Clostridium difficile toxin A excites enteric neurones and suppresses sympathetic neurotransmission in the guinea pig

Y Xia, H Z Hu, S Liu, C Pothoulakis, J D Wood

Abstract

Background and aims—Evidence suggests that the intestinal actions of Clostridium difficile toxin A—stimulation of secretion and motility, and an acute inflammatory response—have a neurally mediated component.

Methods—Direct intracellular electrophysiological recording of electrical and synaptic behaviour in enteric neurones was performed in the submucous plexus of guinea pig small intestine during exposure to the toxin.

Results—Application of toxin A affected both the electrical behaviour of the neuronal cell bodies and inhibitory noradrenergic neurotransmission to the cell bodies. Altered electrical behaviour included depolarisation and increased excitability. Tetrodotoxin or a histamine H1 receptor antagonist did not affect the depolarisation evoked by toxin A. Failure of the histamine antagonist to suppress the actions of toxin A is evidence that its actions were not mediated by degranulation of intramural mast cells. The action of toxin A on neurotransmission was suppression of inhibitory postsynaptic potentials evoked in the neuronal cell bodies by stimulation of sympathetic nerve fibres that synapsed with the cell bodies. The inhibitory postsynaptic potentials were mediated by norepinephrine (noradrenaline) acting at postsynaptic alpha adrenoceptors on the cell bodies. Hyperpolarising responses evoked in the cell bodies by micropressure application of norepinephrine were unaffected by toxin A. This fulfils criteria for a presynaptic inhibitory action of toxin A to suppress release of norepinephrine from sympathetic postganglionic axons.

Conclusions—Results suggest that the neural component of the action of toxin A involves both direct excitation of enteric neurones and suppression of norepinephrine release from postganglionic sympathetic nerve fibres in the enteric nervous system.

(Gut 2000; 46: 481–486)

Correspondence to: Dr J D Wood, Department of Physiology, 302 Hamilton Hall, 1645 Neil Avenue, Columbus, Ohio 43210, USA

Department of Physiology, The Ohio State University, College of Medicine and Public Health, Columbus, Ohio, USA

Y Xia

Division of Gastroenterology, Beth Israel Deaconess Medical Center, Harvard Medical School, Boston, Massachusetts, USA

C Pothoulakis

Accepted for publication 3 November 1999

Keywords: enteric nervous system; enterotoxins; diarrhoea; intestine

Abbreviations used in this paper: EPSP, excitatory postsynaptic potential; IPSP, inhibitory postsynaptic potential; TX-A, C difficile toxin A; TTX, tetrodotoxin.
were removed 20 cm orad to the ileocecal junction. Preparations of the submucous plexus for electrophysiological recording were microdissected as described previously.12 The preparations were mounted in a 2.0 ml recording chamber that was perfused at a rate of 10–15 ml/min with Krebs solution warmed to 37°C and gassed with 95% O₂–5% CO₂ to buffer at pH 7.3–7.4. The composition of the Krebs solution was (in mM) NaCl, 120.9; KCl, 5.9; MgCl₂1.2; Na₂HPO₄1.2; NaHCO₃, 14.4; CaCl₂, 2.5; and glucose, 11.5.

Methods of intracellular recording from the submucous plexus are described in detail elsewhere.10 Transmembrane electrical potentials were recorded with conventional “sharp” microelectrodes. The microelectrodes were filled with 4% biocytin in 2M KCl containing 0.05 M Tris buffer (pH 7.4). Resistances of the electrodes were 80–190 MΩ. The intraneuronal marker (biocytin) was injected by the passage of hyperpolarising current. The preamplifier (M767, World Precision Instruments, Sarasota, Florida, USA) had bridge circuitry for injection of electrical current. All data were recorded on videotape for later analysis. Synaptic potentials were evoked by electrical shocks (0.1–20 Hz) applied focally to interganglionic connectives with 20 μm diameter Teflon insulated platinum wire electrodes.

Actions of pharmacological agents and TX-A were studied by pressure microejection or by application in the superfusion solution. Micropipettes (10 μm tip diameter), manipulated with the tip close to the impaled neurone, were used to microinject the substances. Pressure pulses of nitrogen with predetermined force and duration were applied to the micropipettes through electronically controlled solenoid valves. Duration of the pressure pulses could be increased in increments of 1 ms from 0 to 1 seconds. Stepwise increases in pulse duration were used to assess, in a quasimanner, the dependence of TX-A effects on concentration.

Toxin A was purified to homogeneity from broth culture supernatants of *C difficile* strain 10 463 as described previously.13 Norepinephrine (noradrenaline), phenothaline, tetrodotoxin (TTX), and cimetidine were all obtained from Sigma Biochemicals, St Louis, Missouri, USA. Data are expressed as mean (SEM). Student’s *t* test for unpaired values was used to evaluate significance of differences for means of resting membrane potential and neuronal input resistance.

**Results**

**EXCITATORY ACTIONS OF TX-A**

Application of TX-A was by pressure microejection. Limitations on the supply of TX-A because of constraints of isolation in sufficient quantities restricted ability to obtain concentration–response data by adding the toxin to the superfusion reservoir of the recording chamber. The microejection pipettes contained 0.6 mg/ml TX-A in Krebs solution and were positioned with the tips 20–50 μm from the impaled neurone. Dilution of the ejected toxin occurred rapidly in the 2 ml volume of the tissue chamber which was perfused...
at a rate of 10–15 ml/min. Precise determination of effective concentrations (for example, EC<sub>50</sub>) was not possible with this method.

TX-A, applied by micropressure ejection, evoked excitatory responses in 23 of 24 neurones (nine AH type and 14 S type cells). Micropressure pulses of Krebs solution alone evoked no responses. The effects of TX-A consisted of membrane depolarisation coincident with increased input resistance. Augmented excitability was apparent as spontaneous spike discharge or repetitive discharge during intracellular injection of depolarising current pulses (fig 1). These effects were accompanied by suppression of the characteristic hyperpolarising after potentials in AH type neurones. The amplitude of the depolarising responses increased with increased duration of pressure ejection pulses ranging from 20 to 110 ms (fig 2). This was indicative of concentration dependence of the responses. The effects were reversed as TX-A was washed from the recording chamber.

The excitatory responses to TX-A were reminiscent of excitation of submucous neurones produced by degranulation of antigen sensitised mucosal mast cells. One of the mediators released by mast cell degranulation is histamine, which acts at the histamine H<sub>2</sub> receptor subtype to depolarise and enhance excitability of neurones in the guinea pig submucous plexus.

We used the H<sub>2</sub> histamine receptor antagonist, cimetidine, to determine whether the excitatory responses to TX-A might be a secondary effect of degranulation of mast cells and the release of histamine. Cimetidine is known to block the excitatory actions of exogenously applied histamine and of histamine released by mast cells during antigen exposure in sensitised guinea pig bowel in vitro.

Pretreatment with cimetidine blocked the neuronal excitatory responses to micropressure pulses of histamine, but did not change significantly the depolarising responses to TX-A (fig 3). This suggested that the excitatory action of TX-A was directly on the neurones and not secondary to mast cell activation by the toxin. Excitatory responses evoked by TX-A (membrane depolarisation) were also unaffected by pretreatment with TTX (fig 3). This suggested direct action of the toxin because TTX was expected to block axonal conduction in the plexus and thereby prevent input to the recorded neurone from synaptically connected neurones elsewhere in the microcircuit.

Figure 4 Focal electrical stimulation of sympathetic noradrenergic synaptic inputs to submucous plexus neurones evoked inhibitory postsynaptic potentials (IPSP). (A) Application of C difficile toxin A (TX-A) suppressed the stimulus evoked IPSP. Suppression of the IPSP was concentration dependent as reflected by a progressive decrease in the amplitude of hyperpolarisation with progressively increased duration of micropressure pulses ("spritz") from 20 to 80 ms in a single neurone. (B) Quantitative concentration–response data for eight neurones. The neurone in A had uniaxonal morphology.
TX-A SUPPRESSION OF INHIBITORY POSTSYNAPTIC POTENTIALS

Focal electrical stimulation of sympathetic noradrenergic synaptic inputs to the submucous neurones evoked inhibitory postsynaptic potentials (IPSPs) (fig 4A). Micropressure application of norepinephrine mimicked the hyperpolarising component of the IPSPs (fig 5A). Blockade of the IPSPs (fig 5D–F) by the alpha adrenoceptor antagonist, phentolamine, was consistent with earlier reports that the noradrenergic IPSPs in guinea pig submucosal

Figure 5  The alpha adrenergic blocking drug, phentolamine, blocked both the hyperpolarising responses to micropressure application of norepinephrine (noradrenaline) and stimulus evoked inhibitory postsynaptic potentials (IPSP). (A) Micropressure (“spritz”) application of 10 µM norepinephrine evoked a hyperpolarising response accompanied by decreased input resistance. Decreased input resistance was reflected by decreased amplitude of electrotonic potentials produced by intraneuronal injection of constant current depolarising pulses. (B) Blockade of norepinephrine evoked response by phentolamine. (C) The neurone in A and B had uniaxonal morphology. (D) Focal electrical stimulation of an interganglionic connective evoked an IPSP. (E) Blockade of the IPSP by phentolamine. (F) Reversal of phentolamine induced blockade following washout of the drug. (G) The neurone in D and F had uniaxonal morphology.

Figure 6  Toxin A (TX-A) suppressed stimulus evoked noradrenergic inhibitory postsynaptic potentials (IPSP) but not the hyperpolarising responses to exogenously applied norepinephrine (noradrenaline). (A) Control IPSP. (B) Suppression of IPSP by micropressure pulse of TX-A. (C) Recovery of IPSP after washout of TX-A. (D) Hyperpolarising response to micropressure pulse of norepinephrine. (E) Hyperpolarising response to norepinephrine during exposure to TX-A was unchanged. (F) Hyperpolarising response to norepinephrine after washout of TX-A. All records are from the same neurone. Downward deflections in D–F are electrotonic potentials produced by intraneuronal injection of constant current hyperpolarising pulses.
plexus neurones are mediated by alpha noradrenergic receptors. Application of TX-A from microejection pipettes reversibly suppressed or abolished the stimulus evoked IPSPs (fig 4A). Suppression of the IPSPs was concentration dependent as determined by progressively increasing the duration of micro-pressure pulses of TX-A (fig 4A,B). Blockade of the IPSPs was accompanied by an increase in spontaneously occurring fast excitatory postsynaptic potentials that sometimes reached threshold for action potential discharge (fig 4A). The excitatory postsynaptic potentials probably reflected input from synaptically connected neurones that were excited by the toxin. Micropressure application of norepinephrine evoked phenolamine sensitive hyperpolarising responses associated with decreased input resistance in the submucous neurones (fig 5A,B; fig 6D–F). Application of TX-A had no effect on the hyperpolarisation evoked by norepinephrine while it suppressed the stimulus evoked IPSP in the same neurones (fig 6A–C). In eight neurones, the mean hyperpolarisation to 20 ms duration micropressure pulses of norepinephrine was 30 (2.4) mV in the absence of TX-A and 29.3 (2.3) mV when preceded by pulses of TX-A that suppressed the IPSP. This suggested that the site of action of TX-A was at presynaptic inhibitory sites on the noradrenergic nerve terminals, and that suppression of the IPSPs resulted from inhibition of norepinephrine release from the terminals.

**Discussion**

The results suggest that purified *C difficile* TX-A has direct actions on neurones in the submucous plexus of guinea pig small intestine in relatively low concentrations. Nevertheless, absolute concentration data could not be determined precisely in the present study because of limitations on the amounts of available toxin. Two kinds of actions were revealed. One was an excitatory action at the level of the membranes of the neuronal cell bodies that was reminiscent of the excitatory action of cholera toxin on submucous neurones. The excitatory action of TX-A mimicked slow synaptic excitatory responses characteristically found in enteric neurones. Excitation evoked by TX-A also mimicked the slow excitatory effects produced by paracrine release of histamine from mucosal mast cells. Failure of histamine receptor antagonists and blockade of axonal conduction by TTX to suppress the excitatory responses to TX-A are evidence for direct action at the somal membranes of the neurones. The excitatory action could not be attributed to activation of other neurones that provided synaptic input to the recorded neurone or to secondary effects of mast cell degranulation.

The second action of TX-A was suppression of IPSPs evoked by stimulation of release of norepinephrine from sympathetic postganglionic nerve fibres. This appeared to be a direct presynaptic action that suppressed release of norepinephrine from the noradrenergic terminals because the toxin did not suppress the hyperpolarising action of exogenously applied norepinephrine. The presynaptic inhibitory action of TX-A differed from that of cholera toxin which was reported not to suppress IPSPs in submucous neurones.

An aspect of the pathophysiological significance of the neuronal action of TX-A is illustrated in fig 7. Secretomotor neurones to the intestinal crypts are located in the submucous plexus. When secretomotor neurones fire, they release vasoactive intestinal polypeptide and/or acetylcholine at the neuroepithelial junctions and this stimulates the secretion of water and electrolytes into the crypt lumen. Hyperactivity of the secretomotor neurones is associated with a state of secretory diarrhoea. Two neuronal actions of TX-A are expected to be involved in production of the secretory state. One is direct excitation by the toxin of the secretomotor neurones and/or interneurones that provide excitatory synaptic drive to the secretomotor neurones. The other is the inhibitory action of TX-A on noradrenergic transmission to the secretomotor neurones and/or synaptically connected interneurones. This effectively nullifies any sympathetic braking action on the secretomotor neurones to permit maximal secretomotor firing rates and hyperstimulation of mucosal secretion.

Aside from submucous secretomotor neurones, a major component of sympathetic input to the intestine acts at presynaptic terminals to prevent release of excitatory transmitter substances that mediate fast or slow transmission at synapses within the integrated circuits of the enteric nervous system. These axo-axonal synapses, which have been described ultrastructurally, function to inactivate the excitatory synaptic circuitry that mediates intestinal motility. Application of norepinephrine or electrical stimulation of sympathetic postganglionic fibres in the intestinal mesentery reduces or abolishes fast nicotinic transmission...
in the enteric networks. This occurs without any change in the somal membrane properties during exposure to norepinephrine and without any change in the depolarising responses of the neurones to microinjected acetylcholine. Both findings are evidence that the mechanism of action of norepinephrine is blockade of release of acetylcholine from presynaptic nerve terminals. The presynaptic inhibitory receptors have been identified as the \( \alpha_2 \) adrenoceptor subtype. Based on the neurophysiology of the synaptic interface between the sympathetic and enteric divisions of the autonomic nervous system, the significance of TX-A induced blockade of norepinephrine release at excitatory synapses is expected to be prevention of sympathetic inactivation of the enteric microcircuits that generate intestinal motor activity and other intestinal behaviours during \( C \) difficile enteritis.

This work was supported by National Institutes of Health grants R01 DK-37238 and R01 DK-46941 to JDW.

Clostridium difficile toxin A excites enteric neurones and suppresses sympathetic neurotransmission in the guinea pig

Y Xia, H Z Hu, S Liu, C Pothoulakis and J D Wood

Gut 2000 46: 481-486
doi: 10.1136/gut.46.4.481

Updated information and services can be found at:
http://gut.bmj.com/content/46/4/481

These include:

References
This article cites 20 articles, 6 of which you can access for free at:
http://gut.bmj.com/content/46/4/481#BIBL

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Topic Collections
Articles on similar topics can be found in the following collections
Clostridium difficile (67)
Gastrointestinal hormones (848)

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/