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MECHANISM OF ACTION OF TETANUS TOXIN

ANNUAL\FINAL REPORT

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of TT on any lysosomal protein. and a mitochondrial substrate, is an effect of TT on mitochond idrenal mitochondria. However, by using TT-specific gangliosid bonoclonal antibodies. We conc whosphate salts in the TT prepa D. DISTRIBUTION/AVAILABILITY OF ABSTRAC.	we showed that ria. We confirm we were then un e affinity chrom lude that the de ration which fac	the stimulat ed this find able to show atography, a scribed TT e ilitated Ca+ 21. ABSTRACT SEC Unclassifi	cling of ATP-cling by using any specifi s well as in ffect was du transport	lependent g purified lcity for munoabsor ie to the by mitoch	Ca++ uptake bovine the TT actio ption with presence of condria.

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In the previous annual report, we described our finding that tetanus toxin (TT) irreversibly stimulates ATP-dependent  $Ca^{\pm\pm}$  uptake by human neutrophil lysosomes and we hypothesized that the toxin acts as a kinase capable of either directly phosphorylating the neutrophil lysosomal  $Ca^{\pm\pm}$  ATP-ase or indirectly activating it by phosphorylation of a regulatory intermediate.

Numerous phosphorylation studies on intact neutrophil lysosomes as well as on lysosomal membranes failed to reveal any effect of TT on protein phosphorylation or dephosphorylation.

We then asked the question whether TT acts on lysosomes, since our neutrophil "lysosomal" preparation is actually a microsomal one that consists of two intracellular organelle populations, lysosomes and mitochondria (1). To determine which organelle was affected by TT, we performed Ca<sup>++</sup> uptake experiments using mitochondrial inhibitors.

As shown in Figure 1, either one or both of the mitochondrial inhibitors azida (5 ml) and atractyloside (100 ull) partially inhibited the ATP-dependent Ca<sup>++</sup> uptake of FNN microsomes, indicating that a significant portion of Ca<sup>++</sup> uptake by the proparation was due to mitochondria. The stimulatory effect of TT (5 ug/ml), however, was completely abolished by either one or both azide and atractyloside (Figure 1), strongly suggesting that TT exerts its effect on mitochondria.

Further evidence was provided by the use of the mitochondrial substrate, succinate. As shown in Figure 2, succinate (1 mi) increased the microsomal ATP-dependent  $Ca^{++}$  uptake, which was further increased by the presence of TT (5 ug/ml). Since TT stimulates the succinate-driven increase in ATP-dependent  $Ca^{++}$ uptake, and since only mitochondria can utilize succinate to energize  $Ca^{++}$  uptake, we concluded that TT acts on the mitochondria in our preparation.

To confirm this conclusion, we purified mitochondria from bovine adrenal cortex since previous work demonstrated that adrenal cells could be intoxicated by intracellular injection of TT. Highly purified preparations of bovine adrenal cortex mitochondria were prepared according to well characterized methods (2-5). As illustrated in Figure 3, Ca<sup>++</sup> uptake by adrenal cortex mitochondria in the presence of ATP (1 ml) was increased by increasing concentrations of TT. At 1 ug/ml 1F, the ATP-dependent Ca<sup>++</sup> uptake was significantly increased.

Our next step was to show that ATP-dependent Ca<sup>++</sup> uptake by isolated mitochondria was inhibited by either one or both of the mitochondrial inhibitors anide and atractyloside. As shown in Figure 4, either one or both azide and atractyloside completely inhibited ATP-dependent Ca<sup>++</sup> uptake activity by mitochondria in the presence or absence of TT (5 ug/ml).

We next investigated whether the effect of TT was supported by the nonhydrolyzable ATP analogue AMP-PNP (6). As shown in Figure 5, TT had no effect on mitochondrial  $Ca^{++}$  uptake in the presence of the ATP analogue, indicating a requirement for a hydrolyzable high energy phosphate bond.

Since we showed that the effect of TT on mitochondrial  $Ca^{++}$  uptake requires ATP, our next step was to investigate the mechanism by which it does so. The most tempting hypothesis was that TI acts as a kinase, phosphorylating a protein(s) that participates in  $Ca^{++}$  uptake by mitochondria, or, acting indirectly, phosphorylates a kinase, which in turn phosphorylates another protein. Since all known protein kinases require ATP concentrations in the micromolar range (7-10), the results shown in Figure 6 argue against TT having kinase activity. The studies demonstrated that

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IT stimulated ATP-dependent  $Ca^{++}$  uptake by bovine adrenal cortex mitochondria only at 1 mH and at 0.5 mH ATP. There was little activity of TT at 100 uM ATP and no effect at 10 uH ATP, indicating that TT is not a typical protein kinase.

We next attempted to show a specificity for the stimulatory effect of tetanus toxin (IF) on AIP-dependent  $Ca^{++}$  uptake by mitochondria. To our surprise, boiled toxin, which is biologically inactive, showed the same stimulatory effect. Also, TT dialyzed against the buffer used for the  $Ca^{++}$  uptake experiments showed no stimulatory effect on mitochondrial  $Ca^{++}$  uptake, at least by bovine adrenal mitochondria. A possible explanation is that the active factor is inorganic phosphate  $(F_4)$ , which is abundant in our TT stock preparation, since its buffer is phosphate buffored saline (FBS), and since P<sub>1</sub> is known to activate mitochondrial Ca<sup>++</sup> uptake. When we measured the P<sub>4</sub> concentration of our TT stock preparation, it was found to be 3 ml. We then tested various concentrations of  $P_1$  on mitochondrial ATP-dependent  $Ca^{++}$  uptake and found that 100 uM showed a stimulatory effect and 10 vM had no effect. According to our calculations (from the dilutions of the TT stock we are using) and to the above P<sub>i</sub> measurement, the final concentrations of F; in our experiments are always below 30 uM, and therefore should not have a notable effect on ATP-dependent Ca<sup>++</sup> uptake. Furthermore, scrial dilutions of FBS, equivalent to the dilutions of the TT stock we are using, were compared to the equivalent dilutions of TT. The PBS dilutions increased ATP-dependent Ca<sup>++</sup> uptake, as was expected (since PBS contains  $P_i$ ), but the increases were small compared to the corresponding increases induced by the TT preparation. These results indicated that the observed effect of TT is probably not attributable to the contamination of our TT preparation by phosphate salts.

After the results of the experiments described in the previous paragraph, we realized that we had to show some specificity for the TT effect, and we performed two groups of experiments.

In the first one, we applied our TT preparation to a TT-specific ganglioside  $(GF_b)$  column. We then tested the material eluted from this column. Although there was no TT detectable by protein gel electrophoresis, this material showed the same activity as control TT preparations in increasing ATP-dependent Ca<sup>++</sup> uptake by either human neutrophil microsomes or bovine adrenal cortex mitochondria.

In the second group of experiments, we eliminated the toxin from our preparation by immunoabsorption using protein A sepharose beads and the anti-TT monoclonal autibodies provided by Dr. William Habig. Unfortunately, although all of the toxin present in our preparation was immunoabsorbed as confirmed by protein gel electrophoresis, the material still had the same activity as control TT preparations in augmenting ATF-dependent Ca<sup>++</sup> uptake by either human neutrophil lysosomes or bovine adrenal cortex mitochondria.

The last two groups of experiments confirmed that the effect of IT we have been observing was artefactual. We suspect that this was due to contamination of our TT proparation probably by phosphate salts. We do not wish to further investigate what factor caused the increased ATP-dependent  $Ca^{++}$  uptake, since we know that it definitely is not tetanus toxin.

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## EFFECT OF TETANUS TOXIN IN THE PRESENCE OF SUCCINATE



ATP

ATP+succinate







FIGURE 6 DEPENDENCE OF THE EFFECT OF TETANUS TOXIN ON THE CONCENTRATION OF ATP 10 · ATP-dependent calcium uptake (nmol/min/mg protein) 8 🔲 control Π 🖬 6 4 2 0 2 10 100 500 1000

ATP concentration (µM)