

of blood after bleeding^{15,16}, the doses of vasopressin employed in our experiments will be within the physiological limits.

After bleeding, a significant decrease of the blood flow in the skin has been observed by SAPIRSTEIN et al.¹⁷ and TAKÁCS et al.¹⁸ in the rat. Possibly this phenomenon is caused by a release of vasopressin. This hypothesis is supported by two observations: a significant increase of the vasopressin level in the blood has been observed after bleeding^{15,16}; on the other hand, dogs tolerate bleeding less well after neurohypophysectomy¹⁹.

Zusammenfassung. Vasopressin in den Blutdruck nicht erhöhenden geringen Dosen (0.01 E/kg i.p.) verringert die Hautdurchblutung der Ratte und erhöht die zirkulatorische Resistenz. In höheren Dosen von sogar 1.0, 0.1 E/kg

beeinflusst Vasopressin Minutenvolumen und Zirkulation anderer Organe nicht.

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The Neurogenic Activity of High Potency Substance P

There appears to be general agreement that 'crude' substance P (SP) has an action on the nervous system¹⁻⁶. However, there appears to be some disagreement as to whether the neurotropic action is retained with purification. Thus, STERN and HUKOVIĆ⁷ have reported that as SP is purified up to 270 U/mg, it retains its ability to antagonize morphine-induced analgesia, but loses its ability to produce tranquilization. On the other hand, HAEFELY and HÜRLIMANN⁸ have stated that 'purified' SP is without neurotropic activity.

This paper relates research undertaken in an attempt to determine whether the neurotropic activity of SP is lost when the material is purified to a potency of 10,000 U/mg.

Methods and Materials. In order to provide the most efficient utilization of the small amounts of SP available, the test system used was that previously described from these laboratories⁹, i.e. potentiation of the fourth dorsal root potential (DR IV), by substance P in the presence of lysergic acid diethylamide (LSD).

Decerebrate cats were used in all of these experiments. Decerebration and laminectomy were performed under ether anesthesia. Dorsal root potentials (L7 or S1) were evoked at a frequency of 2.5 cps, using stimuli approximately 50% of maximal and 0.05 msec duration. Ether anesthesia was stopped at least 1 h prior to the administration of SP or LSD. No experiment was initiated until the dorsal root potentials were observed to be constant for at least 30 min.

After a control period of 30 min, LSD, 20 µg/kg, was injected i.v. After a period of approximately 10 min, during which time DR IV was seen to increase in amplitude, and then to remain constant at the new level, SP was injected i.v.

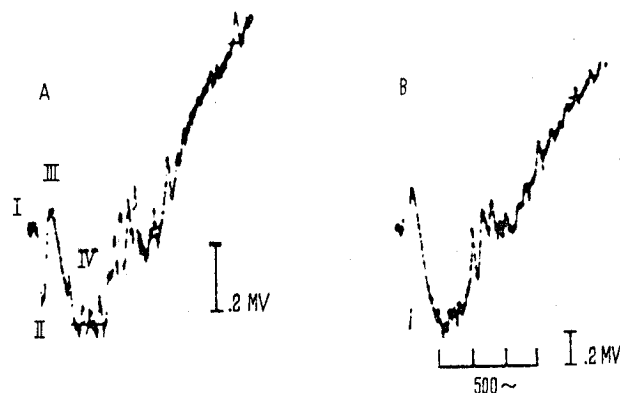
The SP used in these experiments was of three potencies, 10 U/mg, 1,000 U/mg, and 10,000 U/mg⁹, and was purified as described earlier¹⁰.

Results. (1) LSD: In each instance the administration of LSD was followed by an increase in DR IV. This had been described earlier⁹, and no further comment is necessary at this time.

SP: The results of administration of each of the samples of SP can be described together since the effects were qualitatively similar. In each instance SP, 10,000 U/mg, produced a further augmentation of DR IV.

In addition, in certain instances (Figure) this was associated with an increase of the first three dorsal root potentials (i.e. DR I, II, and III). Augmentation of DR I, II, and III with larger doses of SP and presumably consequent to augmentation of DR IV has been observed and discussed in earlier reports⁶.

Using SP of a potency of 10 U/mg, augmentation was seen in three cats using doses ranging from 20-40 U/kg. Using SP of a potency of 1,000 U/mg, the phenomenon



Sample traces of records from the same experiment illustrating the actions of SP (10,000 U/mg) on the dorsal root potential of the cat. Control record (A), illustrates a typical dorsal root potential (DR I-IV) after the administration of LSD, 20 µg/kg, 6 min prior to trace B, SP, 20 U/kg, was injected. Note the change in calibration of the vertical amplifier.

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² G. ZETLER, Arch. Path. Pharmacol. 228, 513 (1956).

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was observed in four cats, the dose range being 30-40 U/mg. Giving SP 10,000 U/mg, the same phenomenon was seen in six cats, the dose range being from 20-70 U/kg. When possible, the duration of these actions of SP was followed and found to be in excess of 30 min. No attempt was made to determine the minimal effective dose.

Discussion. These experiments were undertaken for the experimental purpose of determining if highly 'purified' SP has neurotropic activity, and with the ultimate goal of determining whether or not this activity is lost during purification of the leiotropic fractions, thereby explaining the discrepancy between STERN and HUKOVIĆ⁷ and HAEFELI and HURLIMANN⁸. However, the data presented here show that SP, of relatively high potency, still contains neurotropic activity. Unfortunately, the nature of the test preparation used precludes quantitation of the information obtained, and it is not possible to suggest whether the leiotropic and neurotropic actions are due to the same component.

These data are in support of the conclusions of STERN and HUKOVIĆ⁷ that purified SP has neurotropic activity¹¹.

Zusammenfassung. Gereinigte Präparate von Substanz P mit Aktivitäten von 10, 1000 und 10000 E/mg verstärken das vierte Potential der dorsalen Nervenwurzel im Rückenmark der Katze (Verabreichung nach LSD). Unsere Befunde bestätigen frühere Beobachtungen und legen die Vermutung nahe, dass bei fortschreitender Reinigung ein Teil der neurotropen Aktivität zusammen mit der leiotropen angereichert wird.

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PRO EXPERIMENTIS

Microdetermination of Magnesium and Calcium in Animal Tissue

In connection with studying the metabolism of kidney cortex slices, the necessity arose to determine microgram quantities of Mg and Ca in this material. None of the known methods can satisfactorily remove all the difficulties associated with such determination, which necessitates complete separation of Mg and Ca from each other, and of the two from phosphate and from interfering heavy metals, in particular Fe.

The method presented here is based on the separation of metals on ion exchanger¹. The technique of purifying the resin and the effect of cross-linkage on the strength of the bond of the two metals to the resin was assessed.

Experimental. Reagents: The water used was redistilled from a quartz apparatus. The solutions were kept in polyethylene bottles.

5 × 10⁻³ M EDTA.

Buffer, pH 10.5, 1 M NH₄Cl-NH₄OH.

Buffer, pH 5.6 (20 ml 1 N CH₃COOH + 180 ml 1 N CH₃COONa).

0.06% methanolic solution of Eriochrome Black T or Calcon.

0.1% ethanolic solution of 8-hydroxyquinoline.

0.018% aqueous solution of sodium naphthalohydroxamate².

Dowex 50-X8 (200-400 mesh).

All chemicals were of analytical purity.

Method. The method was tested on standard solutions and applied to the determination of Mg and Ca in the ash of rabbit and rat kidney cortex.

Dry Dowex 50 was left to swell for 24 h and then washed several times by decantation with water. The resin was transferred into 5 × 90 mm columns. Each column was purified by successive washing with: 80 ml 6 N HCl, 15 ml H₂O, 80 ml 6 N HCl, 15 ml H₂O, 40 ml 6 N HCl, and water to a neutral reaction of the eluate, the rate of flow due to gravity being 5 ml/h. This procedure took several days. The resin was then in the H⁺ cycle and ready for use.

The test solution contained 7.3 µg Mg, 2.4 µg Ca, and further 5 µg Fe, 2 µg Zn, 0.5 µg Cu, 0.1 µg Mn and 0.523 mg KH₂PO₄ in 1 ml.

Elution of Mg and Ca. Three types of Dowex 50 with different cross-linkage were investigated for the separation of Mg and Ca. With increasing cross-linkage the strength of bond of Ca rose more rapidly than that of Mg so that, whereas both metals were eluted with 2 N HCl in close sequence from Dowex 50-X4, Dowex 50-X8 permitted elution of Mg selectively with 5 ml 2 N HCl, no Ca being present in the subsequent 4 ml of the eluate and appearing only after elution with 5 ml 3 N HCl. On the other hand, the elution zones of Mg and Ca on Dowex 50-X12 were wide and diffuse.

Determination of Mg and Ca. After adding an aliquote of the test solution, the column of Dowex 50-X8 was washed with 5 ml 1 N HCl; Mg was eluted with 5 ml 2 N HCl and Ca with 5 ml 3 N HCl. The eluates were collected in quartz tubes and evaporated in an electric oven at 100°C. The residues were dissolved in water.

Since the interfering metals are eluted together with Mg, they were removed by adding 0.1% 8-hydroxyquinoline (pH 5.6, acetate buffer) and extracting the chelates formed with chloroform³. Mg was then determined in the aqueous phase by titration with EDTA, using an Agla microsyringe, to Eriochrome Black T (pH 10.5, ammonium buffer)⁴. The equivalence point was determined graphically from curves of extinction changes followed on a Hilger Spekker photocolormeter at 650 mµ.

Calcium was determined photometrically with sodium naphthalohydroxamate at 390 mµ (CF 4 Spectrophotometer) by the indirect method² or by complexometric titration to Calcon (pH 12.5, diethylamine)⁵ similarly as

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