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EFFECT OF EXOGENOUS CALCIUM IONS ON PROSTAGLANDINS OUTPUT FROM RAT HEPATIC TISSUE SLICES

By

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INTRODUCTION

The requirement of extra and/or intracellular CQ++ to stimulate prostaglandins synthesis varies in different tissues.

Poyser, (1985), elicited that prostaglandins synthesis and efflux from their active biosynthetic sites in almost tissues is dependent upon mobilization of Ca++ from intracellular stores. The extracellular Ca++ is essential for release of Ca++ from intracellular stores and essential for replenish these stores., Rilley & Poyser, (1987).

Ernest et al., (1983), reported that the activity of phospholipase seems to be increased by the presence of Ca++. PLA2 is extremly important in the release of arachidonic acid, (which is the most abundent of prostaglandins precursors in almost all the tissues), followed by a rise of prostaglandins biosynthesis. So PLA2 is considered as a regulating and triggering enzyme for prostaglandins biosynthesis.

Prostagladins are not stored preformed but are synthesized and released as required. Therefore the increased efflux reflects the increased biosynthesis and not release of prostglandins as endogenous constituents Ramwell & Shaw, 1970 and Olley & Coceani, (1980).

Synthesis of PGF₂ & may be derived from PGE₂ and/or from PGD₂. The interconversion between PGF₂ & and PGF₂ is dought, Yamamoto, (1983), reported that there is no a direct interconversion between PGE₂ and PGF₂ &. In contrast Ernest et al., (1983) found that reduction of the Ketogroup at C9 in PGE₂ to form PGF₂ & compound is found to take place in guinea pig as well as in human.

The aim of the present work is to study the effect of extracellular Ca++

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dia were used for determination of PGE₂ and PGF₂ ∝ concentraction according to Yalow and Berson (1971), which depends on the competitive binding principles of radio-immunoassay. The Kits were supplied from Dade Bater Travenol Diagnostic Inc.

Statistical analysis:

The results are expressed as ± S. E. M. The data were analysed by unpaired Student's "t" test; the null hypothesis was rejected when P value was leas than 0.05.

RESULTS

Addition of calcium gluconate solution (29 ug/gm wet tissue) resulted in a significant decrease in PGF2 outhigh significant increase in PGE2 output from the hepatic tissue slices to the incubation media.

. I eldst ni betstated in table 1.

DISCUSSION

In this study the addition of calcium gluconate to the incubation media caused a highly significant increase in PGE2 output from the hepatic tissue slices incubated in Kreb's Ringer phosphate buffered solution, this data is coinside with that obtained by Olsen

> on PGE₂ and PGF₂ \propto output from hepatic tissue slices to elucidate the possible role of extracellular Ca++ on the prostaglandins metabolism in hepatic tissue of albino rats.

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Male albino rats weighing 150-200 gms were used in this experimental study. The rats were ted at libidum with milk, bread, carrots and all necesary vitamins and minerals.

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The rats were scarified, the liver tissue slices were excised from the animals, and washed thoroughly with saline. The hepatic tissues were incubated in kreb's Ringer phosphate buttered solution, supplemented by 200 mg/dl glucose to maintain the life of the tissue, according to Maligieri et al., (1975).

Calcium gluconate ampoules 10% produced by Swiss pharma S. A. E. Cairo, were added to the incubation media in a dose of 29 ugm/gm wet tissue.

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After incubation in metabolic shaker for 2 hours at 370C, the liver slices were removed and aliquots of the me-

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et al., (1983), Poyser (1985) and Riley & Poyser (1987), they reported that prostaglandins output is stimulated by Ca⁺⁺ ions. Prostaglandins synthesis and efflux from their active biosynthetic sites are dependent upon mobilization of Ca⁺⁺ from intracellular stores with a direct interaction with the membranes of endoplasmic reticulum, the main site of PLA₂ and prostaglandins synthesizing enzymes. The extracellular Ca⁺⁺ releases the intracellular Ca⁺⁺ from its stores, and is essential for replenish these intracellular stores.

Weis & Malik (1985), elicited that activation of beta- adrenergic receptors will stimulate cardiac PGE₂ and 6keto-PGF₁ synthesis. It is absolutly dependent on extracellular Ca++. Activation of these beta receptors increase transmembrane Ca++ influx, which by activating release of Ca++ from its intracellular stores, stimulatea release of arachidonic acid consequent to activation of phospholipase A₂, making free arachidonic acid available for PGs synthesis.

Recently it has been reported that 6keto PGF1 synthesis in the rat aortic rings elicited by norepinephrine was abolished completely by removal of Ca++ from the mediam and by Ca++ channel blockers, verapamil and nifedipine Stewart et al., 1984).

Cooper and Malik (1986), concluded that norepinephrine requires extracellular as well as intracellular Ca++ to express its rnaximal effect on renal PGs synthesis.

Ca++ that is released from intracellular sites interact with calmodulin. By apecific calmodulin antagonist, rendering calmodulin biologically inactive, basal and stimulated PGE₂ efflux markedly inhibited. In addition calmodulin antagonists might also inhibit PGE₂ efflux by interfering with Ca++phospholipid sensetive protein kinase C. Activation of protein Kinase C might activate release of Ca++ from intracellular stores, thus inducing PGE₂ output, Levis & Weiss, (1976) Picket et al., (1977) and Cooper, & Malik , (1986).

Broekemeier et al., (1985), showed that verapamil exerted a moderate degree of phospholipase A₂ inhibitory activity, whereas the other calcium channel blockers, diltiazem and nifedipine, exhibited only weak inhibition of rat liver mitochondrial phospholipase A₂ activity.

Recently Danon et al., (1986), reported that verapamil, at different concentrations, exerts a dual action on

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hepatic tissue of albino rats.

Addition of calcium ions in the from of calcium gluconate 10% ampoule in a dose of 29 ug/gm wet tissuse to the Kreb's Ringer phosphate media cause a high significant increasre in PGE2 output from hepatic tissue slices. This is associated with a significant decrease in PGF2«. These fingings may give an additional information on the give an additional information on the possible role of Ca++ ions on metabolism of prostaglandins.

From the present study, it is concluded that addition of extracellular Ca++ ions to the incubation media resulted in increase PGE2s synthesis and efflux with decrease PGF2s synthesis and efflux from hepatio tissue of albino rats.

The effect of Ca++ ions on prostaglandins synthesis and release must be put in mined on using drugs which ulter of PG synthesis, efflux or both. An example of these drugs are calgic receptor stimulator as well as alwhich are commonly used in the treatment of many cardio vascular diseases. Their use must be put under observation to avoid undesirable effects as a result of increase or decrease prostaglandins formation.

> cellular phospholipase A₂ activity, thereby stimulating action at low concentration which is extracellular Ca++ dependent and inhibiting action at high concentration and consequently on PGE₂ synthesis by hydronephrotic interstitial cells. While the other calinterstitial cells. While the other calcium channel blocker nitedipine and dilfiazem failed to stimulate PGE₂ syndilfiazem failed to stimulate PGE₂ syn-

Addition of calcium gluconate to the incubation media resulted in a highly significant decrease in PGF2 ~ output, this decrease can be attributed to one of more of the following :

- 1- decrease the synthesis of PGF2 ~ from PGD2.
- 2- increase the conversion of PGF2
 ∞ to PGE2 via stimulation of 9 hydroxydehydrogenase enzyme.
- 3-decrease the conversion of PGE2 to PGF2 & through inhibition of PG-9-Keto-reductase enzyme. Extracellular Ca++ ions may have a role on the activity of these enzymes. This point must be studied in detail in further investigation.

SUMMARY AND CONCLUSION

This vitro study was an attempt to study the possible role of Ca++ ions on metabolism of prostaglandins in Vol. 20, No. 3 & 4 July, & Oct, 1990 Table (1) : Effect of calcium gluconate on PGE2 and PGF2 ∝ output from isolated hepatic tissue incubated for 2 h. at 37°C.

The tested group	PGE ₂ (ng/gm tissue)	PGF ₂ ∝ ng/gm
. Control group mean	29.362	33.625
±S. E. M.	5.058	2.251
. Calcium group Mean	47.451	14.504
±S. E. M.	4.341	2.413
Р	0.05*	0.05*

* Highly significant.

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الملخص العربي

أجرى هذا البحث لدراسة تأثير أيونات الكالسيوم على البروستا جلاندين المستخرج من خلايا كبد الفئران البيضاء.

وقد أدت نتائج البحث إلى أنه اضافة أيونات الكالسيوم إلى محلول كربس رينجر فوسفات المحتضن للخلايا بجرعة ٢٩ ميكروجرام لكل جرام من وزن الأنسجة على هيئة جلوكونات الكالسيوم قد ساعد على زيادة كمية البروستا جلاندين E2 المستخرج من خلايا الكبد زيادة ذات دلالة احصائية عالية. كما أنه أدى إلى نقص كمية البرستاجلاندين تم F2 المستخرج من خلايا الكبد وهذا النقص ذو دلاله احصائية عالية محايدل على أن أيونات الكالسيوم لها دور فعال في عملية تكوين البروستاجلاندينات وتحويل كل من النوعين ته F2 & F2 إلى الآخر.

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