were measured at the range between 200 nm to 700 nm using spectrophotometer (Shimadzu UV-1800) to determine maximum absorbance of respective solutions. The hydrodynamic diameter and zeta potential of C-GNP was measured by using photon correlation spectroscopy or Dynamic Light Scattering (DLS) machine (Beckmann Coulter Delta Nano C TM) to determine its size and stability. XRD measurements of C-GNP was done by XRD analyser (Phillips PW 1830) operating at a voltage of 40 KV and current of 20 mA with Cu-K radiation. FESEM of C-GNP was done using scanning electron microscope (Zeiss EVO 18).

RVV neutralization models

A. Neutralization of minimum lethal dose: The Minimum Lethal Dose (MLD) of RVV was assessed by injection of different amounts of RVV dissolved in 0.2 ml of 0.9% saline water into the tail vein of Swiss male albino mice (20±1 gm) after Theakston and Reid [18]. Various amounts of RVV (1-3 MLD) were mixed with fixed amount of curcumin/C-GNP incubated at 37 ºC for 15 min and injected into the tail vein of Swiss male albino mice (n=4). The MLD neutralization capacity of curcumin/C-GNP was calculated after 24 hrs of observation.

B. Neutralization of minimum edema dose: The Minimum Edema Dose (MED) of RVV was defined as the least amount of RVV when injected (intra-plantar) into Swiss male albino mice (20±1 gm), produced edema in the left paw (100% rise in paw volume). The right paw received saline 0.9% and was treated as control. To assess the anti-edema activity, various doses of RVV (1-5 MED in 0.05 ml) incubated at 37 ºC X 15 min with curcumin/C-GNP were injected (intra-plantar route) in Swiss male albino mice (n=4). The paw edema (volume) was recorded by the use of a digital caliper (Mitutua, Japan) at given time intervals (0,2,4 and 24 hrs), photographed by Canon digital camera (model no. SX 30 IS) and neutralization fold was calculated.

C. Neutralization of minimum hemorrhagic dose: The Minimum Hemorrhagic Dose (MHD) of RVV was defined as the least amount of RVV when injected intradermal (i.d.) into Swiss male albino mice (20±1 gm), produced cutaneous hemorrhage of 10 mm diameter after 24 hrs of observation [18]. To assess the anti-hemorrhagic activity, various doses of RVV (1-3 MHD in 0.05 ml) incubated at 37 ºC for 15 min with curcumin/C-GNP were injected (i.d.) in male Swiss albino mice (n=4). After 24 hrs, the animals were sacrificed, skin was removed and hemorrhagic spot was measured by the use of a digital calliper (Mitutua, Japan). Neutralization fold was calculated, photographs were taken by Canon digital camera (model no. SX 30 IS).

D. Neutralization of minimum defibrinating dose: The Minimum Defibrinating Dose (MDD) was defined as the minimum amount of RVV which, when injected intravenously (i.v.) into male albino mice (20±1 gm, n=4) produces incoagulable blood 2 hour later [18]. Inhibition of this activity was estimated by injecting different amount of RVV (i.v.) and curcumin (50 mg/kg/i.p.) or C-GNP (200 µl/20 gm mice/i.p.). The nature of the blood (clotted/non-clotted) was observed after 2 hour.

E. Neutralization of RVV phospholipase A2 activity: RVV phospholipase A2 (PLA2) activity was estimated by egg yolk coagulation method of Habermann and Neumann [19]. One unit of enzyme activity was defined as the amount of RVV which increased the coagulation time of egg yolk by 1 min. For neutralization of the RVV PLA2 activity, RVV (1-8 units) was incubated with curcumin/ C-GNP at 37 ºC for 15 min and PLA2 activity was assayed. Fold of RVV PLA2 neutralization was calculated in terms of PLA2 units.

F. Neutralization of RVV-induced minimum clotting dose of plasma: The Minimum Clotting Dose of Plasma (MCDP) induced by RVV was determined by Theakston and Reid [18]. Goat plasma was obtained from Government slaughter house in 3.8% sodium citrate in a ratio 1:9 (v/v). Plasma (0.2 ml) was incubated in a water bath at 37 ºC. To each tube, 0.1 ml of RVV in different concentrations (0.1 ml of saline added in control) was added. Finally 0.1 ml of 25 mM CaCl2 was added and clotting time was recorded with the help of a stop watch. Neutralization of RVV coagulant activity was done by mixing RVV (1-2 MCDP) with the fixed amount of curcumin/C-GNP at 37 ºC X 15 min and the clotting time was recorded.

G: Neutralization of RVV-induced serum markers: Animals were divided into Gr.1- sham control, Gr.2- RVV control, Gr.3- curcumin treated and Gr.4- C-GNP treated. Twelve µg (1/5th of MLD through s.c. route) of RVV was injected in Gr. 2, 3 and 4 animals. Gr. 3&4 animals were treated with curcumin (50 mg/kg/s.c.) & C-GNP (100 µl/20 gm/s.c.), respectively. After 18 hours, blood was collected from orbital plexus of animals, serum was prepared and creatinine, urea, Lactate Dehydrogenase (LDH), Serum Glutamic Oxaloacetic Transaminase (SGOT) and serum Acid Phosphatase (ACP) were estimated by biochemical kit method (Spinreac, Spain) according to manufacturer’s instruction.

Statistical analysis: Statistical significance was evaluated by one way Analysis of Variance (ANOVA). Values were expressed as mean ± standard error of mean (n=4) and P<0.05 was considered...
Results

Formation and characterization of C-GNP: C-GNP’s formed was orange color. It was visibly stable at room temperature (28±2 °C) for 15±2 days, incubator (37±1 °C) for 20±3 days and refrigerator (8±1 °C) for 60±5 days. Zeta potential of C-GNP was found to be -26.11 mV. λmax of curcumin and C-GNP was found to be 420 nm and 538 nm, respectively. When excited at 420 nm, curcumin showed fluorescent activity. C-GNP showed a quenching in fluorescence activity after excited at 420 nm. Hydrodynamic diameter of C-GNP was found to be 230-260 nm with polydispersity index of 0.103. X-ray diffraction analysis indicated that C-GNP was composed of crystalline gold. The Field Emission Scanning Electron Microscopy (FESEM) data of C-GNP showed that the particle size was 18-24 nm with nearly spherical shape (Figure 1).

RVV neutralization

A. Neutralization of minimum lethal dose: RVV Minimum Lethal Dose (MLD) was found to be 5 µg/20 gm/i.v and 60 µg/20 gm/ s.c. in male Swiss albino mice. Curcumin and C-GNP did not have any protection against RVV-induced lethality in male albino mice through intravenous and subcutaneous route (Table 1).

B. Neutralization of minimum edema dose: In vivo RVV Minimum Edema Dose (MED) was found to be 2 µg/20 gm/intraplantar. Curcumin gave 1 fold protection against RVV-induced edema, whereas, C-GNP caused 2 fold protection. The edema induced by RVV over a time period of 24 h was significantly neutralized by C-GNP as compared with curcumin. Curcumin gave 10.65% protection in RVV-induced paw edema after 24 h, whereas C-GNP gave 23.82% protection after 24 h (Table 1).

C. Neutralization of minimum hemorrhagic dose: In vivo RVV-induced Minimum Hemorrhagic Dose (MHD) was found to be 20 µg/20 gm/i.d/24 hrs in male albino mice. Curcumin (50 µg) failed to offer any protection against RVV-induced hemorrhage, whereas C-GNP gave 1 fold protection against RVV-induced hemorrhagic activity in Male Albino Mice (Table 1 and Figure 2).

D. Neutralization of minimum defibrinating dose: In vivo Minimum Defibrinating Dose (MDD) of RVV was found to be 2 µg. Treatment with curcumin offered no protection in RVV-induced defibrination activity, whereas treatment with C-GNP offered 1 fold protection in RVV-induced defibrinating activity (Table 1).

E. Neutralization of RVV-induced phospholipase A2 activity: In vitro RVV-induced 1 unit phospholipase A2 activity was found to be 1 µg. Curcumin offered no protection in RVV-induced PLA2 activity. C-GNP was found to neutralize 1 unit of PLA2 activity (Table 1).

F. Minimum clotting dose of plasma: In vitro RVV-induced Minimum Clotting Dose of Plasma (MCDP) was found to be 1 µg. Curcumin offered 1 fold protection in RVV-induced MCDP. There was 1.5 fold protection in MCDP after treatment with C-GNP (Table 1).

Figure 1: Characterization of C-GNP.

(A) Hydrodynamic diameter of C-GNP was found to be 230-260 nm with polydispersity index of 0.103; (B) XRD data showed the presence of crystalline gold in the sample; (C) FESEM diameter of C-GNP was found to be 18-24 nm with nearly spherical shape.

Figure 2: Effect of C-GNP in RVV-induced hemorrhage.

Treatment with C-GNP decreased RVV induced hemorrhage in animals. (A)- Sham control, (B)- RVV control, (C)- C-GNP treated.

Figure 3: Effect of C-GNP in RVV induced changes in serum markers in animal model. Significant change was observed in serum urea, creatinine, Lactate Dehydrogenase (LDH), Serum Glutamic Oxaloacetic Transaminase (SGOT) and serum Acid Phosphatase (ACP) after treatment with C-GNP in RVV-induced animals. *P<0.05; **P<0.01. Values were expressed as mean ± SEM; Gr.1- sham control, Gr.2- RVV control, Gr. 3- curcumin treated, Gr.4- C-GNP treated.
Table 1: Effect of C-GNP in RVV-induced in vivo and in vitro models.

<table>
<thead>
<tr>
<th>RRV's Viper Venom (RVV) neutralizing experiments</th>
<th>RVV (µg)</th>
<th>RVV + curcumin (µg)</th>
<th>RVV + C-GNP (µg)</th>
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<tr>
<td>Minimum Lethal Dose (MLD) (l.v.)</td>
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<td>20 (1 fold)</td>
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<tr>
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<td>1 (1 unit)</td>
<td>1 (1 fold) NP</td>
<td>1 (1 fold)</td>
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<tr>
<td>Minimum Clotting Dose of Plasma (MCDP)</td>
<td>1 (1 unit)</td>
<td>1 (1 fold) 1.5 (1.5 fold) NP</td>
<td>1 (1 fold) 1.5 (1.5 fold) NP 2 (1 fold)</td>
</tr>
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P: Protection; NP: No Protection.

Curcumin offered no protection in MLD, 1 fold protection in MED, no protection in MHD, no protection in MDD, no protection in PLA2 and 1 fold protection in MCDP, whereas C-GNP offered no protection in MLD, 2 fold protection in MED, 1 fold protection in MHD, 1 fold protection in PLA2 and 1.5 fold protection in MCDP.

G: Neutralization of RVV-induced serum markers: There was significant increase in serum urea (53.5%, P<0.01), serum creatinine (41.8%, P<0.05), serum lactate dehydrogenase (78.0%, P<0.01), serum glutamic oxaloacetic transaminase (198.5%, P<0.001) and serum acid phosphatase (84.2%, P<0.01) in group 2 RVV control animals as compared with group 1 sham control animals. Treatment with curcumin in group 3 animals caused significant decrease in serum urea (14.3%, P<0.05), serum creatinine (16.8%, P<0.05), serum lactate dehydrogenase (20.6%, P<0.05), serum glutamic oxaloacetic transaminase (37.6%, P<0.05) and serum acid phosphatase (23.4%, P<0.05) as compared with group 2 RVV control animals. Treatment with C-GNP in group 4 animals caused significant decrease in serum urea (28.9%, P<0.05), serum creatinine (25.3%, P<0.05), serum lactate dehydrogenase (32.4%, P<0.01), serum glutamic oxaloacetic transaminase (50.0%, P<0.01) and serum acid phosphatase (34.1%, P<0.01) as compared with group 2 RVV control animals (Figure 3).

Discussion

ASVS, developed by Dr. Albert Calmett in 1894 is the only treatment for snake bite till date, and there is no alternative available. However, there are many limitations (storage problem, availability in rural areas, dose difficulties, cost factors) and side effects (pyrogenicity, serum sickness, anaphylactic reactions etc.) of using ASVS [20-22]. Many herbs and herbal compounds have been identified to be active against snake venom [12]. Sometimes pure herbal compounds are less effective than the whole herbal extract. So, the question arises: Whether we can increase the efficacy and decrease the toxicity of the anti-snake venom herbal compounds using nanotechnology? Gold nanoparticles are a potential candidate in the field of nanomedicine and nanotherapeutics. In Ayurveda (Indian traditional medicinal system), processed gold and other metals were used as preventive and curative agents against many diseases. Kulkarni has established that ayurvedic processing of metals converted them into nanoparticles [23]. In traditional times, mixture of metallic formulation and herbs/herbal products were very common practice in therapeutics. Gomes et al. Ghosh and Saha have established the increase in efficacy of herbs/herbal constituents after conjugation of gold nanoparticles [14,24-26]. The present study was aimed to introduce a new approach of viper envenomation management (supportive therapy to ASVS) use of C-GGNP’s in animal models.

Several methods that are available for nanoparticle conjugation are adsorption or sodium borohydride method, which increases the efficacy of the herbs/herbal products. Curcumin was first made soluble in alcohol, and then was adsorbed in gold nanoparticle. The physicochemical characterizations of C-GNP were done to speculate the stability, size, shape and conjugation of the particle. Stability study of C-GNP was done by estimating zeta potential of the particles using dynamic light scattering technique. It is the potential difference existing between the particle surfaces and dispersing liquid. Gold nanoparticles bear negative charges; higher negative charge will cause more repulsion force between the particles along with more stability. The result indicated the formation of stable particles in room temperature (28±3 °C), incubator (37±1 °C) and refrigerator (6±1 °C). UV-vis spectrum is sensitive to particle size, shape, local refractive index and its interaction with medium [27]. Curcumin emits fluorescence when excited at its maximum absorbance. After conjugation with nanoparticles, they show fluorescence quenching activity. Joshi et al. showed that, after conjugation there was quenching of fluorescent activity of nanoparticles [28]. Dynamic Light Scattering (DLS) is used to determine the surface distribution profile of small particles in suspension, and it measures its hydrodynamic diameter, measuring the fluctuation of scattered light intensity of small sized particles [29]. Number of peaks of particle size determines its monodispersive or polydispersive nature. Polydispersity index is the parameter by which the particle-size differences can be estimated. In the present study, low polydispersity index confirms the formation of monodispersive C-GNP. Structural characterization of C-GNP was done by X-ray diffraction analysis and the result indicated the presence of crystalline gold in C-GNP. The Field Emission Scanning Electron Microscopy (FESEM) data of C-GNP showed the spherical shaped particle with size 18-24 nm. There was a difference between DLS diameter and FESEM diameter, because DLS diameter measures the water and other solute molecules attached with the particle, whereas the FESEM diameter measures the actual particle’s diameter.

Russell’s viper, mostly found in Indian subcontinent, South-East Asia and China causes a number of deaths throughout the year. The possible mechanism of action for lethality of RVV includes hemolysis, pre-synaptic neurotoxicity, rhabdomyolysis, vasodilation and shock [30]. About 70% of protein content in RVV is phospholipase A<sub>2</sub> with at least 7 isoenzyme forms [30]. High phospholipase activity is associated with high lethality [31]. Zinc-dependent metalloproteinases present in RVV are largely responsible for the hemorrhagic damage [32]. The hemorrhagic toxins directly damage microvessels and cause hemostasis, which provoke profuse bleeding [33,34]. RVV induced hemorrhage is also associated with pathophysiological manifestations including local necrosis, edema and blisters [32].

Earlier studies indicated that curcumin can inhibit phospholipases, metalloproteinases and protein kinase C [35-38] that may act effectively against snake bite. In the present study, the
efficacy of curcumin was increased by tagging with gold nanoparticle. C-GNP did not offer any protection against RVV-induced lethality in animal model, but it neutralized RVV induced edema, defibrination and hemorrhage. Study of venom induced in vitro phospholipase A2 activity and minimum clotting dose of plasma mimics the clinical manifestations of snake bite. C-GNP offers significant protection in RVV induced phospholipase A2 activity and clotting dose of plasma. RV envenomation causes nephrotoxicity, hepatotoxicity and myotoxicity in animal model [26]. Raised levels of urea, creatinine (for nephrotoxicity), AST, ACP (for hepatotoxicity) and LDH (for myotoxicity) were seen in RVV control animals. These markers were significantly reduced after treatment with C-GNP indicating that C-GNP inhibited RVV induced organ damages (liver, kidney, muscle) in animal models.

Previous studies indicated that curcumin has antioxidant and anti-inflammatory properties that help in exerting its protective role in pathophysiology [39]. Saha et al, Gomes et al. has shown that conjugation of gold nanoparticle with herbs/herbal compounds increased their potential against viper and cobra venom in animal model [14,25]. At this stage, it is difficult to point out the molecular mechanism of RVV neutralization by C-GNP, although the in vivo and in vitro studies showed some interesting data which confirmed the effective neutralization of RVV induced local toxic effects. It is likely that C-GNP acts at (1) neutralizing the RVV induced local damages at the vascular bed by inhibiting the pro-oxidant activity of RVV, (2) direct inhibition at the enzymatic level, (3) interference with the cellular markers (proinflammatory markers, antioxidants etc.) (Ghosh and Gomes, unpublished data). Although C-GNP could not neutralize RVV induced lethality, it may act as supportive treatment of snake bite victims. The present study confirmed the effective conjugation of gold nanoparticle with curcumin, which neutralized RVV-induced local toxicity in animal models, thus providing clue for supportive therapy along with ASVS treatment.

Conclusion

Snake venom neutralization using nanotechnology is a new domain of biomedical research which is showing promises for the development of supportive anti snake venom antidote from herbal resources.

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