Abstract. Crotoxin (CrTX), a neurotoxin, is isolated from the venom of South American rattlesnakes and has potent anti-tumor activity. Here, we investigated the antitumor effect of CrTX on the SK-MES-1 human lung squamous cell carcinoma cell line that has acquired resistance to epidermal growth factor receptor tyrosine kinase inhibitors. CrTX caused G1 arrest and p-JNK protein upregulation that resulted in apoptosis of SK-MES-1 cells. SP600125, a specific inhibitor of p-JNK, could rescue SK-MES-1 cells from CrTX-induced apoptosis. CrTX and gefinitib (Iressa) both inhibited the viability and proliferation of SK-MES-1 cells from CrTX-induced apoptosis. CrTX and gefitinib (Iressa) both inhibited the viability and proliferation of SK-MES-1 cells in a dose- and time-dependent manner. The combination of CrTX and Iressa significantly enhanced the antitumor activity of Iressa. In vivo studies revealed that CrTX caused increased damage to blood vessels and reduced tumor size when combined with Iressa. The present study showed that the JNK signal transduction pathway mediated the anti-apoptotic effect of CrTX, and furthermore, CrTX could enhance the antitumor effect of tyrosine kinase inhibitors in cells with acquired resistance.

Introduction

According to the US Centers for Disease Control and Prevention, the number of deaths from lung cancer is higher than any other type of cancer, which holds true for both men and women (1). Its overall 5-year survival rate is only 8-14% following diagnosis and 40% after complete surgical resection. NSCLC represents ~80% of the total lung cancer cases, and the subtypes include adenocarcinoma, squamous carcinoma, and large cell carcinoma (2). The squamous carcinoma subtype accounts for ~50% of lung cancers. Although tyrosine kinase inhibitors have proven useful in combination with chemotherapy against lung cancer, they have demonstrated good efficacy only in 10-15% of lung cancers that possess a sensitizing mutation in the epidermal growth factor receptor (3). Crotoxin (CrTX), the dominant toxin of the South American rattlesnake (Crotalus durissus terrificus) venom (4), was the first snake venom protein to be purified and crystallized. It is a non-covalent complex formed by two non-identical subunits, one acidic (subunit A/crotapotin) and one basic (subunit B/crotactine). Subunit B is a phospholipase A2 (PLA2) formed by a single chain of 122 amino acid residues cross-linked by seven disulfide bonds (5). The classic biological activities normally attributed to CrTX include neurotoxicity, myotoxicity, and nephrotoxicity. Many studies recently have shown that CrTX also has immunomodulatory, anti-inflammatory, antimicrobial and analgesic actions. Studies have also reported that CrTX has antitumor effects (6). Interestingly, CrTX appears to have cytotoxic effect that correlates directly with tumors that express higher levels of the epidermal growth factor receptor (EGFR) though CrTX stimulated tyrosine phosphorylation of the EGFR; the significance of this observation has yet to be established (9).

Our previous studies revealed that the cytotoxic mechanism of action of CrTX may be more complex than previously thought. We have demonstrated that apoptosis and autophagy are activated during CrTX-induced death of cancer cells. These studies included K562 leukemia cells that have resistance to imatinib due to a constitutive tyrosine kinase activity (7). This study was undertaken to evaluate the anti-lung cancer efficacy of CrTX in combination with Iressa and to further establish the molecular mechanisms by which CrTX exerts its antitumor functions by looking at the expression of the pro-apoptotic proteins, caspase-3 and p-JNK, and the apoptosis inhibitor protein, Bcl-2.

Materials and methods

Drug preparation. Crotoxin was supplied by Celtic Biotech Ltd. (Dublin, Ireland). It was purified from Crotalus durissus terrificus.
terricus venom by a combination of size exclusion and anion exchange (4). The identity of the protein was confirmed through molecular weight determination by mass spectrometry (showing averaged signals at 9500 and 14500, in addition to the presence of isofoms). Purity was determined by PAGE and size exclusion >99%. Lethality in mice was determined by the i.p. injection of 0.1 mg with death being recorded within 3 h.

Cell culture and drug treatment. SK-MES-1 cells, a human lung squamous carcinoma cell line with acquired resistance to epidermal growth factor receptors and tyrosine kinase inhibitors, were purchased from the Shanghai Institute of Cell Biology. Cells were cultured in RPMI-1640 medium (Gibco, Rockville, MD, USA) containing 10% heat-inactivated fetal bovine serum (Gibco), 0.03% L-glutamine (Sigma, St. Louis, MO, USA) and 200 U/ml insulin, and incubated in a 5% CO2 atmosphere at 37°C. Cells in mid-log phase were used in all the experiments. To determine the dose- and time-response of SK-MES-1 cells to CrTX, SK-MES-1 cells were plated into 96-well microplates (103 cells/well) and cultured for 24 h. CrTX (12.5, 25, 50 or 100 µg/ml), Iressa (1.25, 2.5, 5 or 10 µmol/l), Iressa 2.5 µmol/l with CrTX (12.5, 25, 50 and 100 µg/ml) or CrTX 25 µg/ml with Iressa (1.25, 2.5, 5 or 10 µmol/l) were added to the culture medium, respectively.

Antibodies against phosphorylated c-Jun N-terminal kinase (p-JNK), Bcl-2 and cleaved caspase-3 were purchased from Cell Signaling Technology (Woburn, MA, USA). SP600125, a specific inhibitor of p-JNK, was purchased from Sigma.

Cell viability assay. Cell viability was assessed with a 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide reduction assay (MTT) (Sigma) 24, 48 and 72 h after drug treatment as previously described (10). The MTT solution was added to the culture medium (500 µg/ml) 4 h before the end of treatment and the reaction was terminated by the addition of DMSO. After incubation at room temperature for 30 min and gentle agitation for 10 min, the absorbance value (A) at 570 nm was read using the Benchmark Microplate Reader (Bio-Rad Laboratories, Hercules, CA, USA). Three independent experiments were performed to generate an average value as the final result for analysis. The percentage of cell death was calculated as following: cell death (%) = (1 - Aexperiment well/Apositive control well) x 100.

Cell colony formation assay. The capability of cell colony formation corresponded to the degree of malignancy of the tumor. The more cell colonies formed, the more malignant the tumor. The tumor was (11). Cells in mid-log phase were digested into a single cell suspension and inoculated into 24-well plates. Each well was inoculated with 200 cells. After 24 h, when the cells adhered to the plate, the medium was removed and various concentrations of CrTX in the medium were applied to treat the cells for 72 h. Subsequently, the medium was changed to fresh RPMI-1640 medium and the cells were incubated for 12 days. Then, cells were fixed with methanol and stained with Giemsa. The colonies including >50 cells were calculated. The experiments were repeated three times independently.

Cell cycle analysis. Flow cytometry (FCM) is usually used to determine cell cycle progress and apoptosis (12,13). Five experimental groups were included in the analysis: control; CrTX (25 µg/ml); SP600125 (10 µmol/l) with CrTX (25 µg/ml); Iressa (5 µmol/l); and CrTX (25 µg/ml) with Iressa (5 µmol/l). SP600125 was added to the culture medium 1 h before CrTX. After being treated for 24 h, cells were harvested using 0.25% trypsin, and 106 cells/sample were aliquoted for analysis. To avoid membrane damage as much as possible, cells were observed with the inverted microscopy after about 2 min of trypsin digestion, and then removed from the flask when the shapes of most cells changed from long spindle-shaped to round. The cells were centrifuged at 5,000 x g at 4°C for 5 min and washed in cooled PBS, fixed with pre-cooled 70% ethanol for 24 h, and treated with 200 µl RNase A (1 g/l) for 30 min. Propidium iodide (PI) was added to a final concentration of 20 µg/ml for 30 min in the dark. Finally, the cell cycle was analyzed with FCM (14).

Detection of apoptosis by Annexin-V staining. Cell treatments were the same as previously described. The cells were harvested after treatment, washed twice with pre-chilled PBS (4°C), and resuspended in 500 µl binding buffer to a final cell concentration of 1X106 cells/ml. They were incubated in the dark with 5 µl Annexin-V and 5 µl PI solution of 20 µg/ml for 30 min. Subsequently, the suspension was analyzed by FCM.

Protein preparation and immunoblotting. Cells were harvested and rinsed with ice-cooled PBS twice. Five volumes of western blotting buffer for each volume of cell pellets was added and the mixture was sonicated on ice (1 sec/ml per sonicate, resting 30 sec between intervals, for a total of 5 times). Microcentrifugation was performed at 10,600 x g at 4°C for 10 min and the supernatant was preserved at -70°C for later use. The protein concentration was determined using the BCA kit (#23225, Pierce, Rockford, IL, USA). Proteins were separated with 10% SDS-PAGE gel, transferred to nitrocellulose membrane and immunoblotted with antibodies against Bcl-2 (1:500, sc-7382, Santa Cruz Biotechnology, Santa Cruz, CA, USA), p-JNK (1:1000, #4668, Cell Signaling Technology), or caspase-3 (1:500, #9662, Cell Signaling Technology) at 4°C overnight. The reaction of primary antibodies was detected with horseradish peroxidase-conjugated secondary antibodies used at a 1:5000 dilution in blocking solution for 1 h at room temperature. Immunoreactivity was detected by enhanced chemiluminescence (SuperSignal West Pico Chemiluminescent Substrate (#34077, Pierce) and visualized by autoradiography. Protein β-actin (1:5000, #A5441, Sigma) was used as a loading control.

Xenotransplantation and drug treatment. All animals used in this study were handled strictly according to the guidelines for use and care of experimental animals approved by the Ethics Committee of Soochow University. Twenty-six female Balb/c nude mice with an average body weight of 18-20 g (aged 4-6 weeks) were housed in a specific pathogen free animal facility. All manipulations (i.e. handling, invasive procedures and tumor volume measurements) were performed in a laminar flow hood under strict sterile conditions. Mice were injected with 106 cells/ml suspended in 0.2 ml PBS into the left axillary space tissue. Mice were monitored daily until detection of the first tumor nodule. Tumor nodules were first noted in eight of the 20 mice (40%) at 3 days after xenotransplantation.
Treatment with CrTX was initiated 11 days after xenotransplantation when the tumor reached 5-6 mm in diameter. Except for 2 no-tumor mice, 24 mice were randomly assigned to one of four experimental groups (n=6 per group): control; CrTX (10 µg/kg, i.p., twice weekly); Iressa (100 mg/kg/day, i.g., daily); and CrTX with Iressa (CrTX 10 µg/kg, i.p., twice weekly with Iressa 100 mg/kg/day, i.g., daily). At the end of 30 days treatment, the mice were sacrificed, and the tumor xenografts were removed, sectioned and analyzed with transmission electron microscopy (TEM).

Statistical analysis. For statistical analysis, all data were presented as mean ± SEM. Statistical analysis was carried out by ANOVA followed by the Dunnett’s t-test, considering P<0.05 to denote significant differences.

Results

CrTX potentiates antitumor effects of Iressa. Both CrTX and Iressa inhibited SK-MES-1 cell viability in a time- and dose-dependent manner. The MTT assay revealed that after 24 h of treatment, the inhibitory rates of CrTX (4 µmol/l) and Iressa (10 µmol/l) on cell viability were 28.3 and 24.3%, respectively. When the incubation time was prolonged to 48 h, the inhibitory rates increased to 48.1 and 37.8%, and reached 53.6 and 55.9%, respectively, after 72 h of treatment. At the lower dose of CrTX (2 µmol/l) and Iressa (5 µmol/l), the inhibitory ratios were only 32.8 and 41.3% with 72 h of treatment, respectively (Fig. 1A and B). With a combination of various doses of Iressa and a constant dose of CrTX, or of various doses of CrTX with a constant dose of Iressa, the inhibitory rates on SK-MES-1 cells increased in comparison with the same doses used alone (Fig. 1C and D). The Q score was 0.85-1.15 suggesting there was an additive effect when CrTX was combined with Iressa. Based on the effectiveness on inhibiting viability of SK-MES-1 cells, the concentrations used in subsequent experiments were 1 µmol/l CrTX with 5 µmol/l Iressa.

The cell colony formation assay also showed that CrTX and Iressa inhibited the colony formation of SK-MES-1 cells in a dose-dependent manner (Fig. 2A and B), and the inhibitory effects were significantly enhanced when they were combined (Fig. 2C and D).

Effects of CrTX on the cell cycle and apoptosis. To understand the mechanisms of CrTX on cell growth inhibition, FCM was used to analyze apoptosis and cell cycle arrest. After treatment for 24 h, the apoptosis rates of SK-MES-1 cells in the control group, the SP600125 with CrTX group, the Iressa group, the CrTX group, and the CrTX with Iressa group were 0.89±0.06, 10.3±0.04, 1.07±1.1, 11.8±1.7 and 19.6±2.8%, respectively. The increase in apoptosis in the SP600125 with CrTX group was not statistically significant from the control group (P>0.05), whereas the apoptosis rate in the CrTX, Iressa and combination-treatment groups increased significantly in comparison to controls (P<0.05, n=3) (Fig. 3A and C). SP600125 protected SK-MES-1 cell from apoptosis indicating that the p-JNK signal transduction pathway may play an important role in the activity of CrTX.

Additionally, from the results of the cell cycle analysis, we found that CrTX significantly increased the number of
Figure 2. Effects of CrTX and Iressa on colony formation of SK-MES-1 cells. Cell colony formation was analyzed 72 h after drug treatment. Fresh RPMI-1640 medium replaced the medium with drugs and the cells were incubated for 12 days. After that, cells were fixed with methanol and stained with Giemsa. The colonies including >50 cells were calculated. (A and B) Colony formation under the effect of CrTX and Iressa individually. (C and D) Colony formation rate with the combination of CrTX and Iressa; n=3. Mean ± SD.

Figure 3. Effects of CrTX on apoptosis and cell cycle of SK-MES-1 cells. (A) Cells were harvested after treatment for 24 h, washed twice with pre-chilled PBS, and resuspended in 500 µl binding buffer to a final cell concentration of 1-5x10^5 cells/ml. They were incubated in the dark with 5 µl Annexin-V and 5 µl PI solution of 20 µg/ml for 30 min. Subsequently, the suspension was analyzed by FCM. (B) After being treated for 24 h, cells were harvested using 0.25% trypsin, and 10^6 cells per sample were aliquoted for analysis. Cells were centrifuged at 5,000 x g at 4˚C for 5 min and washed in cooled PBS, fixed with pre-cooled 70% ethanol for 24 h, and treated with 200 µl RNase A (1 g/l) for 30 min. Propidium iodide (PI) was added to a final concentration of 20 µg/ml for 30 min in the dark. Finally, the cell cycle was analyzed with FCM. (C and D) Statistical analysis of A and B, respectively. *P<0.05, #P<0.01 vs. controls; n=3. Mean ± SD.
SK-MES-1 cells in the G1 phase compared to the control group (Fig. 3B and D).

CrTX increases active caspase-3 and p-JNK but downregulates Bcl-2. SK-MES-1 cells were treated with 1 µmol/l CrTX for 24 h and harvested for western blot analysis of pro-apoptotic proteins. CrTX significantly upregulated cleaved caspase-3 and p-JNK protein but downregulated Bcl-2 protein in SK-MES-1 cells. Pretreatment of SK-MES-1 with SP600125 could effectively inhibit the activation of pro-apoptotic proteins induced by CrTX (Fig. 4).

Antitumor effects of CrTX and/or Iressa in vivo. We evaluated the effects of CrTX, Iressa and CrTX combined with Iressa treatment on the growth of SK-MES-1 cells injected subcutaneously into athymic nude mice. The results showed that treatment with either CrTX or Iressa caused a decrease in tumor size compared to control mice, while treatment of CrTX combined with Iressa induced a significantly greater reduction in tumor weight. The tumor inhibitory rate increased to 35.92% with the drug combination from a <23% maximum with single drug therapy (Fig. 5).

The capillaries in the tumor were analyzed by TEM. In the control group, the capillaries in the interstitial substance of the tumor were plentiful, the shape of the endothelial cells was normal, and the growth of the basal membrane was unabridged. In the experimental group, there were only a few capillaries. The endothelial cells were engorged and some had shrunk. Apoptotic cells were observed in the tumor. In the combined treatment group, the appearance of the endothelial cells was much more pronounced. The vessel wall basal...
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which has endogenous tyrosine kinase activity and is resis
tive toward cell lines expressing an upregulated density of
CrTX has certain specificity toward solid tumors.

Figure 6. Effects of CrTX on capillaries in tumor tissues. Balb/C nude mice
were injected with $10^5$ cells/ml suspended in 0.2 ml PBS into the left axillary
space tissue. Treatments with drugs were initiated 11 days after xenotrans-
plantation and the tumor reached about 5-6 mm in diameter. At the end of
30 days treatment, the mice were sacrificed, and the tumor xenografts were
excised, sectioned and analyzed with transmission electron microscopy.

membrane was ruptured, thinned, and the lumen was narrow;
additionally there was electron-dense material in the vascular
endothelial cells (Fig. 6).

Discussion

CrTX is a potent antitumor agent, which has cytotoxic activity
against a variety of human tumor cell lines in vitro (6,7,15).
CrTX also exerts an anti-inflammatory/analgesic activity
seemingly through the lipoxygenase pathway without nega-
tively impacting its anticancer activity. For both functions,
CrTX requires the PLA2 activity of subunit B and the ability
of the complex to dissociate into its subunits (9,16). Antitumor
inhibition in vivo of 83% has been demonstrated against Lewis
lung carcinoma (17) and MX-1 human mammary carcinoma
(69% growth inhibition). A lower activity (44% growth inhibi-
tion) was observed with HL-60 leukemia cells, suggesting that
CrTX has certain specificity toward solid tumors.

CrTX-induced cytotoxic effects appear to be highly selec-
tive toward cell lines expressing an upregulated density of
EGFRs (9), though CrTX appeared to unexpectedly promote
EGFR phosphorylation. Enhanced EGFR activity in cancer
cells and tumors is associated with increased growth, survival
and angiogenesis of tumors (18). The human CML line K562
which has endogenous tyrosine kinase activity and is resis-
tant to imatinib was also quite sensitive to CrTX (7). In this
study, the antitumor effects of CrTX against SK-MES-1 cells,
possessing resistance to EGFR tyrosine kinase inhibitors, was
dose- and time-dependent. When CrTX was combined with
Iressa, the antitumor effect of Iressa against this tyrosine
kinase resistant cell line was strengthened, which suggests an
additive effect of the two drugs. In vivo, CrTX and Iressa both
had antitumor effects in mice with tumor xenografts; but the
inhibitory rate with the combined treatment was increased by
50%. These data suggest that CrTX stimulation of EGFR phos-
phorylation is a cellular response to the induction of apoptosis
and that the co-utilization of a tyrosine kinase inhibitor blocks
this rescue pathway. It should be noted that the animal treat-
ment protocol for CrTX therapy was not optimized through
the induction of tolerance (19), but Iressa served to sensitize
the tumors permitting low and infrequent dosing of CrTX to
be effective.

There is a close and complex relationship between
apoptosis of tumor cells and cell cycle arrest. Although they
complement each other in inhibiting tumor progression, the
effects of CrTX on inducing apoptosis of tumor cells and
blocking the G1 phase are achieved through different intra-
cellular pathways. FCM showed that CrTX itself significantly
increased the rate of apoptosis of SK-MES-1 cells particu-
larly in the G1 phase indicating that the antitumor effects of
CrTX were closely related to induction of apoptosis and to
cell cycle arrest.

Many researchers have paid more attention to the signaling
pathway mediated by p-JNK, one of the main signal trans-
duction systems regulating apoptosis (20,21). Cells transmit
extracellular signals to the nucleus through this pathway (22).
When the pathway is activated by ultraviolet radiation, nerve
growth factor, satratoxin, or antitumor drugs, JNK is phos-
phorylated. Phosphorylated JNK plays an important role in
apoptosis and cell cycle arrest (23-25). As a central apoptotic
effector, caspase-3 plays a key role in promoting apoptosis
(26). When p17, the active form of caspase-3, is detected, it
indicates that caspase-3 proenzyme is activated and ready to
execute its apoptotic function.

In the present study, the expression of p-JNK and p17
increased during the process of CrTX-induced apoptosis of
SK-MES-1 cells. The use of SP600125 confirmed the role
of the p-JNK signaling pathway. The activation of p-JNK
probably mediated the activation of caspase-3. JNK protein is
normally located in the cytoplasm, and when it is activated,
it aggregates in the nucleus as observed herein. Consistent
with previous data, the activation of caspase-3 is an important
component of the CrTX mechanism (7,15). The expression of
active Bcl-2 was downregulated when the SK-MES-1 cells
were treated with CrTX. All of the above are indicative of the
induction of apoptosis.

Most antitumor drugs exhibit severe host toxicity due to
their poor selectivity toward cancer cells. Iressa, an anti-EGFR
tyrosine kinase inhibitor, is widely used clinically in Asia, and
its therapeutic effect has been confirmed in a specific cohort of
lung cancer subjects. However, it is expensive and its efficacy
is limited and is compounded by primary or secondary drug
resistance which develops with extended treatment periods.
When combined with Iressa, CrTX can significantly increase
its tumor suppressive activity or reduce the required dose of
Iressa suggesting that CrTX appears to be a useful conjunctive
therapy for lung cancer treatment. The present study aids in
the understanding of the mechanism of CrTX-induced apoptosis
in SK-MES-1 cells, and provides new insight into antitumor
combinations for lung cancer therapy.
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References