AD-773 679

FAILURE OF METHYLPREDNISOLONE TO PROTECT LEAD-SENSITIZED RATS AGAINST ENDOTOXIN

Robert B. Jones, et al

Naval Medical Research Institute Bethesda, Maryland

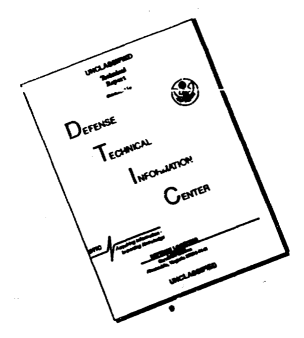
27 June 1973

**DISTRIBUTED BY:** 



National Technical Information Service
U. S. DEPARTMENT OF COMMERCE
5285 Port Royal Road, Springfield Va. 22151

## ISCLAIMER NOTICE



THIS DOCUMENT IS BEST QUALITY AVAILABLE. THE COPY FURNISHED TO DTIC CONTAINED A SIGNIFICANT NUMBER OF PAGES WHICH DO NOT REPRODUCE LEGIBLY.

10 CAL RESEARCH INSTITUTE						
FROM RESEARCH INSTITUTE	1 0110.0	UNCLASSIFIED				
. 1.1. N 20014	Zh. GROU	2h. GROUP				
4 - 6 - 7 - 1 - 7						
TO PROTE	CT BUAD-SENSITIZED R.	FS GGAINST				
1 1 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2						
1 NC PROCEES REPORT						
A A REAL COMMENT, BOOM S. L. WIDE, AND LIT	TZ A. KIESOW					
WELONT - V.:	78. TOTAL NO OF PAGES	7b. NO OF HEFS				
7 (673	75	8				
" p r 14 THAC TO HE GHAN T. NO	98. ORIGINATOR'S REPORT	NUMBER(5)				
h they are the	MR011 .0001 .002 .0007 Report No. 1					
	9 STHER REPORT NO(5) (Any other numbers that may be assigned this report)					
TO THE MINUTION STATEMENT						
	OF THE DIVING A 12 SAME AND A	# 18 7 - <b>**</b> * 2100 - 18 <b>*</b> * 18 * 18 * 18 * 18 * 18 * 18 * 18 *				
HOLDER OF HAS BUEN APPROVED FOR PU	BLUC REDEASE AND SAIR	: 118 PASSABELL				
CE. CME GEAUX GOTES	12 SPONSORING MILITARY	12 SPONSORING MILITARY ACTIVITY				
Nowfiet 1 from: IMPUCTION AND IMMUNITY	1	PURFAU OF MEDICINE AND SUPGERYANCUTE WASHINGTON, D. C.				

Reproduced by
NATIONAL TECHNICAL
INFORMATION SERVICE
U S Department of Commerce
Springfield VA 22151

7D FORM 1473 (PAGE 1)

Ţ

UNCLASSIFIED
Security Classification

Security Classification

Security Classification							
14.	KEY WORDS	LINK A		LINK B		LINK C	
	uel undas	ROLE	₩T	ROLE	wt	ROLE	WT
			!				ı
1.	METHYLPREDNISOLONE	}	}			j	
		<b>'</b>	·		'	! !	
2.	LEAD-SENSITIZED	1	1				
		l	[			1	
3.	?XTS	{	{	(	1	{	
		[	[				
4.	ENDOTOXIN	i	(	į .			
ı		1	(		<b>,</b>		•
		1	}			}	
		1	1		}	}	
		i	1	{			
		1	]		[		
		1	ļ				
		1	i				
		ĺ	ĺ			<b>j</b>	
		į	(		ļ	ļ	
ļ		1	{			ļ	
		{	{			}	
			ſ	1	[	}	
		1	İ			}	
		1	1	{	{	}	)
			j	}	)	}	
		1	}	}	ì		
		1	İ		ĺ		
		1	l	ĺ	[	]	]
		1	İ	{	{	{	
		1	İ	[	ĺ		
		Ì	ļ		ĺ	]	]
		(	{	(	(		,
		İ	1	{	{	1	}
		<b>}</b>	{	1	{	{	}
		ł	1	}	ĺ	{	}
		1	1		)	j	1
		}	}	ł	ļ	•	1
		1	1	ł	İ	1	
		1	}	]	Ì	j	
		1	[	[	l		
		(	1	ĺ	[	[	
l		(	{	{	{	{	}
		1	1	{	(	[	
		1	Ì	ĺ	į	1	
<b>S</b>		{	{	{	(	{	1
l		1	l	ſ	<b>{</b>	<b>f</b>	
Į .	·	1	}	Ì	<b>S</b>	<b>}</b>	
1		1	1	Ì	<b>[</b>	[	
1		}	}	}	]	İ	ĺ
l		1	1	ł	}	•	
[		1	l	l	l	1	
(		1	ł	l	1	i	
(		1	l	{	{	{	
į			l	1	(		
[		{	<b>{</b>	{	{	{	
{			1	1	{	}	
(			ł	ł	}	1	
L		<u> </u>	L				بسيبسيا

DD . 1974. 1473

(BACK)

UNCLASSIFIED

5/N 0102-014-6800

Security Classification

A- 31

## Failure of Methylprednisolone to Protect Lead-Sensitized Rats Against Endotoxin

ROBERT B. JONES, JAMES L. WISE, AND LUTZ A. KIESOW L.

Experimental Medicine Division, Naval Medical Research Institute, Bathesda, Matyland 20014

Received for publication 27 June 1973

FEB 11 1974

Methylprednisolone, in doses that protect normal rats against endotoxin, has no effect in lead-sensitized rats.

TABLE 1. Effect of lead acetate and

methylprednisolone on endotoxin lethality in rats

The administration of lead acetate (PbAc<sub>2</sub>) sensitizes both rats (7) and subhuman primates (4) to minute amounts of bacterial endotoxin. The mechanism of this sensitization has not been defined, although it has been suggested that alterations in either the degradation of endotoxin (8), or in carbohydrate metabolism (3, 4), may be important.

Glucocorticoids are demonstrably effective in protecting normal animals against endotoxin (1, 5). The purpose of the present investigation was to determine if this was also the case in

lead-sensitized animals.

Under light ether anesthesia, femoral cutdowns were performed on male Sprague-Dawley rats weighing 180 to 220 g, and an intravenous injection of 20 mg of PbAc2 dissolved in 0.5 ml of deionized water given. This was immediately followed by an injection of 0.5 ml of Serratia marcescens endotoxin (Difco Laboratories) suspended in 0.15 M NaCl buffered to pH 7.4 with 0.02 M sodium phosphate (PBS), after which 9.5 mg of methylprednisolone (Upjohn Co.) dissolved in 0.5 ml of PBS was also given. Deionized water was used as a control for the PhAc, injections in these experiments. However, other investigators have used sodium acetate with no effect on endotoxin induced mortality (2). PBS was used as a control for the endotoxin and methylprednisolone injections.

The animals were observed for 72 h. although most died within the first 12 h. The mean lethal dose was determined for each group according to the method of Litchfield and Wilcoxon (6). No deaths out of 16 were observed in control animals which received only PbAc, or only methylprednisolone. Two deaths occurred in the 16 animals which received both PbAc, and methylprednisolone, but no endotoxin.

As can be seen from Table 1, methylprednisolone was quite effective in protecting non-lead-treated rats, a single injection causing a fivefold

	Methyl predniso- lone (50 mg/kg°)	No. of animals	Endotoxin LD <sub>50</sub> ° (mg/kg°)
-	-	65	16.5
			(11.6-23.6) <sup>c</sup>
-	+	64	82.7
			(66.4-102.9)
+		152	$0.45 \times 10^{-3}$
			$(0.34 \times 10^{-3} \text{ to } 0.58 \times 10^{-3})$
†	+	178	0.38 × 10 <sup>-3</sup>
			$(0.31 \times 10^{-3} \text{ to } 0.46 \times 10^{-3})$

4 Body weight.

Abbreviation: LD, mean lethal dose.

 $^{\circ}$  Numbers in parenthesis are the 95% confidence limits for each LD  $_{\bullet0}.$ 

increase in the mean lethal dose. However, in the lead-treated rats, which were approximately 37,000 times more sensitive to endotoxin, the methylprednisolone was without any effect.

This failure of a potent glucocorticoid to protect lead-sensitized rats against endotoxin suggests that lead may produce important qualitative, as well as quantitative, differences in the response of an animal to endotoxin. Furthermore, it would seem to indicate that under certain circumstances the efficacy of steroids in the treatment of septic shock may be a function of other, seemingly unrelated, factors.

We gratefully acknowledge the excellent technical assistance of Stanley Shapiro.

This work was supported by the Bureau of Medicine and Surgery, Navy Department Subtask MR011,0001,002,0007

## LITERATURE CITED

 Chedid, L., F. Boyer, and M. Saviard. 1951. Action de la cortisone vis a vis de l'infection experimental avec Salmonella typhi chez la souris. C. R. Acad. Sci. (Paris) 233:713-716.

- 2. Filkins, J. P. 1973. Effects of lead acetate on sensitivity to shock, intravascular carbon and endotoxin clearances,
- shock, intravascular carbon and endotoxin clearances, and hepatic endotoxin detoxification. Proc. Soc. Exp. Biol. Med. 142:471–475.
   Filkins, J. P. 1973. Hypoglycemia and depressed hepatic gluconeogenesis during endotoxicosis in lead-sensitized rats. Proc. Soc. Exp. Biol. Med. 142:915-918.
   Holper, K., R. A. Trejo, L. Brettschneider, and N. R. DiLuzio. 1973. Enchancement of endotoxin shock in the lead sensitized subhuman primate. Surg. Canad. Ob.
- lead-sensitized subhuman primate. Surg. Gynecol. Obstet. 136:593-601.

  5. Kass, E. H. 1960. Effect of corticosteroids and of hormones

- of pregnancy on the lethal action of bacterial endotoxin.
  Ann. N.Y. Acad. Sci. 88:107-115.
  6. Litchfield, J. T., Jr., and F. Wilcoxon. 1949. A simplified method of evaluating dose-effect experiments. J. Pharmacol. Exp. Ther. 96:99-113.
  7. Selye, H., B. Fuchweber, and L. Bertók. 1966. Effect of
- lead acetate on the susceptibility of rats to bacterial
- endotoxins, J. Bacteriol, \$1:884-890.

  8. Trejo, R. A., and N. R. DiLuzio, 1971. Impaired detoxification as a mechanism of lead acetate induced hypersensitivity to endotoxin. Proc. Soc. Exp. Biol. Med. 136:889-893.