Direct effect of croton oil on intestinal epithelial cells and colonic smooth muscle cells

Xin Wang, Mei Lan, Han-Ping Wu, Yong-Quan Shi, Ju Lu, Jie Ding, Kai-Cun Wu, Jian-Ping Jin, Dai Ming Fan

INTRODUCTION

Croton is the fruit of Croton tiglium L, which consists of 34%-57% croton oil and 18% proteins. Croton oil (CO) contains crotonoleic acid, crotonic acid, crotonyl alcohol, 16 kinds of crotonyl alcohol bisester and many kinds of crotonyl alcohol trisester. Also contained in croton oil is 18% proteins. Croton oil (CO) contains crotonoleic acid. Croton oil has been used as a potent stimulant in the making of inflammatory models, especially in models of skin and mucosal inflammation. Croton oil also has a direct effect on the contractility of Guinea pig colonic smooth muscle cells.

CONCLUSION: At a high CO concentration (>20 mg·L⁻¹), cell growth and proliferation are inhibited in a dosage-dependent manner. Increase in cell proliferation and in malignant conversion of the cellular phenotype is observed in cells cultured chronically with CO. COX-2 mRNA expression decreases significantly, while human mitochondrial cytochrome C oxidase subunit II mRNA expression increases markedly in HIEC treated with CO. CO also has a direct effect on the contractility of Guinea pig colonic smooth muscle cells.

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METHODS: Growth curves of HIEC were drawn by MTT colorimetry. The dynamics of cell proliferation was analyzed with flow cytometry, and morphological changes were observed under light and electron microscopy after long-term (6 weeks) treatment with CO. Expression of cyclooxygenase-2 (COX-2) mRNA was detected by dot blot in HIEC treated with CO. Genes related to CO were screened by DD-PCR, and the direct effect of CO on the contractility of isolated guinea pig colonic smooth muscle cells was observed.

RESULTS: High concentration (20 - 40 mg · L⁻¹) CO inhibited cell growth significantly (1, 3, 5, 7d OD sequence: (20 mg·L⁻¹) 0.040±0.003, 0.081±0.012, 0.147±0.022, 0.232±0.016; (40 mg·L⁻¹) 0.032±0.044, 0.056±0.012, 0.104±0.010, 0.189±0.006; OD control 0.031±0.008, 0.096±0.012, 0.173±0.009, 0.300±0.016, P<0.01), which appeared to be related directly to the dosage. Compared with the control, the fraction number of cells in G1 phase decreased from 0.60 to 0.58, while that in S phase increased from 0.30 to 0.34 and DNA index also increased after 6 weeks of treatment with CO (the dosage was increased gradually from 4 to 40 mg · L⁻¹). Light microscopic observation revealed that cells had karyomegaly, less plasma and karyoplasm lopsidedness. Electron microscopy also showed an increase in cell proliferation and in the quantity of abnormal nuclei with pathologic mitosis. Expression of COX-2 mRNA decreased significantly in HIEC treated with CO. Thirteen differential CDNA fragments were cloned from HIEC treated with CO, one of which was 100 percent homologous with human mitochondrial cytochrome C oxidase subunit II. The length of isolated guinea pig colonic smooth muscle cells was significantly shortened after treatment with CO (P<0.05).

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AIM: To investigate the direct effect of croton oil (CO) on human intestinal epithelial cell (HIEC) and guinea pig colonic smooth muscle cells in vitro.

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According to the recent, that dehydrocrotonin (DCTN), an extract of CO, had anti-ulcer effect, especially in ulcers induced by hypothermia, alcohol and pyloric ligation. This effect was supposed to the related to the increase in the release of prostaglandin E2, the non-specific competitive binding of H2 and M receptors of stomach, the reinforcement of mucosal protective action and the weakening of mucosa-imparing factors. So DCTN has the potential to be a new anti-ulcer drug.

It was reported that croton extract could not only suppress tumor growth but also promote tumor development. It has been confirmed that DCTN, which possesses neither genetic toxicity nor cytotoxicity, has anti-tumor activity and can significantly elongate the survival period of mice with S180 sarcoma and ascites neoplasia. Recent research showed that dehydrocrotonin (DCTN), an extract of CO, had anti- ulcer effect, especially in ulcers induced by hypothermia, alcohol and pyloric ligation. This effect was supposed to be related to the increase in the release of prostaglandin E2, the non-specific competitive binding of H2 and M receptors of stomach, the reinforcement of mucosal protective action and the weakening of mucosa-imparing factors. So DCTN has the potential to be a new anti-ulcer drug.

Croton is a strong cathartic; trace of croton can cause severe...
diarrhea because of its potent in accelerating gastrointestinal motility. Now, Chinese medicine is the hot spot in the development of new drugs promoting gastrointestinal motility. However, its mechanism in inducing diarrhea and accelerating gastrointestinal motility is not yet clarified.

The aim of this research is to observe the effect of CO on intestinal epithelial cells and isolated colonic smooth muscle cells; investigate the effect of CO on biological properties of intestinal epithelia, mRNA differential display and intestinal smooth muscle cell contraction; and further clarify its molecular mechanism so as to lay the foundation of the development of new drugs promoting gastrointestinal motility.

**MATERIALS AND METHODS**

**Cell lines and culture**

Human intestinal epithelial cell line (HIEC), presented as a gift by Dr. Ren DQ from Department of Radiation Medicine, Fourth Military Medical University, was grown in RPMI 1640 medium with 100mM·L⁻¹ fetal bovine serum. CO dissolved in dimethyl sulfoxide (DMSO) was added into the culture medium when the medium was exchanged. The end-concentration of CO was 4mM·L⁻¹; then the concentration increased progressively to 40mM·L⁻¹ in 6 weeks.

**Effects of CO on HIEC growth (MTT)**

Add HIEC into 96 well plate (200µL/well); and add CO of different concentrations (4, 20, 40, 80 mg·L⁻¹) and control DMSO (0.2, 1, 2, 4mM·L⁻¹) to corresponding groups, with three wells in each group. Put the culture plates into culture cassette, then take out one 96-well plate each time (1, 3, 5, 7d), add 20µL MTT solution (5g·L⁻¹) into each well, and resume culturing at 37°C for 4 h. Afterwards, suck and throw away the culture medium from the 96-well plate and add 150µM DMSO to the plate. Detect OD in each well with photometer (wave length: 490nm) and draw the growth curve.

**Flow cytometry analysis of HIEC proliferation treated with CO**

Harvest HIEC in log phase growth that was treated with CO for 6 weeks. Cell proliferation cycle of HIEC was analyzed with flow cytometry after HIEC was routinely digested, washed twice with PBS and fixed with 700mL·L⁻¹ ethanol. DNA index (DI) was calculated with the formula: DI = experimental group cells G0/1 control cells G0/1 DNA.

Judgment standard of ploidy: DI = 0.95-1.05 (DI = 1 ± CV, CV is 0.05); diploid; DI ± 1.00, between 0.90-1.10: proximal diploid; DI>1.10 or DI<0.9: increased or decreased heteroploid DNA content.

**Morphological changes of HIEC treated with CO**

Changes in cell morphology were observed dynamically under inverted microscope. HIEC in log phase growth that was treated with CO for 6 weeks was harvested, and its proliferation cycle was analyzed with flow cytometry after routinely digested, washed twice with PBS, centrifuged for 5 min at 1200-1400 r·min⁻¹, fixed with glutaraldehyde for 30 min at 4°C, and prepared into ultrathin section for transmission electron microscopic observation.

**Expression of COX-2 mRNA in HIEC(Dot blot)**

Total RNA was extracted from control and experimental cells treated with CO for 6 wk. RNA samples were pointed onto the nitric cellulose membrane, and specific COX-2 oligonucleotide probes were end-labeled. Expression of COX-2 mRNA was detected with dot blot.

**mRNA differential display of HIEC treated with CO(DD-PCR)**

Gene expressions of HIEC treated with CO for 6 weeks and of control were compared through DD-PCR. Primers were designed according to the literature: 3’ end anchor primer H-T11C: 5’- AAGCTT TTTTTTTTTC-3’, 8 strips of 5’ random primers: H-AP, H-APs. The first chain of cDNA was synthesized by reverse transcription; then with RT products as templates, and PCR was performed with 8 strips of 5’ end random primers individually. Products of RT-PCR were run on 60g·L⁻¹ urea degeneration polyacrylamide gels until the indicator arrived at the basal part of the gel at 200-300V for 6-8 h. The gels were stained with modified silver-stain. Differential cDNA fragments were cut, recovered and purified for secondary amplification with PCR, and amplification products were isolated by agarose gel electrophoresis and ligated with pUC-Tm vector after recovery and purification. Recombinant plasmids were transfected into DH5α Escherichia coli and the bacteria were plated and incubated at 37°C overnight. Positive clones were picked out and the plasmids were extracted, followed by restriction enzyme cleavage identification. Further identification of the positive clones included reverse Northern blot and sequencing by automatic sequencer. Homology of the sequences of the differential expression cDNA fragments was searched in Gene Bank Database.

**Direct effect of CO on Guinea pig colonic smooth muscle cells**

Guinea pig colonic smooth muscle cells were isolated by modified methods reported by Bitar and Makhlouf. Activity of the smooth muscle cells was detected with trypan blue, and the number of the cells were calculated under invert microscope. The number of smooth muscle cells with vitality must exceed 90% for experiment. CO was dissolved in dimethyl sulfoxide (DMSO) to make 4, 2, and 0.25 g·L⁻¹ CO solutions. Ten µL CO solutions of different concentrations were added into 200µL colonic smooth muscle cell suspensions, the end-concentration of CO being 0.14, 0.07, and 0.009 g·L⁻¹, respectively. The cells were incubated with CO at 30°C for 30 s and 80µL of 25g·L⁻¹ glutaraldehyde was added to terminate the reaction.

The length change of the smooth muscle cells was detected with micrometer under invert microscope. The average length was calculated from every 50 cells randomly. The contractile response of the smooth muscle cells was measured by the decrease of average length of cells treated with CO compared with that of the control. The formula is:

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\text{Percentage of cell contractile change} = \left( \frac{\text{average length of experimental cells} - \text{average length of control cells}}{\text{average length of control cells}} \right) \times 100\%.
\]

**RESULTS**

**Effect of CO on the growth and cell cycle-phase of HIEC**

High dosage (80 mg·L⁻¹) CO resulted in the death of all HIEC; moderate dosage (20-40 mg·L⁻¹) significantly inhibited cell growth (1.3, 5, 7d OD sequence: (20 mg·L⁻¹) 0.040±0.003, 0.081±0.012, 0.147±0.022, 0.024±0.016; (40 mg·L⁻¹) 0.033±0.044, 0.056±0.012, 0.104±0.010, 0.189±0.006; OD control 0.031±0.008, 0.096±0.012, 0.173±0.009, 0.300±0.016, P<0.01) in a dose-dependent manner (IC₅₀ is 52 mg·L⁻¹); and low dosage (4 mg·L⁻¹) had no effect on cell growth (P>0.05). The flow cytometry showed that S phase HIEC quantity significantly increased, G1/M phase cell quantity decreased and DI rose to 1.15 after HIEC was treated with CO for 6 weeks.

**Changes of HIEC morphology treated with CO**

Morphologic changes of HIEC were dynamically observed under inverted microscope in cells incubated with CO. Initially many cells exhibited high fluorescence intensity, however, some cells showed low intensity and some even were not stained.
sheet shedding, intercellular space increased, toxic granules accumulated in the cytoplasm, and cell growth was significantly delayed. The growth behavior of the cells improved after treatment with CO for 11d. After the dosage of CO was progressively increased to 40 mg·L−1 at the 6th week, the following phenomena occurred: fusiform cells significantly increased, cell bodies shrank, the nuclei were enlarged, the cytoplasm was reduced, karyoplasm lopsidedness was developed, and cell growth was accelerated. Under transmission electron microscope, we observed many wide apophyses in cell membrane, abundant microvilli, enlarged nuclei, multiplied nucleoli, and abnormal nuclei with pathologic mitosis (Figure 1).

**Expression of COX-2 mRNA in HIEC**

Expression of COX-2 mRNA was significantly down regulated in HIEC after treated with CO for 6 week (Figure 2).

**DD-PCR**

Some cDNA fragments found in HIEC treated with CO were up-regulated, down-regulated, newly born or deleted in comparison of experimental cells with controls (Figure 3). Thirteen different cDNA fragments were successfully cloned; among the 13 positive clones, 2 fragments were significantly up-regulated and 11 down-regulated in HIEC treated with CO, which was confirmed through reverse Northern blot analysis. Sequencing and homology demonstrated that one over-expressed cDNA fragment is 100% homologous with the homeobox gene of mitochondrial cytochrome C oxidase subunit-2 gene.

**Direct effect of CO on guinea pig colonic smooth muscle cells**

Free Guinea pig colonic smooth muscle cells appeared in fusiform shape with diverse length. Some of them were relaxed while others were at different phases of contraction (Figure 4). The length of normal guinea pig colonic smooth muscle cells ranged from 60µm to 160µm, with an average length being (102±23)µm. After incubated with different concentrations of CO (0.14, 0.07, 0.009g·L−1), the average length of colonic smooth muscle cells became (72±27)µm, (74±26)µm and (72±17)µm, respectively, while that of the control cells was (104±22)µm (Figure 5). The percentages of decrease in cell length were 29.5%, 28% and 29.6%, respectively, significantly smaller than that of the control (P<0.05).

**DISCUSSION**

Our study showed that high dosage of CO could result in the death of HIEC; middle dosage could inhibit cell growth in a dose-dependent manner; low dosage had no effecton cell growth. Meanwhile, after long-term continuous incubation with CO, the cycle from G1 to S of HIEC was shortened, the quantity of G1 phase cells was reduced, the quantity of S phase cells was increased, and DI was raised. We found that the principal phenomena of HIEC treated with CO included cell degeneration, necrosis, diminished ability of cells adhesion, and delayed cell growth. As the drug action persisted, degenerated cell quantity decreased and cell proliferation was accelerated. We also observed that HIEC treated with CO proliferated faster than control cells, and abnormal nuclei with pathologic mitosis were observed under electron microscope. These phenomena indicated that the cytotoxicity of high concentration CO might be the major cause for cell growth suppression. CO had the potential effect on promoting cell over-proliferation and increasing the content of heteroploid DNA when HIEC was incubated with CO for a long time.
COX-2 is an isoenzyme of cyclooxygenase discovered recently. Under normal conditions, COX-2 expression was low or non-existent in most tissues and cells in human body, but its expression could be promptly enhanced when cells were stimulated by a variety of factors. Recent studies showed that COX-2 is closely related to the development and progression of many kinds of human tumors, especially gastrointestinal tumors, such as colon cancer and gastric cancer. It has been confirmed that the expression of COX-2 is non-existent in most tissues and cells in human body, but its expression could be promptly enhanced when cells were stimulated by a variety of factors. Recent studies showed that COX-2 is closely related to the development and progression of many kinds of human tumors, especially gastrointestinal tumors, such as colon cancer and gastric cancer. It has been confirmed that the expression of COX-2 increases more significantly in colon cancer tissues than in normal colon tissues, and its expression is higher in many kinds of gastric cancer cell lines and tissues than in their normal counterparts. The incidence and mortality of colon cancer were decreased with the long-term intake of non-steroidal anti-inflammatory drugs (NSAIDS), a COX-2 inhibitor, which is effective on familial adenomatous polyposis. These data indicated that COX-2 might play an important role in the pathology of these tumors. The specific mechanism of COX-2 induced tumorigenesis is yet unclear, but it is suspected to be related to the following factors: catalysis of carcinogen synthesis; catalysis of prostaglandin (PG) synthesis, which can regulate the expression of genes and accelerate DNA synthesis and cell proliferation; inhibition of cell apoptosis; enhancement of adhesion between cells and extracellular matrix; promotion of tumor angiogenesis; inhibition of immunological response. The expression of COX-2 mRNA was detected in HIEC treated with CO for 6 weeks, and the results revealed its significant down-regulation. The relationship between CO-induced tumorigenesis and down-regulation of COX-2 in HIEC remains to be elucidated.

Thirteen differential expression gene fragments were detected in HIEC treated with CO for long time through mRNA differential display. Therefore, these genes may be related to the biological effects of CO on HIEC. One of the genes is human mitochondria cytochrome C oxidase subunit II gene, whose expression is high in CO treated HIEC. Human mitochondria cytochrome C oxidase consists of subunit I, II and III, which contributes substantially to the respiratory function of mitochondrion. In addition, two notably highly expressed genes were separated from Cisplatin-resistant human head and neck squamous epithelium cancer by Higuchi through DDRT-PCR, and one of them is again the human mitochondria cytochrome C oxidase subunit II gene, which is putatively related to Cisplatin-resistance of tumor cells. We have also separated the highly expressed mitochondria cytochrome C oxidase subunit II gene from vincristine-resistant human gastric cancer cell SGC7901/VCR by DDRT-PCR (to be published separately). This suggests that mitochondria cytochrome C oxidase subunit II gene is closely correlated with drug-resistance. We found that CO-resistance of HIEC could be induced by gradually increasing the concentration of CO. However, the mechanism of CO-resistance in HIEC is unclear. As normal intestinal epithelial cells express certain multidrug resistance genes and proteins, such as P-glycoprotein (P-gp) in small intestinal epithelial cell line, it demands further investigation about whether the CO-resistance is mediated by P-gp or mitochondria cytochrome C oxidase subunit II gene product.

Croton possesses many kinds of drug actions; however, its clinical application is largely limited to treating constipation, while its effect in promoting gastrointestinal motility is neglected. The neglect is not only due to its complexity in components and severe side effects, but also to the ignorance of its molecular mechanism in promoting gastrointestinal motility. Pol found that CO could accelerate gastrointestinal movement of mice at 3h after gastric administration. When CO was administrated to dogs, it could quickly evoke a rapid complex wave similar to the interdigestive myoelectric complex (IMC) III phase, although its contractile time was shorter its velocity of conduction faster than normal IMC. However, it was not affected by vague nerve, which showed no relationship with α and ( receptors. IMC cycle shortening induced by CO was the principal cause of diarrhea. Albuquerque made a series of gastrointestinal movement experiments on animals with CO. The results showed that CO and its preparations could directly affect intestinal smooth muscles and rat diaphragm muscles and cause them to contract. Other researchers reported that senna could induce diarrhea and accelerate gastrointestinal movement by enhancing the synthesis and release of endogenous prostaglandin E2 (PGE2). Although we do not know whether the gastrointestinal motility enhancement of CO is related to PGE2, the significant down-regulation of COX-2 expression in HIEC with long-term treatment of CO indicated that the mechanism of CO-promoted motility is different from that induced by senna. Furthermore, our experiment confirmed that the average length of colonic smooth muscle cells was significantly shortened when they were incubated with CO of different concentrations, which showed that CO had a direct contractile effect on colonic smooth muscle cells.
